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RESEARCH ARTICLE

Diaphragm sniff ultrasound: Normal values, relationship with sniff nasal pressure and accuracy for predicting respiratory involvement in patients with neuromuscular disorders

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Abstract

Background

In patients with neuromuscular disorders, assessment of respiratory function relies on forced vital capacity (FVC) measurements. Providing complementary respiratory outcomes may be useful for clinical trials. Diaphragm sniff ultrasound (US) is a noninvasive technique that can assess diaphragm function that may be affected in patients with neuromuscular disorders.

Purpose

We aimed to provide normal values of sniff diaphragm ultrasound, to assess the relationship between sniff diaphragm US, vital capacity (VC) and sniff nasal pressure. Additionally, we aimed to evaluate the diagnostic accuracy of sniff diaphragm US for predicting restrictive pulmonary insufficiency.

Materials and methods

We included patients with neuromuscular disorders that had been tested with a sniff diaphragm US and functional respiratory tests. Healthy subjects were also included to obtain
normal diaphragm sniff ultrasound. We performed diaphragm tissue Doppler imaging (TDI) and time movement (TM) diaphragm echography combined with sniff maneuver.

Results
A total of 89 patients with neuromuscular diseases and 27 healthy subjects were included in our study. In patients, the median age was 32 years [25; 50] and the median FVC was 34% of predicted [18; 55]. Sniff diaphragm motion using TM ultrasound was significantly associated with sniff nasal pressure, both for the right hemidiaphragm (r = 0.6 p <0.0001) and the left hemidiaphragm (r = 0.63 p = 0.0008). Right sniff peak TDI velocity was also significantly associated with FVC (r = 0.72, p<0.0001) and with sniff nasal pressure (r = 0.66 p<0.0001). Sniff diaphragm ultrasound using either TM mode or TDI displayed significant accuracy for predicting FVC<60% with an area under curve (AUC) reaching 0.93 (p<0.0001) for the right sniff diaphragm ultrasound in TM mode and 0.86 (p<0.001) for right peak diaphragm TDI velocity.

Conclusion
Sniff diaphragm TM and TDI measures were significantly associated with sniff nasal pressure. Sniff diaphragm TM and TDI had a high level of accuracy to reveal respiratory involvement in patients with neuromuscular disorders. This technique is useful to assess and follow up diaphragm function in patients with neuromuscular disorders. It may be used as a respiratory outcome for clinical trials.

Introduction
Respiratory and cardiac failures are the main causes of morbidity and mortality in patients with neuromuscular disorders. Assessment and monitoring of respiratory function relies on functional pulmonary tests. Forced vital capacity (FVC) is used as an indicator of global respiratory function. However, vital capacity (VC) is a late indicator of respiratory muscle involvement in neuromuscular disorders since VC may be normal until a significant decrease in respiratory muscle strength occurs [1]. Sniff nasal pressure and maximal inspiratory pressures, tests used to assess inspiratory muscle strength, may be affected despite a normal FVC [2, 3]. Providing complementary respiratory outcomes may be useful for clinical trials. Among the respiratory muscles, the diaphragm is the main inspiratory muscle. Its assessment classically relies on a trans-diaphragmatic pressure measurement. Since this test is invasive and not comfortable, it cannot be used routinely. Recent research has focused on ultrasound to noninvasively assess diaphragm muscle function [4, 5, 6]. Other techniques have been suggested to assess respiratory muscle function. These techniques include a ratio of maximal expiratory and inspiratory pressures (PEmax/PImax) [7] and the maximum relaxation rate (MRR) of inspiratory muscles [8]. Diaphragm function can be assessed using ultrasound at rest, during deep inspiration or during a sniff maneuver [9]. However, this technique is limited in the assessment of the left diaphragm during deep inspiration, due to loss of the ultrasound picture of the left diaphragm during deep inspiration. Sniff is a physiological maneuver that is used during fluoroscopy for the assessment of diaphragm function [10], it can be used to investigate motion of the right and left diaphragm, and it can be coupled with ultrasound. Additionally, in cardiology, tissue Doppler imaging (TDI) is used to assess subclinical myocardial impairment [11, 12]. Diaphragm sniff ultrasound may be affected earlier in patients with neuromuscular disorders. Little is known about the
potential application of this technique in diaphragm muscle analysis. A subset of patients with muscular dystrophy may have subclinical diaphragm impairment with normal VC values. In this study, we aimed to complete the following:

- To provide normal values for diaphragm peak sniff velocity using TDI and sniff diaphragm motion using time movement (M mode) ultrasound.
- To evaluate the relationship between diaphragm sniff ultrasound and sniff nasal pressure, as well as the relationship between sniff ultrasound and FVC.
- To determine whether peak TDI velocities may be reduced in neuromuscular patients with normal FVC.
- To assess the diagnostic accuracy of diaphragm sniff TDI and sniff M mode ultrasound for predicting impairment of respiratory muscles among patients with neuromuscular disorders.

**Materials and methods**

We retrospectively included patients with neuromuscular disorders followed at Raymond Poincare hospital, a tertiary neuromuscular reference center specialized in the cardiac and pulmonary management of patients with neuromuscular disorders. We included patients who benefited from a sniff diaphragm ultrasound (right and/or left) and we recorded the functional respiratory tests available that included sniff nasal pressure and/or FVC. Diaphragmatic ultrasound was performed in patients with echocardiography, by the same operator (AF).

We recorded the following parameters from the medical records: age, sex, body mass index (BMI), type of disease, peripheral skeletal muscle insufficiency (Walton score), cardiac drugs, steroids, left ventricular ejection fraction (LVEF), history of documented sleep apnea (from polygraphy or polysomnography), functional lung tests that included forced FVC, decubitus FVC, forced expiratory volume in one second (FEV1), inspiratory capacity (IC), expiratory residual volume (ERV), maximal inspiratory pressure (MIP), maximal expiratory pressure (MEP), sniff nasal pressure and arterial partial carbon dioxide pressure (PaCO2). The study was performed in compliance with the ethical principles formulated in the Declaration of Helsinki and was approved by the French National Agency regulating Data Protection (commission nationale de l'informatique et des libertés). All data were fully anonymized before analysis. We also included a healthy control group that underwent diaphragm ultrasound imaging, to obtain reference diaphragm ultrasound sniff values.

**Lung function tests**

Pulmonary function testing was performed in all patients as a part of the routine evaluation, including spirometry that was performed according to ATS/ERS recommendations[13], using a Vmax 229 Sensormedics System (Yorba Linda, CA, USA) with the patient in the upright position [14]. Measurements are expressed as percent of predicted values [15]. Pulmonary function and pressure measurements were performed by different experimental technicians who worked in the same respiratory muscle laboratory.

**Sniff nasal pressure and maximal inspiratory pressure and expiratory pressure measurements**

Maximal inspiratory pressure (MIP) and sniff nasal pressure (SNIP) were both measured from functional residual capacity (FRC) in a standard manner (sitting position) and maximal expiratory pressure (MEP) measured from total lung capacity, according to previously described
methods. MIP is an isometric maneuver, while SNIP is a quasi-isometric maneuver [3, 16].
MIP was measured with a flanged mouthpiece with the maneuvers repeated at least four times
or until two identical readings were obtained [17]. SNIP was measured during at least 10 and
up to a maximum of 20 sniffs in a standard manner according to previously described methods
[3]. Briefly, the plug used to obstruct the nostril was an eartip usually used for auditory evoked
potentials (Eartips, 13 mm, Nicolet, Madison, WI). This plug was connected to a pressure
transducer via a catheter (see below). The length of this catheter was reduced to the minimum
length possible. Air leak was detected by obstructing the other nostril during an inspiratory
maneuver and when present it was eliminated by adding waxed earplug material. Detailed
instruction on how to perform the sniff maneuver was not necessary and may be counterpro-
ductive [18]. Patients were vigorously coached during the test maneuvers.

All pressure signals were measured using a differential pressure transducer (Validyne,
Northridge, CA), amplified by a carrier amplifier (Validyne), and passed through an analog-
digital board to a computer running Acqknowledge software (Biopac System, Santa Barbara,
CA) which allows visual feedback to improve the sniff efficiency. The signal was digitized at
100 Hz. Subjects received strong verbal encouragement with visual feedback, as previous stud-
ies have suggested [19]. Values of MIP, MEP and SNIP were expressed in cmH₂O.

Diaphragm ultrasound technique

Diaphragm ultrasound was performed in the supine position by the same operator (AF), who
was blinded to the pulmonary function tests. We used the liver window for the right hemi-
diaphragm analysis and the spleen window for the left hemidiaphragm analysis. The trans-
ducer was placed in the anterior subcostal region between the midclavicular and anterior
axillary lines so that the ultrasound beam reached the posterior part of the diaphragm. After
visualization of the diaphragm, an M mode was applied perpendicularly to the hemidiaphragm
to measure diaphragm motion during inspiration and during a sniff maneuver. Patients were
in a semirecumbent position (45˚) during the procedure. In normal subjects, the normal inspi-
ratory motion is caudal and the operator recorded the M mode trace moving upward (dia-
phragm moving toward the probe). With the sniff maneuver, the operator recorded a sharp
upstroke in the normal situation due to the high speed of diaphragm movement (Fig 1). For

![Diaphragm ultrasound displacement from the subcostal view in a control subject and in a DMD patient](https://doi.org/10.1371/journal.pone.0214288.g001)
sniff peak diaphragm TDI recording, we used the liver window or the spleen window. Using a cardiac probe, we activated the tissue Doppler imaging modality with the beam positioned perpendicular to the diaphragm motion; then, we recorded the peak sniff inspiratory velocity (cm/s) (Figs 2 and 3). Diaphragm thickness was also recorded from the apposition zone ultrasound, using a linear probe positioned in the midaxillary line perpendicular to the chest wall. Thickness was measured at the end of a quiet expiration.

**Statistical analysis**

Continuous variables are reported as the median ± interquartile range [IQR] and compared using nonparametric tests due to their distribution; categorical variables are described by number of subjects and percentage and compared by Fisher’s exact test. Spearman correlations were performed to evaluate associations between continuous variables. Discrimination performances of sniff M mode and sniff TDI regarding FVC < 60% and FVC < 30% were assessed using receiver-operator characteristic (ROC) analyses, and corresponding ROC curves were drawn. C-statistics corresponded to the areas under corresponding ROC curves. Figures were created and statistical analyses were performed using GraphPad Prism (GraphPad Software, Inc., State of California, USA).

**Results**

**Clinical and respiratory functional tests and diaphragm ultrasound**

A total of 89 patients with genetically confirmed neuromuscular diseases were included in our study (35% Duchenne muscular dystrophy (DMD), 29% myotonic dystrophy type 1, 12% sarcoglycanopathies). The median age was 32 years [25; 50] and the median BMI was 22 kg/m².
In this sample, 57% of patients were wheelchair bound. Patients disclosed a restrictive respiratory insufficiency pattern with a median FVC of 34% of predicted [18; 55]. In total, 63% of patients were on home mechanical ventilation (HMV). Inspiratory muscles were affected as attested by a median inspiratory capacity (IC) of 59% of predicted [16; 74], a median MIP of 22 cmH20 [14; 32] and a median sniff nasal pressure of 22 cmH20 [14;32]. Diaphragm ultrasound parameters were significantly reduced in patients. Table 1 summarizes the clinical and respiratory data in the patients. Table 2 summarizes the diaphragm ultrasound data in the patients vs the control group.

**Relationship between sniff ultrasound and sniff nasal pressure**

Diaphragm sniff time movement (TM) ultrasound motion was significantly associated with sniff nasal pressure, both for the right hemidiaphragm (r = 0.6 p <0.0001) and the left hemidiaphragm (r = 0.63 p = 0.0008) (Fig 4). Indeed, diaphragm peak sniff TDI velocities were significantly associated with sniff nasal pressure (Fig 5).

**Relationship between diaphragm sniff ultrasound and forced vital capacity**

Using the global respiratory function assessment, diaphragm sniff ultrasound TM motion was significantly associated with FVC. This finding involved the right hemidiaphragm as well as the left hemidiaphragm (Figs 6 and 7). We also compared sniff M mode with FVC expressed in ml and observed a similar relationship to FVC expressed in %, which takes into account height (Fig 8).

Right sniff peak TDI was also significantly associated with FVC (r = 0.72, p<0.0001) (Fig 9). Supine FVC was also significantly associated with sniff diaphragm right TM motion (r = 0.59, p = 0.0003, n = 34). Additionally, Supine FVC was significantly associated with right
diaphragm sniff TDI velocity (r = 0.59, p = 0.0007, n = 29). The level of the relationship between the sniff diaphragm TM motion and FVC remained similar in patients with DMD vs patients with DM1 vs patients with other myopathies (Fig 10).

**Sniff diaphragm TDI velocities and TM motion in patients with FVC>80%**

Figs 6, 7 and 9 provide an overview of the distribution of the right sniff TM motion and the right sniff peak TDI velocity in patients with FVC>80%. Among those with FVC>80%, 5 (83.3%) patients had a right diaphragm sniff TM motion under the median value obtained in controls. In patients with FVC>80%, 3 (60%) patients had right sniff TDI peak velocity under the median right diaphragm sniff TDI peak velocity obtained in the control group.

### Table 1. Clinical and respiratory data in patients.

| Parameters                          | Number (%) or median [IQR] |
|-------------------------------------|----------------------------|
| Neuromuscular Diseases (N = 89)     |                            |
| DMD                                 | 31 (35%)                   |
| DM1                                 | 26 (29%)                   |
| Sarcoglycanopathy                   | 11 (12%)                   |
| Mitochondrial disease               | 4 (4.5%)                   |
| Congenital myopathy                 | 4 (4.5%)                   |
| Spinal muscular atrophy             | 3 (3%)                     |
| Pompe disease                       | 2 (2%)                     |
| Desminopathy                        | 1 (1%)                     |
| Other diseases                      | 7 (8%)                     |
| Age (y)                             | 32 [25;50]                 |
| Wheelchair bound                    | 51 (57%)                   |
| Walton score                        | 6 [3;7]                    |
| BMI (kg/m²)                         | 22 [14; 27]                |
| Sleep apnea (N, %)                  | 24 (27%)                   |
| FVC (%)                             | 34 [18;55]                 |
| Decubitus FVC (%)                   | 30 [18;47]                 |
| IC (%)                              | 59 [16;74]                 |
| FEV1 (ml)                           | 1280 [530;1765]            |
| FEV1/VC                             | 91 [81;97]                 |
| PEF (l/s)                           | 3 [2;4]                    |
| ERV (ml)                            | 420 [110;640]              |
| PaCO2 (mmHg)                        | 46 [42;50]                 |
| MIP (cmH20)                         | 22 [14;32]                 |
| MEP (cmH20)                         | 22 [15;36]                 |
| Sniff nasal pressure (cmH20)        | 22 [14;32]                 |
| HMV                                 | 56 (63%)                   |

Data are presented as number (percentage) or median [IQR].
BMI: body mass index; IC: inspiratory capacity (%); PEF: peak expiratory flow; FEV1: forced expiratory volume in one second; MIP: maximal inspiratory pressure; MEP: maximal expiratory pressure; PaCO2: arterial partial carbon dioxide pressure (mmHg); FVC: forced vital capacity (%); HMV: home mechanical ventilation; DMD: Duchenne muscular dystrophy; DM1: myotonic dystrophy type I. Y = years.

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Accuracy of sniff ultrasound for the prediction of respiratory muscle impairment in patients with muscular dystrophy

Using receiver operating characteristic (ROC) curves, sniff diaphragm ultrasound using either TM mode or TDI accurately discriminated those with FVC < 60%; the area under the curve (AUC) reached 0.93 (p < 0.0001) for the sniff right TM mode diaphragm ultrasound and 0.86 (p < 0.001) for right peak diaphragm TDI velocity. In patients with significant respiratory insufficiency (FVC < 30%), the AUC remained high but with a slight decrease in the AUC of the right diaphragm ultrasound TDI (0.76, p < 0.017). The M mode right diaphragm motion value cutoff to predict a FVC < 60% was 25 mm with a sensitivity of 100% and a specificity of 64%. The M mode right diaphragm motion value cutoff to predict a FVC < 30% was 10.5 mm with a sensitivity of 84% and a specificity of 84%.

Table 2. Diaphragm ultrasound in the patient and control groups.

| Ultrasound parameters               | Patient group (N = 89) | Control group (N = 27) | p    |
|-------------------------------------|-----------------------|------------------------|------|
| Sex (F)                             | 11 (12%)              | 14 (50%)               | 0.14 |
| Age (y)                             | 32 [25;50]            | 31 [26;59]             | 0.43 |
| BMI (kg/m²)                         | 22 [14;27]            | 22 [20;25]             | 0.46 |
| Right deep inspiratory motion (mm)  | 23 [10;38]            | 72 [62;88]             | <0.001|
| Left deep inspiratory motion (mm)   | 23 [11;36]            | 62 [61;97]             | 0.002|
| Right sniff TM diaphragm motion (mm)| 12 [6;19]             | 32 [19;36]             | <0.001|
| Left sniff TM diaphragm motion (mm) | 13 [9;18]             | 33 [23;42]             | 0.001|
| Peak right sniff TDI diaphragm velocity (cm/s) | 6 [4;9]          | 13 [10;14]             | <0.001|
| Peak left sniff TDI diaphragm velocity (cm/s) | 7 [5.5;11]         | 12 [11;13]             | 0.06 |
| Right diaphragm thickness (mm)*     | 1.8 [1.5;2.1]         | 1.5 [1.3;1.9]          | 0.11 |
| Left diaphragm thickness (mm)*      | 1.6 [1.4;2]           | 1.5 [1.1;1.9]          | 0.25 |

Data are presented as number (percentage) or median [IQR].

F: female; BMI: body mass index; TM: time movement; TDI: tissue Doppler imaging.

*: at rest. p = p value.

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Accuracy of sniff ultrasound for the prediction of respiratory muscle impairment in patients with muscular dystrophy

Using receiver operating characteristic (ROC) curves, sniff diaphragm ultrasound using either TM mode or TDI accurately discriminated those with FVC < 60%; the area under the curve (AUC) reached 0.93 (p < 0.0001) for the sniff right TM mode diaphragm ultrasound and 0.86 (p < 0.001) for right peak diaphragm TDI velocity. In patients with significant respiratory insufficiency (FVC < 30%), the AUC remained high but with a slight decrease in the AUC of the right diaphragm ultrasound TDI (0.76, p < 0.017). The M mode right diaphragm motion value cutoff to predict a FVC < 60% was 25 mm with a sensitivity of 100% and a specificity of 64%. The M mode right diaphragm motion value cutoff to predict a FVC < 30% was 10.5 mm with a sensitivity of 84% and a specificity of 84%.
Fig 5. Relationship between right diaphragm sniff peak TDI velocity and sniff nasal pressure in patients. Sniff TDI peak inspi = sniff peak diaphragm tissue Doppler imaging velocity.

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Fig 6. Relationship between right sniff TM mode and FVC in patients. FVC: forced vital capacity (%).

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Fig 7. Relationship between left sniff TM mode and FVC in patients. FVC: forced vital capacity (%).
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Fig 8. Relationship between right sniff TM mode and FVC (ml) in patients. FVC: forced vital capacity.
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Fig 9. Relationship between right sniff TDI peak velocity (cm/s) and FVC in patients. FVC = forced vital capacity (%).

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Fig 10. Relationship between right sniff diaphragm TM motion and FVC in DMD, DM1 and other myopathies. DMD = Duchenne muscular dystrophy; DM1 = myotonic dystrophy type I; others = other myopathies; FVC = forced vital capacity (%); sniff M mode = right sniff diaphragm time movement motion (mm).

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The right peak TDI velocity value cutoff to predict a FVC<60% was 7.5 cm/s with a sensitivity of 84% and a specificity of 89%. The right peak TDI velocity value cutoff to predict a FVC<30% was 6.5 cm/s with a sensitivity of 90% and a specificity of 56%. Figs 11 and 12 show the ROC curves of the sniff diaphragm ultrasound for predicting FVC<60% and for predicting FVC<30%, respectively.

Discussion

In the present study, we describe for the first time the diaphragmatic sniff ultrasound pattern in patients with muscular dystrophy, investigating the relationship between sniff diaphragm ultrasound and sniff nasal pressure as well as the relationship with FVC. We also report the accuracy of sniff diaphragm ultrasound for predicting moderate and/or severe respiratory muscle impairment. The main findings are the following:

- Sniff diaphragm ultrasound motion using TM mode was significantly reduced in patients with myopathies and was correlated with FVC and sniff nasal pressure.
- Sniff peak diaphragm ultrasound TDI velocity was significantly reduced in patients with myopathies and was correlated with FVC and sniff nasal pressure.

![ROC curves](https://doi.org/10.1371/journal.pone.0214288.g011)

**Fig 11.** ROC (receiver operating characteristic) curves for predicting a FVC<60% in patients with neuromuscular disorders. TDI right: peak sniff TDI velocity (cm/s) at the right hemidiaphragm; M mode left: left diaphragm motion during a sniff maneuver (mm); M mode right: right diaphragm motion during a sniff maneuver (mm); FVC = forced vital capacity (%).
Diaphragm peak sniff TDI velocities and diaphragm sniff TM motion may be impaired in patients even with FVC > 80%.

Lung volumes in patients with neuromuscular disorders are affected by weakness of the respiratory muscles [20], and FVC provides a global evaluation of respiratory muscle function. In patients with NMD, the classic pattern of restriction is a reduction of FVC [21]. FRC may be normal or decreased, and RV may increase [22]. In our study, FVC was significantly associated with sniff diaphragm ultrasound. Interestingly, we found that sniff diaphragm ultrasound may predict restrictive impairment in patients with muscular dystrophy and was associated with sniff nasal pressures. Classically, MIP and sniff nasal pressures have been used to assess inspiratory muscle strength in respiratory laboratories [20]. Low values may be due to muscle failure, however, they may also be due to a lack of motivation, lack of coordination, peribuccal leaks and fatigue [23]. The sniff test is a natural effort maneuver that solves leak problems, reduces the risk of fatigue and can be combined with diaphragm ultrasound. However, the sniff value can be underestimated in cases of nasal obstruction [23]. Our data support the fact

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Fig 12. ROC curves for predicting a FVC < 30% in patients with neuromuscular disorders. M mode right: right diaphragm motion during a sniff maneuver; M mode left: left diaphragm motion during sniff maneuver (mm); TDI right: peak sniff TDI velocity (cm/s) recorded at the right hemidiaphragm; FVC: forced vital capacity (%).

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that sniff TDI can be used to detect early subclinical diaphragm involvement in patients with neuromuscular diseases. Indeed, the usefulness of TDI is to depict early muscle involvement in a pathological setting. Similar to the way TDI is used in cardiology to detect early myocardopathy before the onset of a reduced ejection fraction [12], TDI may be used in pneumology and combined with sniff maneuver to detect early muscle involvement in myopathy before the onset of reduced FVC. In the early stages of diaphragm involvement, FVC remains normal, since FVC is a late marker of respiratory muscle involvement. In our study, in the control group, the median normal peak sniff TDI was 13 cm/s for the right hemidiaphragm and 12 cm/s for the left hemidiaphragm, and some patients had a reduced peak sniff TDI velocity, and at the same time, had a forced VC that was in the normal range. These findings are interesting since they could prompt monitoring in patients with abnormal sniff TDI. Patients with neuromuscular disorders presenting abnormal diaphragm peak TDI velocity may benefit from close pulmonary test monitoring. In normal subjects, during the sniff maneuver, inspiratory muscles are short at higher speed, and the pressure measured in the esophagus is closely related to the pressure in the mouth, nasopharynx and nose [24]. Heritier et al [3] reported an excellent correlation between sniff nasal pressure and sniff esophageal pressure (r 0.99, p < 0.001). Diaphragm ultrasound emerged recently in the literature as a noninvasive method to assess diaphragm function [25, 26]. Our data in the control group regarding sniff TM diaphragm motion were similar, even higher than the data reported by Boussuges et al [27], 16 mm in women and 18 mm in men [27]. Various factors (sex and height) may influence normal diaphragm ultrasound, as reported by Scarlata et al [25]. In our study, we also compared sniff M mode with FVC expressed in ml and observed a similar relationship to FVC expressed in %, which takes into account height. No data have been reported regarding sniff diaphragm TDI velocities in the literature. In our study, the correlation between peak sniff TDI velocity and sniff nasal pressure was high (r = 0.66 p < 0.0001) but was relatively reduced in comparison with the correlation between sniff esophageal pressure and sniff nasal pressure reported by Heritier et al [3] (r = 0.99). This discrepancy may be explained by the fact that the peak sniff TDI velocity recorded the contribution of one side of the diaphragm and the peak velocity may be influenced by the pressure interactions between the chest and abdomen, the geometry and pattern of the ribcage. Moreover, the sniff maneuver used during ultrasound was different from the method used in pulmonary laboratories since the patient has two nostrils open during diaphragm sniff ultrasound examinations. Finally, the AUC for sniff diaphragm ultrasound using TDI or TM mode was significantly high for predicting FVC <60% and/or FVC <30%. However, in patients with FVC <30%, the sniff right TDI velocity displayed a lower AUC (0.76, p 0.017), in comparison with sniff right TM mode diaphragm displacement (AUC 0.86, p <0.0001). This difference can be explained by the lower TDI velocity recorded for the dia-

**Clinical perspective**

The assessment and monitoring of diaphragm function requires dedicated equipment and specialized techniques, which are not available in all centers. A single sniff diaphragm ultrasound measuring using TDI or TM mode can help physicians detect patients with asymptomatic restrictive pulmonary insufficiency and early diaphragm involvement in patients with neuromuscular disorders. Diaphragm sniff TDI may be used as a presymptomatic test to reveal early diaphragm involvement in patients with neuromuscular disorders. These findings may help clinicians select patients for regular respiratory test evaluation during follow-up, particularly in patients with TDI velocity impairment. Our study provided normal sniff diaphragm ultrasound values. These findings can be used as reference values for other studies in the
neuromuscular field. Sniff diaphragm ultrasound can be used as a screening test and could be used as a monitoring tool for future biotherapy assessment. In patients with significant respiratory insufficiency, it would be appropriate to focus on the right TM mode technique for monitoring patients. Finally, studies that assess economic performance and nursing technical requirements need to be performed with this new technique. In the future, this radiological technique may be considered as an additional test in the arsenal of pulmonary exploratory functional tests used in a multiparametric approach in the neuromuscular field.

**Limits**

This study is limited by its retrospective design. Additionally, sniff ultrasound is a volitional test and data depend on patient motivation and fatigue. The other limitation relies on the technical aspect of Doppler imaging. Indeed, velocity recorded with TDI depends on beam angulation and the beam direction should be perpendicular to diaphragm displacement to obtain the maximal TDI peak velocity. Moreover, patient position may affect respiratory function tests [28]. Here, with US, patients were in a semirecumbent position (45°) that would amplify diaphragm dysfunction and US measurements [27, 29]. The supine position is preferred, because there is less overall variability, less side-to-side variability, and greater reproducibility. Additionally, diaphragm excursion is known to be greater in the supine position for the same volume inspired than in the sitting or standing positions because the abdominal viscera more easily moves the diaphragm in this position, and the relationship between inspired volume and diaphragm movement has been shown to correlate better in the supine position than in the sitting position. The supine position also exaggerates any paradoxical movement and limits any compensatory active expiration by the anterior abdominal wall, which may mask paralysis [4]. According to Brown et al [29], intraobserver reliability was excellent (>0.93) for all body positions tested. Because many of our patients were not able to maintain the strict supine position for different reasons (e.g., tolerability), we systematically used the semirecumbent position (45°). Nevertheless, despite the difference in position between respiratory function measurements (90°) and diaphragmatic ultrasound measurements (45°), significant correlations were observed. Moreover, we compared nasal pressure and diaphragm velocity and motion. These latter measures depend on other factors that include abdominal pressure and compartment and history of abdominal surgery, and we did not have esophageal pressure for the reference method. Finally, nasal pressure does not necessarily reflect esophageal pressure in some patients with neuromuscular disease due to the inability to generate a significant transnasal pressure [30].

**Conclusion**

Sniff diaphragm TM and TDI are significantly associated with sniff nasal pressure. Sniff diaphragm TM and TDI had high accuracy for depicting respiratory involvement in patients with neuromuscular disorders. This noninvasive diaphragm evaluation can reveal early diaphragm involvement. These techniques are useful to assess and follow up diaphragm function in patients with neuromuscular disorders and may be used as a respiratory outcome for clinical trials.

**Author Contributions**

**Conceptualization:** Abdallah Fayssoil, Bernard Clair.

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References

1. Braun NM, Arora NS, Rochester DF. Respiratory muscle and pulmonary function in polymyositis and other proximal myopathies. Thorax. 1983 Aug; 38(8):616–23. PMID: 6412385

2. Fauroux B, Aubertin G, Cohen E, Clément A, Lofaso F. Sniff nasal inspiratory pressure in children with muscular, chest wall or lung disease. Eur Respir J. 2009 Jan; 33(1):113–7. https://doi.org/10.1183/09031936.00050708 PMID: 18799509

3. Hérilier F, Rahm F, Pasche P, Fitting JW. Sniff nasal inspiratory pressure. A noninvasive assessment of inspiratory muscle strength. Am J Respir Crit Care Med. 1994 Dec; 150(6 Pt 1):1678–83

4. Sarwal A, Walker FO, Cartwright MS. Neuromuscular ultrasound for evaluation of the diaphragm. Muscle Nerve. 2013 Mar; 47(3):319–29 https://doi.org/10.1002/mus.23671 PMID: 23382111

5. Haji K, Royse A, Green C, Botha J, Canty D, Royse C. Interpreting diaphragmatic movement with bedside imaging, review article. J Crit Care. 2016 Aug; 34:56–65 https://doi.org/10.1016/j.jcrc.2016.03.006 PMID: 27288611

6. Boon AJ, Harper CJ, Ghahfarokhi LS, Strommen JA, Watson JC, Sorenson EJ. Two-dimensional ultrasound imaging of the diaphragm: quantitative values in normal subjects. Muscle Nerve. 2013 Jun; 47(6):884–9 https://doi.org/10.1002/mus.23702 PMID: 23625789

7. Fregonezi G, Azevedo IG, Resqueti VR, De Andrade AD, Gualdi LP, Aliverti A, et al. Muscle impairment in neuromuscular disease using an expiratory/inspiratory pressure ratio. Respir Care. 2015 Apr; 60(4):533–9. https://doi.org/10.4187/respcare.03367 PMID: 25587161

8. Evangelista MA, Dias FAL, Dourado Júnior MET, do Nascimento GC, Sarmento A, Gualdi LP, et al. Noninvasive assessment of respiratory muscle strength and activity in Myotonic dystrophy. PLoS One. 2017 Jun 8; 12(6):e0177318. https://doi.org/10.1371/journal.pone.0177318 PMID: 28594857

9. Naik LY, Sondekoppam RV, Jenkin Tsui J, Tsui BC. An ultrasound-guided ABCDE approach with a sniff test to evaluate diaphragmatic function without acoustic windows. Can J Anaesth. 2016 Oct; 63(10):1199–200 https://doi.org/10.1007/s12630-016-0685-z PMID: 27324892

10. Kodric M, Trevisan R, Torregiani C, Cifaldi R, Longo C, Cantarutti F, et al. Inspiratory muscle training for diaphragm dysfunction after cardiac surgery. J Thorac Cardiovasc Surg. 2013 Mar; 145(3):819–23. https://doi.org/10.1016/j.jtcs.2012.07.087 PMID: 22938776

11. Cameli M, Mondillo S, Solari M, Righini FM, Andrei V, Contaldi C, et al. Echocardiographic assessment of left ventricular systolic function: from ejection fraction to torsion. Heart Fail Rev. 2016 Jan; 21(1):77–94. https://doi.org/10.1007/s10741-015-9521-8 PMID: 26712329

12. Ommen SR, Nishimura RA, Appleton CP, Miller FA, Oh JK, Redfield MM, et al. Clinical utility of Doppler echocardiography and tissue Doppler imaging in the estimation of left ventricular filling pressures: A comparative simultaneous Doppler-catheterization study. Circulation. 2000 Oct 10; 102(15):1788–94 PMID: 11023933
13. Miller MR, Hankinson J, Brusasco V, Burgos F, Casaburi R, Coates A, et al. Standardisation of lung function testing: the authors’ replies to readers’ comments. The European respiratory journal 2010; 36 (6):1496–1498. https://doi.org/10.1183/09031936.00130010 PMID: 21119213

14. Quanjer PH, Tammeling GJ, Cotes JE, Pedersen OF, Peslin R, Yernault JC. Lung volumes and forced ventilatory flows. Report Working Party Standardization of Lung Function Tests, European Community for Steel and Coal. Official Statement of the European Respiratory Society. Eur Respir J Suppl 1993; 16:5–40. PMID: 8499054

15. Quanjer PH, Stanojevic S, Cole TJ, Baur X, Hall GL, Culver BH, et al. Multi-ethnic reference values for spirometry for the 3-95-yr age range: the global lung function 2012 equations. The European respiratory journal 2012; 40(6):1324–1343. https://doi.org/10.1183/09031936.00080312 PMID: 22743675

16. Black LF, Hyatt RE. Maximal respiratory pressures: normal values and relationship to age and sex. Am Rev Respir Dis 1969; 99(5):696–702. https://doi.org/10.1164/arrd.1969.99.5.696 PMID: 5772056

17. Wilson SH, Cooke NT, Edwards RH, Spiro SG. Predicted normal values for maximal respiratory pressures in caucasian adults and children. Thorax 1984; 39(7):535–538. PMID: 6463933

18. ATS/ERS Statement on respiratory muscle testing. Am J Respir Crit Care Med 2002; 166(4):518–624. https://doi.org/10.1164/rccm.166.4.518 PMID: 12186831

19. Laporta D, Grassino A. Assessment of transdiaphragmatic pressure in humans. J Appl Physiol 1985; 59(8):1469–1476. https://doi.org/10.1152/jappl.1985.59.5.1469 PMID: 3158636

20. Black LF, Hyatt RE. Maximal static respiratory pressures in generalized neuromuscular disease. Am Rev Respir Dis. 1971 May; 103(5):641–5. https://doi.org/10.1164/arrd.1971.103.5.641 PMID: 5579907

21. Vincken WG, Elleker MG, Cosio MG. Flow-volume loop changes reflecting respiratory muscle weakness in chronic neuromuscular disorders. Am J Med. 1987 Oct; 83(4):673–80. PMID: 3674055

22. de Troyer A, Borenstein S, Cordier R. Analysis of lung volume restriction in patients with respiratory muscle weakness. Thorax. 1980 Aug; 35(8):603–10. PMID: 7444828

23. Prigent H, Lejaille M, Falaize L, Louis A, Ruquet M, Fauroux B, et al. Assessing inspiratory muscle strength by sniff nasal inspiratory pressure. Neurocrit Care. 2004; 1(4):475–8. https://doi.org/10.1385/NCC:1:4:475 PMID: 16174953

24. Koulouris N, Mulvey DA, Larocca CM, Sawicka EH, Green M, Moxham J. The measurement of inspiratory muscle strength by sniff esophageal, nasopharyngeal, and mouth pressures. Am Rev Respir Dis. 1989 Mar; 139(3):641–6. https://doi.org/10.1164/ajrccm/139.3.641 PMID: 2923363

25. Sciarlata S, Mancini D, Laudisio A, Benigni A, Antonelli Incalzi R. Reproducibility and Clinical Correlates of Supine Diaphragmatic Motion Measured by M-Mode Ultrasonography in Healthy Volunteers. Respiration. 2018; 96(3):259–266 https://doi.org/10.1159/000489228 PMID: 30114702

26. Boussuges A, Brégeon F, Blanc P, Gil JM, Poiré D. Characteristics of the paralysed diaphragm studied by M-mode ultrasonography. Clin Physiol Funct Imaging. 2018 Oct 16

27. Boussuges A, Gole Y, Blanc P. Diaphragmatic motion studied by m-mode ultrasonography: methods, reproducibility, and normal values. Chest. 2009 Feb; 135(2):391–400 https://doi.org/10.1378/chest.08-1541 PMID: 19017880

28. Lo Mauro A, Aliverti A. Physiology of respiratory disturbances in muscular dystrophies. Breathe (Sheff). 2016 Dec; 12(4):318–327

29. Brown C, Tseng SC, Mitchell K, Roddey T. Body Position Affects Ultrasonographic Measurement of Diaphragm Contractility. Cardiopulm Phys Ther J. 2018 Oct; 29(4):166–172. https://doi.org/10.1097/CPPT.0000000000000083 PMID: 30319315

30. Haight JS, Cole P. The site and function of the nasal valve. Laryngoscope. 1983 Jan; 93(1):49–55. PMID: 8823174