Assessment of the voluntary activation of the diaphragm using cervical and cortical magnetic stimulation

T. Similowski, A. Duguet, C. Straus, V. Attali, D. Boisteanu, J-Ph. Derenne


ABSTRACT: The twitch occlusion technique is a promising tool for use in assessing central drive to the diaphragm and determining maximal transdiaphragmatic pressure (Pdi) from submaximal efforts. Its clinical use is limited by difficulties inherent to bilateral electrical stimulation (BES) of the phrenic nerves. This study was designed to revisit the technique using cervical magnetic stimulation (CMS). In addition, the effects of a voluntary contraction on diaphragm response to magnetic stimulation of the cortex (CxMS) were studied.

Seven volunteers aged 23–33 yrs were studied. Pdi was determined at relaxed functional residual capacity (FRC) in response to BES (Pdi,p-ES) and CMS (Pdi,p-CMS), and the effects of an increasing voluntary contraction (Pdi,vol) were assessed. The same procedure was applied to CxMS.

\[ P_{\text{di},p-\text{CMS}} \text{ at relaxed FRC was } 27.5 \pm 2.2 \text{ cmH}_2\text{O (mean±SEM), about 20% higher than } P_{\text{di},p-\text{ES}}, \text{ as reported previously. } P_{\text{di},p-\text{CMS}} \text{ linearly decreased with } P_{\text{di},\text{vol}}, \text{ and six out of seven subjects were capable of producing voluntary contractions sufficient to extinguish the twitch. More complex patterns were observed with CxMS. }\]

Cervical magnetic stimulation provides diaphragmatic twitch occlusion data very similar to bilateral electrical phrenic stimulation. Magnetic stimulation, be it cervical or cortical, could probably be helpful for the assessment of central and peripheral mechanisms of diaphragmatic dysfunction in the clinical setting.

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The degree to which a skeletal muscle can be activated during attempted maximal voluntary efforts can be assessed using the technique of twitch interpolation. First described by MERTON [1], it involves the superimposition of twitches produced by electrical stimulation on a voluntary contraction. The amplitude of the response to peripheral nerve stimulation decreases monotonically as the strength of the underlying contraction increases. It is commonly held that this contraction stems from maximal activation when it completely suppresses the response to nerve stimulation [1,2]. Co-operative subjects are able to fully activate most limb muscles [1–3], but with regards to the diaphragm this issue has been controversial. A commonly accepted idea is that the diaphragm does not produce its maximal force during inspiratory manoeuvres, but rather during expulsive tasks when there is co-contraction of abdominal muscles [4,5]. Although conflicting results have been published [6], several studies applying the twitch occlusion technique to the diaphragm suggest that normal subjects can voluntarily produce maximal diaphragmatic contraction using inspiratory as well as expulsive efforts [7,8]. This has been demonstrated in a comprehensive and elegant study by GANDEVIA and co-workers [9], who, in addition, first raised the possibility of testing maximal voluntary drive to respiratory muscles by motor cortical stimulation [10].

Besides these physiological considerations, the twitch occlusion technique could be highly relevant clinically, since it could be used to evaluate the intensity of the central command to the diaphragm. Indeed, in animals, estimates of central drive derived from twitch occlusion closely mirror central drive as reflected by recordings of phrenic nerve activity [11]. In humans, the twitch interpolation technique is the only test among those used to assess respiratory muscle function that can separate peripheral from central diaphragmatic fatigue or weakness [12,13]. In this respect, the failure of asthmatic patients to achieve maximal diaphragmatic activation [14], and the link of this failure to depression of mental state [15] have led to the idea that reduced voluntary drive to breathe could be a source of unexplained ventilatory failure in asthma. Conversely, the ability of patients with severe chronic obstructive pulmonary disease (COPD) to activate their diaphragm fully [16] argues against central inhibition of respiration as a mechanism of CO₂ retention or respiratory failure.

Another salient feature of the twitch occlusion technique is that, because the response to nerve stimulation is linearly related to the intensity of the underlying contraction, it permits accurate assessment of maximal muscle strength under conditions where the patient is unable or unwilling to co-operate [7].
Table 1. – Characteristics of the subjects, and transdiaphragmatic pressure data at FRC

<table>
<thead>
<tr>
<th>Subject No.</th>
<th>Age (yrs)</th>
<th>Sex</th>
<th>Height (cm)</th>
<th>Weight (kg)</th>
<th>$P_{di,p-ES}$ (cmH₂O)</th>
<th>$P_{di,p-CMS}$ (cmH₂O)</th>
<th>$P_{di,max}$ (cmH₂O)</th>
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<tbody>
<tr>
<td>1</td>
<td>29</td>
<td>M</td>
<td>165</td>
<td>65</td>
<td>25.2</td>
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<td>166</td>
<td>64</td>
<td>22.6</td>
<td>27.2</td>
<td>123.8</td>
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<tr>
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<td>23</td>
<td>F</td>
<td>163</td>
<td>52</td>
<td>17.6</td>
<td>20.1</td>
<td>79.0</td>
</tr>
<tr>
<td>5</td>
<td>25</td>
<td>M</td>
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<td>19.8</td>
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<tr>
<td>6</td>
<td>33</td>
<td>F</td>
<td>187</td>
<td>85</td>
<td>31.5</td>
<td>35.0</td>
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<td>M</td>
<td>165</td>
<td>70</td>
<td>26.0</td>
<td>32.0</td>
<td>155.0</td>
</tr>
</tbody>
</table>

Mean 23.3 27.5 126.2
SEM 1.9 2.2 16.0

FRC: functional residual capacity; M: male; F: female; $P_{di,p-ES}$: amplitude of the transdiaphragmatic twitch pressure elicited by bilateral electrical phrenic nerve stimulation at relaxed FRC; $P_{di,p-CMS}$: amplitude of the transdiaphragmatic twitch pressure elicited by cervical magnetic stimulation at relaxed FRC; $P_{di,max}$: amplitude of the transdiaphragmatic twitch pressure generated by maximal, static, pure diaphragmatic contractions.

For all these reasons, the twitch occlusion technique has been endorsed by the 1989 National Heart, Lung and Blood Institute (NHLBI) workshop on respiratory muscle fatigue as a potentially major diagnostic tool [17]. However, its usefulness is restricted by difficulties inherent to bilateral supramaximal stimulation of the phrenic nerves [17]. The present study was, therefore, designed to revisit the assessment of the voluntary drive to the diaphragm using noninvasive and easy to use cervical and cortical magnetic stimulations.

Material and methods

Subjects

Seven healthy volunteers (4 males and 3 females; aged 23–33 yrs) participated in the study (table 1), after approval of the Local Ethical Committee. Two of them had previous experience of respiratory muscle experiments, considerable in one case. All subjects were studied sitting on a chair, abdomen unbound. They had been informed of the purpose and methods of the study.

Measurements

Electromyograms. Surface recordings of the right and left costal diaphragmatic electromyographic activity (RAi and LAi, respectively) were obtained using skin-taped silver cup electrodes placed in the 7th and 8th right and left intercostal spaces and connected to a Dantec 2000 electromyograph (Dantec Electronic, Denmark). In four subjects, a surface electromyogram of one sternomastoid muscle was also obtained (Esm) (see Discussion).

Pressures. Oesophageal and gastric pressure ($P_{oes}$ and $P_{ga}$) were measured with two balloon-catheters (thinned-walled balloons sealed over a distally side-holed polyethylene catheter, 50 cm length and 1.7 mm internal diameter (ID)). Transdiaphragmatic pressure ($Pa$) was obtained on-line by connecting these catheters to a Validyne MP45 differential pressure transducer (±200 cmH₂O; Validyne, Northridge, USA). Pressure signals were fed to a personal computer and digitized through a Data Translation® DT 2801-A A/D board. $Pa$ was continuously displayed to the subject on a Hewlett-Packard oscillograph.

Rib cage (RC) and abdomen (AB) displacements. RC and AB displacements were studied in two subjects (Nos 6 and 7). Magnetometry and inductance plethysmography being unreliable during magnetic stimulation, tightly fitting air-filled tubings were taped around the upper rib cage and abdomen and connected to two differential pressure transducers (±2 cmH₂O; Validyne, Northridge, USA). Their position was adjusted during diaphragmatic twitches produced by selective bilateral electrical stimulation of the phrenic nerves (see below). The AB tubing was set approximately at the level of the umbilicus, in such a way as to record a positive change in pressure during diaphragmatic twitches (i.e. AB expansion). The RC tubing was set as high as possible on the rib cage, in such a way as to record a negative change in pressure during diaphragmatic twitches (i.e. upper rib cage inward movement).

Stimulations

Bilateral electrical stimulation (BES) of the phrenic nerves was performed according to the usual technique, using bipolar electrodes connected to a constant current stimulator delivering square-wave shocks of 0.1 ms duration (Curamètre®; Bio-Industry, Outreau, France). The nerve once spotted, the intensity of BES was progressively increased until a plateau in the amplitude of the corresponding compound diaphragmatic action potentials (CDAP) was reached. It was then further increased by 10–20%, to ascertain supramaximality.

Magnetic stimulation was carried out using a first generation Magstim 200 stimulator (The Magstim Co., Sheffield, UK) equipped with a 90 mm circular coil (1.5 Tesla, 0.1 ms square-wave pulses). Cervical magnetic stimulation (CMS) was performed as reported previously [18]. As with BES, its supramaximal nature was judged according to the relationship between stimulation intensity and CDAP amplitude. Because a plateau in CDAP was not always observed, the peak-to-peak amplitudes
of the mean.

The CMS-related CDAPs (maximal intensity) were systematically compared with those of the supramaximal BES-related CDAPs (fig. 1).

Magnetic stimulation of the cortex (CxMS) was performed with the coil positioned over the vertex, 1 to 3 cm behind the mid-auricular plane [20], at maximal stimulation intensity. The electromyographic (EMG) response to CxMS is henceforth termed motor evoked potential (MEP).

The $P_{di}$ responses to BES, CMS, and CxMS will henceforth be termed $P_{di,p-ES}$, $P_{di,p-CMS}$ and $P_{di,c}$, respectively ("p" standing for peripheral, and "c" for central).

**Procedures**

**Training.** Before the experiments, all subjects were allowed time enough to learn how to generate and control $P_{di}$ using visual feedback. Maximal $P_{di}$ ($P_{di,max}$) was then determined [5].

**Stimulations.** To preclude twitch potentiation, the subjects were first asked to breathe quietly and remain relaxed for 20 min [21]. BES, CMS and CxMS were performed at functional residual capacity (FRC), after airway occlusion, with the diaphragm relaxed. CMS and CxMS were then superimposed upon graded isovolumetric diaphragmatic contractions ($P_{di,vol}$) ranging 10–100% of $P_{di,max}$.

Values in the results section are mean±standard error of the mean.

**Peripheral stimulation**

With increasing CMS intensity, a clear plateau was observed in all cases with respect to $P_{di,p-CMS}$, but only in four with respect to CDAPs. Figure 1, according to BLAND and ALTMAN [19], shows that the peak-to-peak amplitudes of the CDAPs produced by CMS at maximal intensity were not different from the peak-to-peak amplitudes of the CDAPs produced by supramaximal BES. Therefore, it is likely that supramaximality was consistently achieved with CMS in this series of experiments.

At FRC, with the diaphragm relaxed, mean±SEM $P_{di,p-CMS}$ was $27.5±2.2\text{ cmH}_2\text{O}$, whereas $P_{di,p-ES}$ was $23.3±1.9\text{ cmH}_2\text{O}$ (table 1). The difference was entirely accounted for by higher twitch $P_{tes}$ values.

Phrenic conduction time ranged 5.5–7 ms, and was similar with BES and CMS. In some subjects, the shape of the CMS-related CDAP was different from that of the BES-related CDAP.

In the two subjects in whom they were studied (Nos. 6 and 7), AB displacements induced by BES and CMS were similar in pattern and amplitude. BES-related RC displacements consisted in sharp paradoxical inward movements starting from the relaxed position. The RC pressure signal in response to CMS began with a slight increase, which was followed by a negative swing ascertaining an inward movement, but the latter was of lower amplitude and slower time course than with BES (fig. 2).

When CMS was superimposed upon a voluntary diaphragmatic contraction, the amplitudes of the corresponding twitches linearly decreased according to:

$$P_{di,p-CMS} = a - b P_{di,vol}$$

The $Y$-intercept "a" of Equation (1) amounted to $26.6±2.1\text{ cmH}_2\text{O}$ (range $19.7$–$33.4$), not significantly different from the actual $P_{di,p-CMS}$ at FRC (table 1; paired $t$-test). The slope "b" of Equation (1) was $0.24±0.03$.

Individual $P_{di,p-CMS} - P_{di,vol}$ relationships are depicted in figure 3a. In six subjects, $P_{di,p-CMS}$ could be brought to zero, suggesting maximal diaphragmatic activation [7, 8]. In subject No. 5, a positive $P_{di,p-CMS}$ persisted whatever the effort made (see Discussion).
Fig. 3. – a) Effects of a voluntary diaphragmatic contraction (P_{di,vol}) on the amplitude of the transdiaphragmatic pressure response to cervical magnetic stimulation (P_{di,p-CMS}), in the seven subjects. The data points correspond to the expression both of P_{di,vol} and P_{di,p-CMS} as percentage of their maximum values. All subjects but No. 5 were able to produce a P_{di,vol} extinguishing the response to peripheral phrenic nerve stimulation. b) Effects of a voluntary diaphragm contraction (P_{di,vol}) on the amplitude of the transdiaphragmatic pressure response to cortical magnetic stimulation (P_{di,c}), in the seven subjects. P_{di,vol} is expressed as percentage of its maximal value. P_{di,c} is expressed in percentage of P_{di,p-CMS} at relaxed FRC. In subjects Nos. 1–3, 6 and 7, P_{di,c} continuously increased with P_{di,vol} and could be fitted by a linear function. In subjects Nos. 4 and 5, P_{di,c} first increased with P_{di,vol} (facilitation) and then decreased (twitch occlusion) (see text for details).
observed, and a groups. No response of the relaxed diaphragm was for this response to appear. As a result, P_{di,vol} did not affect the amplitude and the latency of the CDAPs (fig. 4).

Cortical magnetic stimulation

CxMS activated several upper and lower limb muscle groups. No response of the relaxed diaphragm was observed, and a P_{di,vol} of about 10% P_{di,max} was needed for this response to appear. In subjects Nos. 1–3, 6 and 7, P_{di,c} continuously increased with P_{di,vol} (fig. 3b), according to:

\[ P_{di,c} = c + d P_{di,vol} \]  \hspace{1cm} (2)

As a result, P_{di,c} twitches persisted at P_{di,vol} levels corresponding to the suppression of P_{di,p-CMS}. In subjects Nos. 6 and 7, these twitches were associated with an outward AB displacement and a complex RC response, including an inward movement, making a diaphragmatic contribution likely (see Discussion).

It should be noted that because the absence of response of the relaxed diaphragm was probably due to the lack of power of the stimulator and not to a physiological mechanism [22], the corresponding data points have been discarded in order to compute the regression equations given in figure 3b. The slope of Equation (2) reached significance in the five subjects. Of note, the r values were lower for CxMS than for CMS, but were not improved by fitting nonlinear models (e.g. power function) to the data.

With increasing P_{di,vol}, the amplitude of the MEPs increased and multiple EMG responses could be observed, whereas latencies tended to decrease (fig. 4).

In subjects Nos. 4 and 5, P_{di,c} initially increased with P_{di,vol}, and then markedly decreased (fig. 3b).

Discussion

The main finding of this study is that CMS can provide diaphragmatic twitch occlusion data very similar to those obtained with BES. The study further suggests that, given that some points are elucidated, CxMS could be used in the assessment of the central drive to respiratory muscles. In addition, it provides information which may contribute to explain the differences in P_{di} observed with CMS and BES. This aspect will be examined first.

Contribution to the understanding of the differences between CMS and BES

CMS is a noninvasive and easy-to-use nonvolitional test of diaphragmatic contraction [18, 23]. It is associated with P_{di} values higher than those obtained with BES, due to a larger P_{oes} component ([23], and this study). Because CMS provokes diaphragmatic contraction via stimulation of cervical roots rather than of the phrenic nerve itself, it implies co-contraction of various muscles innervated by the C_{3}-C_{6} roots [24], as well as muscles innervated by the Xth cranial nerve. Therefore, it has been postulated that the better "efficiency" of CMS-induced diaphragmatic contraction in terms of P_{oes} was due to decreased rib cage distorsility as a result of the prior action of extradiaphragmatic muscles (e.g. sternomastoid or scaleni muscles) [23, 25].

There are other possibilities: 1) a direct contribution of CMS-related neck muscle contraction to P_{oes} (for such an explanation to hold, it has to be assumed that the P_{oes} produced by neck muscles contraction is not or is only partially transmitted to the abdominal cavity, and can therefore contribute to the build-up of P_{di}); or 2) partial recruitment of the diaphragm by BES due to incomplete aggregation of fibres constituting the phrenic nerve at the site of stimulation in the neck (this can be due to anatomical variability: on the one hand, the phrenic nerve in the neck is at times made of a complex of rootlets that unite to constitute a single nerve only in the thorax [26]; on the other hand an accessory phrenic nerve from the fifth cervical root sometimes adds to the main trunk...
in the mediastinum [27]. In both cases, CMS would activate all phrenic fibres, but not BES).

Our AB and RC displacement data in subjects Nos. 6 and 7 (fig. 2), showing a reduced and slower inward movement of RC with CMS, substantiate the rib cage stabilization hypothesis. In this respect, the initial positive pressure "bump" observed at the very beginning of RC movements as assessed by the devices used in this study is interesting to discuss. If it corresponded to RC expansion resulting from the action of neck muscles, there would be a direct contribution of the latter to $P_{oes}$, and the CMS induced $P_{oes}$ swing would probably be biphasic [18]. Such a pattern is not observed in practice. Rather, a small positive $P_{oes}$ deflection often precedes the CMS induced $P_{oes}$ negative swing (see, for example, fig. 2 in [23]). Rib cage stabilization with CMS could, therefore, be the result of initial expiratory action on the rib cage, via contraction of the trapezius or pectoralis muscles for example. The increase in volume of these muscles associated with their contraction would then explain the initial positive RC "bump".

**Effects of an underlying contraction on the diaphragm response to CMS**

$P_{di,p}$-CMS linearly decreased with increasing $P_{di,vol}$, whereas the latency and the amplitude of the corresponding CDAPs did not change. Such a pattern confirms that CMS produces diaphragmatic responses through stimulation of peripheral nervous structures (the roots) rather than of a central one (the spinal cord). In the latter case, increasing $P_{di,p}$-CMS and increasing amplitude of CDAPs would have been expected with $P_{di,vol}$, together with reduced CDAP latency, as a result of spinal facilitation [28].

As stated in the introduction, the twitch occlusion technique has interesting potentialities for studies of diaphragmatic function [17]. However, the consistency of the stimulus delivered to the nerves by BES is often difficult to maintain during intense inspiratory manoeuvres recruiting neck muscles. Being by nature much easier to maintain than BES in this setting, CMS could promote the clinical use of information derived from the twitch occlusion. We feel that the data provided by this study are a step in this direction. Indeed, the influence of $P_{di,vol}$ on $P_{di,p}$-CMS was comparable to what is described with BES: 1) the y-intercept of the $P_{di,vol}$ - $P_{di,p}$-CMS relationship (26.5±2.1 cmH$_2$O) was not significantly different from the $P_{di,p}$-CMS values obtained at relaxed FRC (27.5±2.2 cmH$_2$O); 2) the slope of the relationship, corresponding to the twitch-to-tetanus ratio, was 0.24±0.03, which is very close to values reported for the diaphragm and other mammalian muscles [29]; and 3) 6 out of 7 subjects were able to produce voluntary efforts suppressing the response to CMS. As we are confident that CMS was actually supramaximal (see Results), this suggests that these efforts were produced by maximal or close to maximal diaphragmatic command (see below).

In addition, we performed a BES twitch occlusion procedure in subject No. 5, who failed to extinguish $P_{di,p}$-CMS: the results were impossible to distinguish from those for CMS.

Given that CMS provides valid diaphragmatic twitch occlusion data, and taking into account the differences between $P_{di,p}$-ES and $P_{di,p}$-CMS discussed above, one could further submit that the "magnetic" twitch occlusion could in fact give a more realistic description of diaphragmatic properties than the "electric" one. The reasons for this contention are as follows. When $P_{di}$ manoeuvres are performed with visual feedback, neck muscles often stay relaxed during low intensity contractions, and are co-activated during high intensity ones. BELLEMARE and BIGLAND-RITCHIE [7] reported that neck muscle contractions became significant for $P_{di,vol}$ values of about 70% of $P_{di,max}$, and then increased with $P_{di,vol}$. In the four subjects where $E_{sm}$ was recorded (see Methods), we observed such a recruitment pattern. Neck muscle activation should markedly decrease upper rib cage distortion.

Thus, during a BES twitch occlusion procedure, the stimulation-induced diaphragmatic contraction will interact with a highly distortable rib cage at low $P_{di,vol}$ values, and with a much stiffer rib cage at high $P_{di,vol}$ values.

The corresponding improved efficiency of diaphragmatic contraction probably explains the linearity observed by BELLEMARE and BIGLAND-RITCHIE [7] in the $P_{di,p}$-$P_{di,vol}$ relationship. They noted that the slope of this relationship abruptly decreased after approximately 70% of $P_{di,max}$ (see fig. 7 in [7]), which led them to state that $P_{di,max}$ extrapolated from submaximal efforts using the twitch occlusion principle could slightly underestimate the actual value. The CMS-related rib cage stabilization discussed previously should result in a more homogeneous "environment" for diaphragmatic contraction at all levels of $P_{di,vol}$ and, therefore, reduce this problem. In most of our subjects (fig. 3a), we did not observe the slope change described by BELLEMARE and BIGLAND-RITCHIE [7], and the extrapolated values for $P_{di,max}$ closely matched the actual values observed during attempted maximal efforts (126.2±16 vs 122.2±19.2 cmH$_2$O, respectively; NS).

**Effects of an underlying contraction on the diaphragmatic response to cortical magnetic stimulation**

To our knowledge, all published studies dealing with diaphragmatic response to CxMS [20, 30–32] mention the need of a voluntary contraction for a diaphragmatic response to appear. This is probably simply a consequence of the insufficient power of first generation stimulators used in previous studies, as well as in the present one.

The increase in $P_{di,c}$ with $P_{di,vol}$ depicted in figure 3b, together with the shortening in latency and the increase in amplitude of MEPs depicted in figure 4, is consistent with the phenomenon of facilitation which has been well-described for other muscles [28]. However, $P_{di,c}$ continued to increase with $P_{di,vol}$ over the whole range of $P_{di,vol}$ studied in 5 out of 7 subjects (fig. 3b). This represents an important difference from limb muscles, where it is generally admitted that a voluntary contraction amounting to 10–20% of maximal strength is sufficient to maximize the effects of facilitation [33]. Only two subjects (Nos. 4 and 5) exhibited a somewhat expected pattern, namely increasing $P_{di,c}$ up to a certain level of $P_{di,vol}$ (facilitation) followed by a tendency for $P_{di,c}$
Firstly, mechanisms of diaphragmatic facilitation could differ from those of limb muscles facilitation. In fact, facilitation in limb muscles depends partly on the pre-activation of spinal motoneurones by information carried by Ia fibres [34]. Since the diaphragm contains few spindles, it is conceivable that motoneurone preactivation requires higher levels of tension than for limb muscles. Such a hypothesis seems to be supported by the continuous decrease of MEP latency with P\textsubscript{di,vol}, suggesting persistent increasing facilitation.

Secondly, the role of muscles other than the diaphragm could be invoked. In this respect, the progressive recruitment of inspiratory and expiratory muscles during P\textsubscript{di,vol} manoeuvres of increasing intensity would lead to their increasingly stronger contraction in response to CxMS, again because of the occurrence of facilitation. This contraction, under certain conditions, could contribute to P\textsubscript{di,c}. Gandevia et al. [9] have already reported the persistence of P\textsubscript{di,c} at P\textsubscript{di,vol} levels where the peripheral response to phrenic nerve stimulation was abolished. Using BES, they have shown that normal, motivated subjects can produce diaphragmatic contractions intense enough to extinguish the twitch, both during expulsive and inspiratory efforts. In spite of this complete peripheral twitch occlusion, electrical transcranial cortical stimulation evoked a large P\textsubscript{di} response. It was larger when the efforts were expulsive than inspiratory, but existed in both cases. As was the case in our study, this “extra” P\textsubscript{di} response at times exceeded the peripheral twitch at relaxed FRC, and, with expulsive manoeuvres, was predominantly due to a rise in gastric pressure.

To explain their data, Gandevia et al. [9] emphasized the fact that maximal activation of the diaphragm, as ascertained by the twitch occlusion, was not mandatorily associated with maximal activation of its synergistic in-series muscles (e.g. the inspiratory intercostals in the case of inspiratory manoeuvres, or the abdominal muscles in the case of expulsive manoeuvres). These muscles, therefore, could still be recruited by CxMS in spite of maximal diaphragmatic activation, and could contribute to P\textsubscript{di} if transmission of the corresponding \( P_{GA} \) to the thorax or of \( P_{OES} \) to the abdomen was incomplete. Such an incomplete pressure transmission through the diaphragm was demonstrated by direct stimulation of abdominal muscles: their contraction did not produce P\textsubscript{di} when the diaphragm was relaxed, but a P\textsubscript{di} swing appeared when it was contracted. If this hypothesis is correct CxMS could provide an estimate of the degree of global activation of synergistic respiratory muscles during voluntary respiratory manoeuvres.

However, it should be noted that the RC and AB displacement data collected in subjects Nos. 6 and 7 in the present study, although very fragmentary and preliminary, could cast doubt on the theory presented above and bring the diaphragm back into play. Indeed, P\textsubscript{di,c} responses at P\textsubscript{di,max} were not associated with an inward movement of the abdominal wall, as would have been expected with abdominal recruitment, but rather with an outward movement suggesting a diaphragmatic contraction. There was also some degree of RC inward movement.

Suggesting a mechanism by which the diaphragm, supposedly fully activated, could still respond to CxMS, is not simple. Because we did not use the most sensitive technique [35] to ascertain the reality of twitch occlusion in our subjects, one could argue that diaphragmatic activation at the highest values of P\textsubscript{di,vol} was not actually maximal. In this case, a positive P\textsubscript{di,c} would be expected, but its amplitude would be very small. Alternatively, BES could have combined excitatory and inhibitory effects [36], which would limit the corresponding P\textsubscript{di} output. The persistence of P\textsubscript{di,c} would then be explained by the lack of inhibitory effects with CxMS. This remains highly speculative. Confirmation of the role of the diaphragm in development of P\textsubscript{di,c} at P\textsubscript{di,max} on a larger number of subjects on the one hand, and other types of investigation, such as collision tests, on the other hand are obviously needed to clarify this point.

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References


