

Diaphragmatic Compound Muscle Action Potentials Recorded From Over the Sternum and Lateral Chest Wall: a Comparative Study

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Abstract

Objectives:

To report normative data for diaphragmatic compound muscle action potentials (DCMAPs) recorded from over the sternum and lateral chest wall (LCW).

Method:

The phrenic nerve of seventy-three healthy subjects was bilaterally stimulated at the posterior border of the sternocleidomastoid muscle. DCMAPs from over the sternum and LCW were recorded (inspiration/expiration).

Results:

Normative values of sternal and LCW DCMAPs were presented. Significant differences were found between sternal and LCW potentials in the mean latency, amplitude, and area ($p < 0.001$). The duration did not differ between the two sites. Differences were found between inspiration and expiration, right and left sides, and men and women.

Regression analysis showed a relation between latency of sternal and LCW potentials and age. Latency (LCW potentials) and amplitude and area (sternal/LCW potentials) were related to gender. Amplitude (LCW potentials/inspiration) and area (sternal potentials/inspiration) were related to chest circumference ($p = 0.023$ and 0.013 respectively). Area (sternal potentials/expiration) was related to the BMI ($p = 0.019$).

Conclusions:

Our normative values for sternal and LCW DCMAPs are provided. Notable differences in the DCMAPs parameters were detected between the two recording sites, inspiration and expiration, right and left, and men and women.

Significance:

The technique of phrenic nerve should be standardized.

Introduction

Phrenic nerve conduction study has found increasing application in the diagnosis of phrenic neuropathy associated with different medical and surgical conditions [1-12].

Percutaneous electric stimulation of the phrenic nerve at the neck with recording of the diaphragmatic compound muscle action potential (DCMAP) by surface electrodes is easy, quick, and non-invasive technique [2]. Amplitude, latency, area, and duration are measures used to evaluate phrenic nerve integrity [13-16].

Several studies have been conducted in healthy individuals to describe diaphragmatic potentials and normative values have been provided. The influence of age, gender, height, body mass index (BMI), chest circumference, and other subjects' characteristics were shown [13-16]. Values during inspiration and expiration were also provided [13-15].

The phrenic nerve can be stimulated at the supraclavicular fossa [13-15, 17, 18] or at the posterior border of the sternocleidomastoid muscle at the level of the cricoid or thyroid cartilage [19-21]. DCMAPs are commonly recorded from over the sternum with the active electrode (G1) fixed 5 cm above the xiphoid process and the reference

electrode (G2) placed 16 cm from G1 on the chest margin ipsilateral to the stimulated phrenic nerve [13-16] or from over the lateral chest wall (LCW) with recording electrodes placed in the 6th, 7th, or 8th intercostal space along the anterior axillary line [11, 19, 21-24]. Diaphragmatic potentials are less commonly recorded from the 7th, 8th, or 9th intercostal space on the anterior or anterolateral chest [25, 26].

A number of studies demonstrated that different positions of the recording electrodes contribute to variability of phrenic nerve conduction study parameters. Swenson and Rubenstein [17] found significant differences in the amplitude of potentials recorded from over the xiphoid and lateral chest wall. Dionne et al [18] recorded diaphragmatic potentials from over six different recording sites and reported similar finding. McKenzie and Gandevia [20] found variation in the latency of DCMAP recorded at different sites. They concluded that recording electrodes at each site are relatively selective for the adjacent portion of the diaphragm [20] that is innervated by separate nerve branches [27].

The results from these studies are significant when evaluating segmental lesions of diaphragmatic innervation resulting from different medical or surgical conditions [20]. These studies were however conducted on a limited number of healthy individuals (20, 11, and 3 subjects) and did not define reference values for DCMAP parameters at different sites [17, 18, 20].

Aim of the work:

The aim of this study was to report normative data for DCMAP parameters recorded from over the sternum (sternal DCMAP) and lateral chest wall (LCW DCMAP) in healthy individuals and highlight factors that may contribute to variations in the measurements at the two sites.

Subjects And Methods

Seventy-three healthy individuals (48 men and 25 women) participated in the study. Their age ranged from 18 to 61 years (mean: 37.66 ± 11.51) with almost equal number of subjects in each decade. Subjects were recruited from hospital personnel (students, employees, and relatives of patients). None of the subjects was current or ex-smoker. Exclusion criteria included pregnancy, chest wall abnormalities, and the presence of any medical condition. None of the participants was involved in any sport activity.

Their mean chest circumference (measured at the fourth intercostal space/auxiliary level) [28] was 93.96 ± 7.77 cm, height was 1.69 ± 0.08 m, and weight was 72.84 ± 12.95 kg. The mean BMI was 25.73 ± 5.08 kg/m². Two subjects (2.9%) were underweight, thirty-two (45.7%) had normal weight, twenty-four (34.3%) were overweight, and twelve (17.1%) were obese.

The study was approved by the local ethics committee of the Faculty of Medicine, Alexandria University and an informed written consent was obtained from all participants.

Phrenic nerve conduction study:

Electrophysiological studies were conducted on Nicolet VikingQuest version 11 USA electromyography machine. During electrophysiological examination, the skin was kept warm around 32-34°C. A bipolar surface stimulator was used to provide stimulation. Self-adhesive surface electrodes were used for recording. The ground electrode was a disc electrode placed between the stimulating and the recording electrodes.

Phrenic nerve conduction study was performed bilaterally. Subjects were lying supine with the head slightly elevated and rotated to the opposite side to the nerve under stimulation. The phrenic nerve was supramaximally stimulated at the posterior border of the sternocleidomastoid muscle at the level of the cricoid cartilage with the anode placed proximal to the cathode. Rectangular pulses of 0.2–0.5ms duration were used. Measurements were made separately during normal inspiration and expiration. Co-stimulation of the brachial plexus was observed as indicated by muscle contractions, arm movement, paraesthesia (reported by the subject), a short latency <5 ms, low amplitude and initially positive response. In such case, the cathode was repositioned medially to preferentially stimulate the phrenic nerve.

The DCMAP was recorded from over the sternum with the active electrode placed 5 cm superior to the tip of the xiphoid process and the reference electrode placed 16 cm from the xiphoid process at costal margin ipsilateral to the stimulation site (sternal recording). DCMAP was also recorded from over the lateral chest with the active electrode placed over the 7th intercostal space at the anterior axillary line and the reference electrode at the 8th intercostal space ipsilateral to the stimulation site (LCW recording). Two responses were recorded at each site and the average values of the two responses were calculated.

The filter band width was set at 2 Hz to 10 kHz, sweep speed at 5 ms/division, and sensitivity at 200-2000 μV /division. Measured parameters included the onset latency, amplitude (peak to peak), duration (negative phase), and area. Side-to-side differences were calculated for all parameters. Side-to-side amplitude difference was expressed as a ratio between the two sides.

Peripheral neuropathy was excluded electro-physiologically [29].

Statistical analysis:

Data were analyzed using IBM SPSS software package version 20.0. (Armonk, NY: IBM Corp) The Kolmogorov-Smirnov test was used to verify the normality of distribution and all parameters were normally distributed. Quantitative data were described using range, mean, and standard deviation. Significance was judged at the 5% level. Cut-off values were calculated as $\text{mean} \pm 2\text{SD}$. **Paired t-test** was used to compare between two periods. **Student t-test was used** to compare between two studied groups. **Pearson coefficient was used** to correlate between two normally distributed quantitative variables. Multivariate regression analysis was used to estimate the relationship between different dependent and independent variables.

Results

The phrenic nerve could not be stimulated in 3 subjects (short/obese neck). Data presented are pooled data (140 nerves from right and left sides in 70 subjects). All parameters were normally distributed, so transformation of data was not required. Mean, normative, and inter-side difference values of sternal and LCW potentials during inspiration and expiration are presented in table 1.

The latency of sternal potentials was significantly shorter than LCW potentials with significantly higher amplitude and larger area. The duration did not differ significantly between the two recording sites, table 2. DCMAP amplitude was significantly higher during inspiration. The mean latency, duration, and area were significantly higher during expiration, table 2. The latency and duration were significantly longer on the right side (except for latency of sternal potentials during inspiration) whereas the amplitude was significantly higher on the left side (except for sternal potentials during expiration). The area did not differ between the two sides, table 3. Parameters from right and left

sides were strongly/very strongly correlated, table 3 The mean latency, amplitude, and area were significantly higher in men. The duration did not differ significantly between the two sexes, table 4

Table 5 shows the results of correlation analysis between DCMAP parameters and subjects' characteristics. The latency of diaphragmatic potentials correlated positively with age (inspiration/expiration), height (inspiration/expiration), and chest circumference (sternal potentials during inspiration). The amplitude of sternal and LCW potentials (during inspiration/expiration) correlated negatively with the BMI and that of LCW potentials (inspiration) correlated negatively with chest circumference.

The duration of the sternal potentials correlated positively with height (inspiration) and negatively with BMI (inspiration/expiration) and chest circumference (expiration). The duration of the LCW potentials correlated positively with height (inspiration/expiration).

The area of sternal and LCW potentials correlated negatively with the BMI (inspiration/expiration) and that of sternal potentials correlated positively with height (inspiration/expiration).

Multiple regression analyses were done between different parameters of DCMAP (dependent variables) and subjects' criteria (independent variables), table 6.

Latency of sternal and LCW potentials (inspiration/expiration) was related to age ($p < 0.001$). Latency of LCW potentials (inspiration/expiration) and amplitude and area of sternal and LCW potentials (inspiration/expiration) were related to gender. Amplitude of LCW potentials and area of sternal potentials (inspiration) were related to chest circumference ($p = 0.023$ and 0.013 respectively). Area of sternal responses (expiration) was related to the BMI ($p = 0.019$). Duration of LCW responses (inspiration/expiration) was related to height ($p = 0.003$ and 0.002 respectively).

An example of DCMCPs recorded from over the sternum and lateral chest wall in one of the studied subjects is shown in figure 1.

Discussion

None of the participants was current- or ex-smoker to avoid the impact of smoking on the lungs (chronic obstructive pulmonary disease or emphysema). None of the subjects had peripheral neuropathy.

Overall, our mean and normative values of DCMAPs parameters were comparable to those reported in the literature. Differences are mostly related to variations among studies in the site of stimulation or signal recording or how the parameter was measured.

The mean latency values of sternal potentials [7.08 ms (inspiration) and 7.14 ms (expiration)] are higher than values reported by Chen et al (6.5 ms) [13], Maranhão et al [6.12 (inspiration) and 6.42 ms (expiration)] [15], Resman-Gäspěrsč and Podnar [6.55 (inspiration) and 6.59 ms (expiration)] [14], Vincent et al (6.59 ms) [16], and Dionne et al (6.6 ms) [18], They all stimulated the nerve at a lower level (supraclavicular fossa).

The mean latency values of LCW potentials [7.29 ms (inspiration) and 7.34 ms (expiration)] are close to values reported by Luo et al (7.3 (left side) and 7.6 ms (right side) [23], Dionne et al (7.14 ms) [18], and Mckenzie and Gandevia [7.68 (right side) and 7.92 ms (left side)] [20]. They adopted the same stimulation and recording montages as the current work.

Our upper limits of the latency [8.34 (sternal/inspiration), 8.48 (sternal/expiration), 8.63 (LCW/inspiration), 8.64 ms (LCW/expiration)] are comparable to values by Vincent et al [(8.36 ms (sternal))] [16] and Resman- Găspărsă and Podnar [7.92 ms (sternal/inspiration and expiration)] [14]. They are higher than values reported by Maranhão et al (6.34 ms (sternal/inspiration) and 6.42 (sternal/expiration) [15]. Chen et al found that the suggested normal latency limit is <8 ms (sternal) [13]. Variation in limit values among studies is again related to variation in stimulation and recording sites.

Our mean amplitude values are higher than values reported in the literature [13-18, 20]. This is because we measured the amplitude from peak-to-peak as opposed to other studies where the amplitude was measured from onset-to-peak. Highest mean amplitude values were recorded from over the sternum (1.4 (inspiration) and 1.16 mV (expiration). Lower limits of the amplitude [0.60 (sternal/inspiration), 0.46 (sternal/expiration), 0.34 (LCW/inspiration), 0.32 mV (LCW/expiration)] are comparable to values by other studies (range: >0.3-0.7 mV) [13-15]. Swenson and Rubenstein [17], Johnson et al [30], and Vincent et al [16] on the other hand reported much lower normative values [0.1, 0.12, and (0.14 right and 0.11 left) mV respectively). As illustrated by Maranhão et al [15], the wide range of phrenic nerve amplitude creates a great problem in determining a lower normal limit.

Mean and normative values of the duration of sternal DCMAP in the current work [14.54 and 20.82 ms (inspiration) and 17.74 and 24.12 ms (expiration)] are within the range of values reported by Chen et al [13], Resman- Găspărsă and Podnar [14], Maranhão et al [15], and Vincent et al. [16] They all recorded the potentials from over the sternum. No data is available in the literature on the normative value of the duration of potentials recorded from over the LCW.

Mean values of the area of sternal DCMAP [7.27 mV.ms (inspiration) and 8.12 mV.ms (expiration)] are close to values in the studies by Chen et al [13], Resman-Găspărsă and Podnar [14], and Maranhão et al. [15] Our normative values [2.47 mV.ms (inspiration) and 3.32 mV.ms (expiration)] are however lower than other studies (lower limit value > 4 mV.ms) [13-15]. This variation is mostly related to how the limit normal was determined (mean-2SD as in the current work compared to 5th percentile limit in other studies). Our values are higher than that by Vincent et al [16] (mean= 3.05 mV.ms and limit normal=0.87 mV.ms). Lower amplitude values have contributed to lower area values in their study.

The means values of the area of DCMAP from over LCW ([5.59 mV.ms (inspiration) and 6.19 mV.ms (expiration)] are comparable to value by Dionne et al (6.41 mV.ms) [18]. Data on the normative values from costal diaphragm is scanty in the literature.

Our mean and normative values of the interside differences in latency, amplitude, duration, and area were presented. These values are useful in assessment of patients with a unilateral phrenic nerve lesion as those caused by trauma or surgery especially when the absolute values of both sides are within the normal range. Our values are consistent with those in the studies by Chen et al [13] and Resman-Găspărsă and Podnar. [14] Vincent et al [16] reported higher interside differences in the latency, amplitude, and duration of potentials recorded from over the sternum. Swenson and Rubenstein [17] also reported higher mean interside amplitude differences of potentials recorded from over the sternum and LCW which may be related to differences in the placement of recording electrodes.

Differences between sternal and LCW DCMAPs:

Comparative studies showed that the latency, amplitude, and area varied significantly between sternal and LCW potentials. The duration did not differ between the two recording sites.

The latency of sternal potentials was significantly shorter than LCW potentials. Swenson and Rubenstein [17] reported similar finding. This can be expected considering the shorter length of the anterior (sternal) branch. McKenzie and Gandevia [20] in their illustrative work provided the distance from the branching points of phrenic nerve to different recording sites, which should correspond to the length of different branches. Distance corresponding to the anterior branch was shorter than that for lateral branch on the right (6.7 and 13.8 cm) and left sides (2.8 and 8.8 cm respectively).

The amplitude of sternal potentials was significantly higher than LCW potentials. The finding is consistent with that by Dionne et al [18] who recorded diaphragmatic potentials from over six different sites. The highest amplitude was obtained from over the sternum. They related this finding to the orientation of the electrical dipole of the diaphragm (different from limb muscles). They assumed that G1 and G2 are both active, with the DCMAP representing out-of-phase summation of opposite polarity activity at the two electrodes, and this in part has accounted for larger amplitude at the sternum as it was the only technique with G1 positioned above the xiphoid.

Amplitude difference between sternal and LCW potentials can also be due to variation between the two sites in the distance between the active and reference recording electrodes [31] with higher amplitude of sternal potentials being related to longer distance between recording electrodes compared to LCW recording site (16 and 3.5-5 cm respectively).

Recording of DCMAP from over the sternum was recommended by Chen et al [31] and Dionne et al [18] as this electrode position gives the maximum amplitude. It also allows the easiest and most rapid study to perform as it does not involve rib counting or multiple electrodes repositioning especially in obese subjects and in patients in intensive care unit where chest tubes and catheters are frequently encountered.

Contrary to our finding, Swenson and Rubenstein [17] found smaller but easily recorded potentials at the sternum. Maximum peak was recorded over the anterolateral chest at the intersection of the axillary line with transverse plane through the xiphoid. In their study both xiphoid and costal recordings had their reference electrodes placed below the umbilicus which may have accounted for their results.

The duration did not differ significantly between sternal and LCW potentials. This parameter reflects the range of conduction velocities of conducting nerve fibers [32] and is not expected to differ between DCMAPs recorded from over the two sites. Significantly larger mean area of sternal potentials is consistent with significantly higher amplitude values of sternal compared to LCW potentials with no difference in the duration between them.

Despite the given advantages of recording from over the sternum (higher amplitude and easier study with no need for rib counting or repositioning of the electrodes), [13, 18] it cannot be concluded that this montage should be the standard one for recording DCMAPs.

The diaphragm is a specialized muscle that demonstrates distinct muscular subvolumes (neuromuscular compartments) in which the intramuscular phrenic nerve distribution (branching) is confined [27]. Recording electrodes at each recording site are relatively selective for the subjacent portion and record activity from the underlying portion of the diaphragm [20] that is innervated by separate nerve branches [27].

The presence of significant differences between potentials recorded from over the sternum and lateral chest wall in the current work and in other studies [17, 18, 20] as well as the evidence by some electrophysiological studies that one branch but not the other can be affected in different medical conditions [33, 34] highlights the importance of assessing the conduction along different nerve branches in every case referred for electrophysiological evaluation. It also highlights the need to provide normative data for potentials recorded from different diaphragm subvolumes innervated by different branches of the nerve.

Differences between DCMAPs recorded during inspiration and expiration:

We found variation in DCMAP parameters with respiratory cycle. The mean latency, duration and area were significantly lower, and the mean amplitude was significantly higher during inspiration compared to expiration. Findings are mostly related to changing lung volumes during movement of the diaphragm in the respiratory cycle and changing physiological properties of the diaphragm during contraction [13, 35].

Similar to our finding, Maranhão et al [15] found shorter latency of DCMAPs during inspiration. Resman-Găspěrsč and Podnar [14] found significant difference in peak but not the onset latency between inspiration and expiration. DCMAPs were recorded from over the sternum in both studies. Latency of CMAP represents the summated durations of impulse propagation along the nerve fiber, time delay across the neuromuscular junction, and depolarization time across the muscle [29]. Increased muscle fibers conduction velocity during contraction (inspiration) therefore decreases the overall time needed for impulse propagation and hence decreases the response latency.

Significantly higher amplitude, shorter duration and smaller area during inspiration come in agreement with the results of different studies. Resman-Găspěrsč and Pondar [14] found similar changes in the amplitude and duration during inspiration. Changes in the area were however insignificant. Chen et al [13] reported increase in the amplitude and decrease in the duration with inspiration. Maranhão et al [15] demonstrated the same results although they did not conduct a statistical comparison. On the other hand, Swenson and Rubenstein [17] reported higher DCMAP amplitude during expiration. They recorded the potentials at the level of the 7th intercostal space with the reference electrode below the umbilicus.

The amplitude value of diaphragmatic potentials depends on the orientation of recording electrodes and the distance between recording electrodes and the diaphragm which change during the respiratory cycle [13]. Chen et al suggested that during inspiration the diaphragm flattens which changes the angle, the moving dipole meets at the recording electrodes [13] which according to the theory of volume conduction would result in increased CMAP amplitude [36]. Resman-Găspěrsč and Pondar added that shortening (shorter path) and thickening (faster conduction) of diaphragmatic muscle during contraction make larger amplitude and shorter conduction time and shorter duration [14].

In a similar context, it was proved by musculoskeletal ultrasound that thickness of the diaphragm in healthy individual increases by 28-96% during inspiration relative to expiration [37, 38]. This in turn increases conduction velocity of muscle fibers due to diminished resistance of thick fibers and improves synchronization of the electrical potentials. In addition, muscle fibers shortening bring them closer to the recording electrodes, thus muscle contraction increases temporal and spatial summation of muscle fibers action potentials leading to increased CMAP amplitude and decreased duration [39].

We found significantly smaller area during inspiration due to significant decrease in the duration that overweighed the increase in amplitude during inspiration. Maranhão et al [15] demonstrated decrease in the area of DCMAPs during inspiration. They did not however conduct a statistical comparison. Resman-Găspěrsč and Pondar [14] found that difference in the area between inspiration and expiration is not significant, still smaller during inspiration. They considered the area to be more useful than the amplitude due in part to its insensitivity to respiratory cycle. The current study was conducted on larger number of healthy individuals (70 subjects) compared to that by Resman-Găspěrsč and Pondar [15] (29 healthy subjects). It is evident that there is an assent among studies that the area of DCMAPs is smaller during inspiration. The variation is about the extent of the difference whether it is significant or not. The exact change of the area with respiration is to be further evaluated.

Differences between DCMAPs recorded on right and left sides:

We found significantly longer latency of diaphragmatic potentials on the right side. Mier et al [2] reported similar finding but did not mention an explanation. Delhez et al [40] and Katayama et al [41] on the other hand showed longer latency on the left side. They attributed their finding to longer anatomical course of the left phrenic nerve. Maranhão et al [15], Mckenzie and Gandevia [20], Vincent et al ⁽¹⁶⁾ and Chen et al [13] did not find significant differences between both sides.

The course as well as the length of the right and left phrenic nerves varies. Jiang et al measured the full length and the length of thoracic part of the phrenic nerve on both sides and both were shorter on the right side. The intramuscular branches were however longer on the right side in higher number of their studied corpses [42]. McKenzie and Gandevia measured the conduction distance from entry point of phrenic nerve branches into the diaphragm to motor points adjacent to different recording sites, which should correspond to length of intramuscular branches. All distances were longer on the right side [20]. This can be expected given that the right phrenic nerve has almost as straight course. It enters the diaphragm close to the esophageal hiatus [27, 42], thus at a point at longer distances from the anterior and lateral chest walls than the left phrenic nerve. The conduction along these branches (thin) is slower than conduction along the nerve trunk [29]. Both factors (the length and conduction velocity along the nerve trunk and the branches) may have accounted for the longer latency of the recorded potentials on the right side.

The amplitude of DCMAPs (except for sternal potentials during expiration) was significantly lower on the right side. Resman-Găspěrsč and Podnar similarly found lower amplitude of sternal potentials on the right side during inspiration but not expiration and attributed the finding to the higher position of right hemidiaphragm, which is lifted by the liver (longer and thinner muscle fibers) [14]. Absence of a difference between right and left sternal potentials during expiration can be expected because at the xiphoid and during expiration, anatomical and physiological differences between the two sides are kept to minimum.

The duration of diaphragmatic potentials was significantly longer on the right side. Given that electrical potentials can propagate easily through thickened muscle fibers with lower resistance, (Kim BJ) it can be speculated that the duration of DCMAPs from the right hemidiaphragm (thinner fibers) is longer than the left. The area did not differ between the right and left sides. Similar findings were found by Resman-Găspěrsč and Podnar [14]. This is quite expected as lower amplitude and higher duration on the right side are equated by higher amplitude and lower duration on the left side.

Despite the difference in DCMAP parameters between the right and left sides, we found strong/highly strong right-to-left correlation in all parameters ($P < 0.001$) indicating that if one side deviated from the mean value the other side tended also to deviate in the same direction. Mier et al also found significant correlation between right and left phrenic nerve conduction time ($r = 0.81$, $p < 0.001$) [2]. Swenson and Rubenstein found constancy of only the onset latency. They did not find right-to-left amplitude correlation [17]. Similar to their findings, Maranhão et al did not find consistent right-to-left correlation [15]. The presence of strong/highly strong right-to-left correlation indicates that in unilateral lesion, the opposite side would serve as a reasonable standard for comparison.

Differences between DCMAPs recorded in men and women:

Measured parameters were different between men and women. The mean latency, amplitude, and area of DCMAPs were significantly increased in men. The duration did not differ significantly between both sexes. Maranhão et al reported substantial differences in amplitude, latency, and duration of DCMAPs between men and women [15]. Resman-Găspărsă and Pondar [14] and Vincent et al [16] found significantly higher amplitude in men. Vincent et al [16] found that the latency was different after adjustment for age and BMI.

Differences between men and women in parameters of DCMAPs are mostly related to variations in the anthropometric measures between both sexes (longer nerves contributing to longer latency and greater muscle mass contributing to higher amplitude in men) [14, 16]. Larger area in men is due to significantly higher amplitude and longer duration (not to significant level) in men compared to women. Only the duration of the DCMAPs did not differ significantly between men and women. This finding is expected given that the duration reflects range of conduction velocities of nerve fibers [32] which should not differ by gender.

The correlation and regression analyses between different parameters of phrenic nerve conduction study and subjects' demographic and anthropometric data

Correlation and regression analyses revealed that latency of DCMAPs is significantly related to age which is consistent with data from several studies [13-16] and can be attributed to dropout of largest fibers, segmental degeneration, and reduced internodal length with aging [43, 44]. No correlation was found between age and other parameters. The same was reported by Maranhão et al [15] and Vincent et al [16] and can be explained by the exclusion of elderly individuals from enrollment in the study (oldest individual was 61 years). Prominent electrophysiological changes in amplitude, duration, and area are seen in individuals above the age of 60-65 years [45].

Regression analysis showed significant relation between amplitude and area of DCMAPs and gender. The finding is consistent with the presence of significant differences between men and women in the amplitude and area of DCMAPs and can be explained by larger muscle mass in men. The latency of LCW (but not sternal) potentials was also related to gender. The contribution is indirect and is in part related to longer nerves in men (more evident for the lateral branch).

The latency of DCMAPs was also found to correlate with height which can be explained by increased length of nerves with relative conduction slowing in tall individuals [29]. Our results agree with Resman-Găspărsă and Podnar [14] and Maranhão et al. [15] Results of the regression analysis did not however show a relation between latency of

DCMAPs and height. This indicates that height is not an independent contributor to the latency of DCMAPs. McKenzie and Gandevia found that conduction distance does not directly relate to the subject's height and conduction velocity is not uniform along the main nerve trunk [20]. In one study, the length of the sternum was used as a surrogate for the height. Nevertheless, it was not found to correlate with the latency of DCMAPs [14]. It is evident that the length of the phrenic nerve cannot only be represented by subjects' height as the intramuscular branches which vary in thickness and size also contribute to the conduction distance and latency.

The duration of DCMAPs correlated with height. This is because relative conduction slowing in tall individuals decreases the synchronization of muscle fibers potentials which increases the duration of CMAP [46]. This is further supported by the results of the regression analysis where duration of only the LCW recordings was found to be related to height. Lateral intramuscular branch is longer than sternal (anterior) branch [20] which should magnify the effect of increased length on the duration of the potentials.

The amplitude and area of sternal and LCW potentials (during inspiration and expiration) correlated negatively with the BMI. This is most likely due to amplitude attenuation by the thicker subcutaneous tissue in the person with higher BMI [47, 48]. Such a relation was not shown in the results of regression analysis which indicate that BMI is not an independent contributor to amplitude and area of the DCMAPs yet stills a factor. Factors as the mass of the diaphragm (not measured in the current work) may have major contribution to the amplitude and area than the BMI.

Absence of a correlation between the latency of DCMAPs and BMI is plausible since the latency represents the conduction along the fastest fibers regardless the number of axons [29]. Thus in clinical setting, the fastest fibers appear to conduct equally in thin and obese individuals.

Chest circumference did not correlate with most of the recorded parameters. It only correlated positively with the latency of sternal potentials during inspiration and expiration and negatively with amplitude (of LCW DCMAP) and area (of sternal DCMAP) during inspiration and duration of sternal potentials during expiration. Multiple regression analysis revealed that BMI is related to the amplitude of LCW responses and the area of sternal responses during inspiration ($p=0.023$ and 0.013 respectively).

Our results are contradictory to that by Resman-Gäspěřšć and Podnar [14] who found significant positive correlation between chest circumference and the CMAP area and Chen et al [13] who found significant positive correlation between chest circumference and the amplitude of diaphragmatic potentials and attributed this to the greater diaphragmatic muscle mass and more flattened diaphragms in persons with larger chest circumference. These contradictory findings may be because we measured the chest circumference at the level of the 4th intercostal space while Chen et al [13] measured the circumference at the level of the xiphoid process. Also, the intervening breast tissue in women may have accounted for differences in the measures of chest circumference between the two levels. The rib cage cross-section area may be a better measure to correlate with DCMAPs.

Technical aspects:

Participants were tested in the supine position which is characterized by larger diaphragmatic excursion than the upright position. In establishing reference values for labs or comparing patients or in follow up studies, the same decubitus must be adopted [49]. The current required for supramaximal stimulation was less than 50 mA in the majority of subjects with pulse duration of 0.2 ms.

We stimulated the nerve at the posterior border of the sternocleidomastoid muscle at the level of cricoid cartilage. This stimulation site is easy to locate, less painful and it is easy to avoid brachial plexus co-stimulation by moving the electrode more medial [2, 23, 50]. It is to be mentioned that, the stimulation site can vary from subject to subject and may even vary between the left and the right sides of one subject due to asymmetry [14].

The phrenic nerve was generally easy to stimulate. Difficulties, especially on the left side, were sometimes encountered possibly due to anatomical variation of the phrenic nerve that may pass through the anterior scalene muscle or there may be medial or lateral displacement of the phrenic nerve [51]. This difficulty was overcome by increasing current intensity and/or medial repositioning of the stimulating electrode.

The technique was also challenging, and the nerve could not be stimulated in 3 subjects with short or obese neck (despite more rotation and/or extension of the neck was tried). In clinical practice, if percutaneous electric stimulation could not be performed due to technical problems, needle electric stimulation or magnetic stimulation of the phrenic nerve is to be tried [23, 25].

Stimulation was performed during quiet respiration (at the end of inspiration and expiration). Generally, the phases of respiration may create a technical problem during performing phrenic nerve conduction study. However, if supramaximal stimulation to the phrenic nerve is repeated several times during quiet respiration and the two highest amplitude potentials are obtained, the phases of respiration are not an influential factor [52].

Conclusions And Recommendations:

The results of the current work showed that there are notable differences in the parameters of DCMAP between the sternal and LCW sites, inspiration and expiration, right and left sides, and men and women.

Normative values of diaphragmatic potentials along different branches should be provided. Moreover, the effect of age, gender, height, and BMI should be considered. The technique should be standardized regarding phases of respiration and the identification of the parameters to be measured. This is essential for valid comparison between patients and healthy subjects or during follow up and for comparisons between different labs and different studies.

Declarations

- The authors did not receive support from any organization for the submitted work.
- The authors have no competing interests to declare that are relevant to the content of this article.
- Ethics approval: The study was approved by the local ethics committee of the Faculty of Medicine, Alexandria University and the procedures used in this study adhere to the tenets of the Declaration of Helsinki.
- Consent to participate: Informed consent was obtained from all individual participants included in the study.

Author contributions

All authors contributed to the study conception and design. Material preparation, data collection and analysis were performed by D.M. and R.E. The first draft of the manuscript was written by G.Y. and N.E. and all authors commented on previous versions of the manuscript. All authors read and approved the final manuscript.

References

1. Shaw RK, Glenn WWL, Hogan JF, Phelps ML (1980) Electrophysiological evaluation of phrenic nerve function in candidates for diaphragm pacing. *J Neurosurg Sep*;53(3):345–54. doi: 10.3171/jns.1980.53.3.0345.
2. Mier A, Brophy C, Moxham J, Green M. (1987) Phrenic nerve stimulation in normal subjects and in patients with diaphragmatic weakness. *Thorax* 42:885–8.
3. DeVita MA, Robinson LR, Rehder J, Hattler B, Cohen C. (1993) Incidence and natural history of phrenic neuropathy occurring during open heart surgery. *Chest* 103(3):850–6
4. Bolton CF (1994) Assessment of Respiratory Function In the Intensive Care Unit. *Can. J. Neurol. Sci.* 21: S28-S34.
5. Macia F, Le Masson G, Rouanet-Larriviere M, Habonimana D, Ferrer X, Marthan R, Lagueny A (2003) A prospective evaluation of phrenic nerve conduction in multifocal motor neuropathy and chronic inflammatory demyelinating polyneuropathy. *Muscle Nerve* 28: 319–323.
6. Martínez SMF, Tirado EA, Corona MÁC, Bialik PS (2012) Estimulación eléctrica del nervio frénico en pacientes con ventilación mecánica prolongada. *An Med (Mex)* 57 (1): 32–38.
7. Lakhani M, Saiful F, Parikh V, Goyal N, Bekheit S, Kowalski M (2014) Recordings of diaphragmatic electromyograms during cryoballoon ablation for atrial fibrillation accurately predict phrenic nerve injury. *Heart Rhythm Mar*;11(3):369–74.
8. Maranhão AA, Rodrigues MMJ, Carvalho SR, Caetano MR, Compagnoni IM, Carnio TC, Ribeiro D (2018) Is phrenic nerve conduction affected in patients with difficult-to-treat asthma? *Arq Neuropsiquiatr* 76(3):177–182
9. Sen BK, Pandit A (2018) Phrenic Nerve Conduction Study in the Early Stage of Guillain–Barre Syndrome as a Predictor of Respiratory Failure. *Annals of Indian Academy of Neurology* 21(1):57–61
10. Mamede de Carvalho, Susana Pinto, Michael Swash (2018) Diaphragm motor responses to phrenic nerve stimulation in ALS: Surface and needle recordings. *Clin Neurophysiol Feb*;129(2):349–353
11. Reynaud V, Prigent H, Mulliez A, Durand M, Lofaso F (2021) Phrenic nerve conduction study to diagnose unilateral diaphragmatic paralysis. *Muscle & Nerve* 63:327–335.
12. Pinto S, De Carvalho M (2021) Phrenic nerve study as outcome in clinical trials for amyotrophic lateral sclerosis. *Amyotrophic Lateral Sclerosis and Frontotemporal Degeneration* 22: 9–13
13. Chen R, Collins S, Remtulla H, Parkes A, Bolton, CF (1995) Phrenic nerve conduction study in normal subjects. *MUSCLE NERVE* 18:330–5
14. Resman-Găspăresc A, Podnar S (2008) Phrenic nerve conduction studies: technical aspects and normative data. *Muscle Nerve* 37: 36–41
15. Maranhão AA, Carvalho SRS, Caetano MR, Alamy AH, Peixoto EM, Filgueiras PDP (2017) Phrenic nerve conduction studies: normative data and technical aspects. *Arq Neuropsiquiatr* 75(12):869–874,
16. Vincent M, Court-Fortune I, Costes F, Antoine J, Camdessanché J (2019) Phrenic nerve conduction in healthy subjects. *Muscle Nerve* 59:451–456
17. Swenson MR, Rubenstein RS (1992) Phrenic nerve conduction studies. *MUSCLE & NERVE* 15:597–603
18. Dionne A, Parkes A, Engler B, Watson BV, Nicolle MW (2009) Determination of the best electrode position for recording of the diaphragm compound muscle action potential. *Muscle Nerve* 40: 37–41
19. Davis JN (1967) Phrenic nerve conduction in man. *J Neurol Neurosurg Psychiatr* 30: 420–426.

20. McKenzie DK, Gandevia SC (1985) Phrenic nerve conduction times and twitch pressures of the human diaphragm. *J. Appl. Physiol.* 58(5):1496–1504
21. Similowski T, Mehiri S, Duguet A, Attali V, Straus C, Derenne JP (1997) Comparison of magnetic and electrical phrenic nerve stimulation in assessment of phrenic nerve conduction time. *Journal of Applied Physiology* 82(4):1190–9
22. Luo Y, Polkey M, R Lyall R, Moxham J (1999) Effect of brachial plexus co-activation on phrenic nerve conduction time. *Thorax* 54:765–770.
23. Luo YM, Polkey MI, Johnson LC, Lyall RA, Harris ML, Green M, Moxham J (1998) Diaphragm EMG measured by cervical magnetic and electrical phrenic nerve stimulation. *J Appl Physiol* 85(6): 2089–99)
24. Ito H, Ito H, Fujita K, Kinoshita Y, Takanashi Y, Kusaka H (2007) Phrenic Nerve Conduction Study in the Early Stage of Guillain–Barre Syndrome as a Predictor of Respiratory Failure. *Acta Neurol Scand* 116: 255–258
25. Vérin E, Straus C, Demoule A, Mialon P, Derenne J, Similowski T (2002) Validation of improved recording site to measure phrenic conduction from surface electrodes in humans. *J Appl Physiol* 92: 967–74
26. Demoule A, Morelot-Panzini C, Prodanovic H, Cracco C, Mayaux J, Duguet A, Similowski T (2011) Identification of prolonged phrenic nerve conduction time in the ICU: magnetic versus electrical stimulation. *Intensive Care Med* 37:1962–8
27. An X, Yue B, Lee J, Lee M, Lin C, Han S (2012) Intramuscular distribution of the phrenic nerve in human diaphragm as shown by Sihler staining. *Muscle Nerve* 45: 522–6
28. Olsén MF, Lindstrand H, Broberg JL, Westerdahl E (2011) Measuring chest expansion; A study comparing two different instructions. *Advances in Physiotherapy* 13: 128–132)
29. Preston DC, Shapiro BE (2013) Polyneuropathy. In: Preston DC, Shapiro BE, editors. *Electromyography and neuromuscular disorders: clinical-electrophysiologic correlations*. 3rd ed. London; New York: Elsevier Saunders p. 384–416.
30. Johnson NE, Utz M, Patrick E, Rheinwald N, Downs M, Dilek N, Dogra V, Logigian EL (2014) Visualization of the diaphragm muscle with ultrasound improves diagnostic accuracy of phrenic nerve conduction studies. *MUSCLE & NERVE* 49: 669–675.
31. Shapiro BE, Katirji B, Preston DC (2014) Clinical Electromyography (Nerve Conduction Studies and Needle Electromyography). In: Katirji B, Kaminski HJ, Ruff RL, eds. *Neuromuscular Disorders in Clinical Practice*. 2nd ed. New York, NY: Springer, New York p. 89–152.
32. Ferrante MA, Teresa Spiegelberg B, Tsao B (2015) Principles of Nerve Conduction Studies and Needle EMG. AANEM 61st Annual Meeting [Internet]. [Cited 2015. Available from: https://www.aanem.org/getmedia/e9cca38c-9a6d-4ee4-bde1-ee0538e91b3/Regional-Meeting_NCS-and-EMG.pdf
33. Elsayy NA, ELwakil HS, Mourad S, Zeid M (2007) Respiratory dysfunction in uremic patients: could it be related to phrenic neuropathy? *Egypt Rheumatol.* 29(1):67–92
34. El-Tantawi GA, Imam MH, Morsi TS (2015) Phrenic Nerve Conduction Abnormalities Correlate with Diaphragmatic Descent in Chronic Obstructive Pulmonary Disease. *COPD.* 12(5):516–24.)
35. Wait JL, Nahormek PA, Yost WT, Rochester DP (1989) Diaphragmatic thickness-lung volume relationship in vivo. *J Appl Physiol* 67(4):1560–8.)
36. Dumitru D, DeLisa JA (1991) AAEM Minimonograph #10: volume conduction. *Muscle Nerve* 14(7):605–24.

37. Thimmaiah VT, Geetha M, Jain K (2016) Evaluation of thickness of normal diaphragm by B mode ultrasound. *Int J Contemp Med Res.* 3:2658–60
38. Kim M, Lee K, Cho J, Lee W (2017) Diaphragm Thickness and Inspiratory Muscle Functions in Chronic Stroke Patients. *Med Sci Monit.* 23:1247–53
39. Kim BJ, Date ES, Park BK, Choi BY, Lee SH (2005) Physiologic changes of compound muscle action potentials related to voluntary contraction and muscle length in carpal tunnel syndrome. *J Electromyogr Kinesiol.* 15(3):275–81
40. Delhez L (1979) Significance of electromyography of the diaphragm in respiratory physiopathology. *Electrodiagn Ther* 16(1):7–22.
41. Katayama Y, Senda M, Kaneda D, Ozaki T (2018) Difference between the Right and Left Phrenic Nerve Conduction Times, Latency, and Amplitude. *Acta Med Okayama.* 72(6):563–6
42. Jiang S, Xu WD, Shen YD, Xu JG, Gu YD (2011) An anatomical study of the full-length phrenic nerve and its blood supply: clinical implications for endoscopic dissection. *Anat Sci Int.*86(4):225–31
43. Lexell J (1997) Evidence for Nervous System Degeneration with Advancing Age. *J. Nutr.* 127: 1011S–1013S
44. Palve SS, Palve SB (2018) Impact of Aging on Nerve Conduction Velocities and Late Responses in Healthy Individuals. *Journal of Neurosciences in Rural Practice* 9 (1):112–116
45. Kurokawa K, Mimori Y, Tanaka E, Kohriyama T, Nakamura S (1999) Age- related change in peripheral nerve conduction: compound muscle action potential duration and dispersion. *Gerontology* 45:168–173.
46. Kimura J (2001) Principles and variations of nerve conduction studies *Electrodiagnosis in diseases of nerve and muscle: principles and practice.* 3rd ed. New York: Oxford University Press p. 91–129
47. Saadeh PB, Crisafulli CF (2003) EMG RESPIRATORY STUDIES.
48. Buschbacher RM (1998) Body mass index effect on common nerve conduction study measurements. *Muscle Nerve.* 21(11):1398–404.
49. Takazakura R, Takahashi M, Nitta N, Murata K (2004) Diaphragmatic motion in the sitting and supine positions: Healthy subject study using a vertically open magnetic resonance system. *J Magn Reson Imaging.* 19(5):605–9
50. Markand ON, Kincaid JC, Pourmand RA, Moorthy SS, King RD, Mahomed Y, et al (1984) Electrophysiologic evaluation of diaphragm by transcutaneous phrenic nerve stimulation. *Neurology.* 34(5):604–14
51. Lee JH, Kim HT, Choi IJ, Heo YR, Jung YW (2020) An unusual anatomical variant of the left phrenic nerve encircling the transverse cervical artery. *Folia Morphol (Warsz).*
52. Bolton CF (1993) AAEM minimonograph: clinical neurophysiology of the respiratory system. *Muscle Nerve.* 16(8):809–18

Tables

Table (1)

The mean, range and normative- interside difference values of sternal and LCW DCMAP parameters during inspiration and expiration

		Min.- Max.	Mean ± SD	Limit Values			
Latency (ms):	Sternal DCMAP:						
	Inspiration	5.9 - 8.6	7.08 ± 0.63	8.34	0-0.9	0.31 ± 0.20	0.71 (10.6%)
	Expiration	5.7 - 9	7.14 ± 0.67	8.48	0-0.9	0.28 ± 0.21	0.7 (10.4%)
	LCW DCMAP:						
	Inspiration	5.9 - 9.1	7.29 ± 0.67	8.63	0-0.7	0.29 ± 0.20	0.69 (10.3%)
	Expiration	5.9 - 9.6	7.34 ± 0.65	8.64	0-0.9	0.31 ± 0.21	0.73 (11.1%)
Amplitude (mV):	Sternal DCMAP:						
	Inspiration	0.6 - 2.4	1.4 ± 0.4	0.60	0-35%	14 ± 10	34%
	Expiration	0.4 - 2.2	1.4 ± 0.4	0.46	0-36%	15 ± 10	35%
	LCW DCMAP:						
	Inspiration	0.4 - 1.9	1.02 ± 0.34	0.34	0-43%	16 ± 11	38%
	Expiration	0.4 - 1.7	0.9 ± 0.29	0.32	0-43%	16 ± 10	36%
Duration (ms):	Sternal DCMAP:						
	Inspiration	8.44 - 21.3	14.54 ± 3.14	20.82	0-5.9	2.27 ± 1.5	5.27 (30%)
	Expiration	11 - 26	17.74 ± 3.19	24.12	0-5.2	2.25 ± 1.4	5.15 (25.7%)
	LCW DCMAP:						
	Inspiration	9.1 - 20.65	14.89 ± 2.96	20.81	0-5.3	2.21 ± 1.4	5.1 (29.97%)
	Expiration	10.4 - 24.1	17.77 ± 3.02	23.81	0-6	2.1 ± 1.5	5.1 (25.1%)
Area (mV.ms):	Sternal DCMAP:						
	Inspiration	2.9 - 12.9	7.27 ± 2.4	2.47	0-6.4	1.24 ± 1.13	3.5 (31.6%)
	Expiration	2.3 - 13.5	8.12 ± 2.4	3.32	0-7.6	1.23 ± 1.29	3.81 (28.8%)
	LCW DCMAP:						

Inspiration	2.2 - 12.8	5.59 ± 2.13	1.33	0-3.3	1.07± 0.76	2.59 (33.5%)
Expiration	2.6 - 13.4	6.19 ± 2.12	1.95	0- 3.85	1.05 ± 0.82	2.69 (33.2%)

Values determined from 140 phrenic nerves in 70 normal subjects.

Normative values determined as mean±2SD

Interside amplitude and area differences are expressed as the ratio between the two sides.

Table (2)

Comparison between sternal and LCW DCMAPs during inspiration and expiration

Parameter	Respiratory phase	Sternal DCMAP	LCW DCMAP	Paired t-test	P value
Latency (ms):	Inspiration	7.08 ± 0.63	7.29 ± 0.67	4.473	<0.001*
	Expiration	7.14 ± 0.67	7.34 ± 0.65	4.473	<0.001*
	t-test	2.468	2.069		
	(P₁ value)	(0.015*)	(0.04*)		
Amplitude (mV):	Inspiration	1.40 ± 0.40	1.02 ± 0.34	14.8	<0.001*
	Expiration	1.16 ± 0.35	0.90 ± 0.29	11.59	<0.001*
	t-test	12.276	10.659		
	(P₁ value)	(<0.001*)	(<0.001*)		
Duration (ms):	Inspiration	14.54 ± 3.14	14.89 ± 2.96	1.49	0.138
	Expiration	17.74 ± 3.19	17.77 ± 3.02	0.106	0.919
	t-test	18.25	18.649		
	(P₁ value)	(<0.001*)	(<0.001*)		
Area (ms.mV):	Inspiration	7.27 ± 2.40	5.59 ± 2.13	10.27*	<0.001*
	Expiration	8.12 ± 2.4	6.19 ± 2.12	10.35*	<0.001*
	t-test	7.774	7.746		
	(P₁ value)	(<0.001*)	(<0.001*)		
*: Statistically significant at p ≤ 0.05					
P: Comparison between sternal and LCW DCMAPs					
P ₁ : Comparison between DCMAPs during inspiration and expiration					

Table (3)

Comparison between right and left DCMAPs recorded from over the sternum and LCW during inspiration and expiration

Parameter	Recording site (Respiratory phase)	Right side	Left side	Paired t-test		Pearson coefficient	
				Test value	P value	r	P value
Latency (ms)	Sternal DCMAP:						
	Inspiration	7.11 ± 0.62	7.05 ± 0.65	1.297	0.199	0.831	<0.001*
	Expiration	7.20 ± 0.65	7.08 ± 0.69	3.103	0.003*	0.881	<0.001*
	LCW DCMAP:						
	Inspiration	7.36 ± 0.66	7.22 ± 0.68	3.618	0.001*	0.879	<0.001*
	Expiration	7.41 ± 0.63	7.28 ± 0.67	3.044	0.003*	0.849	<0.001*
Amplitude (mV)	Sternal DCMAP:						
	Inspiration	1.36 ± 0.42	1.44 ± 0.38	2.516	0.014*	0.775	<0.001*
	Expiration	1.16 ± 0.35	1.17 ± 0.35	0.415	0.679	0.747	<0.001*
	LCW DCMAP:						
	Inspiration	0.99 ± 0.31	1.05 ± 0.36	2.267	0.027*	0.783	<0.001*
	Expiration	0.87 ± 0.27	0.93 ± 0.31	2.747	0.008*	0.804	<0.001*
Duration (ms)	Sternal DCMAP:						
	Inspiration	14.90 ± 3.15	14.17 ± 3.11	2.212	0.031*	0.641	<0.001*
	Expiration	18.10 ± 3.42	17.38 ± 2.92	2.259	0.027*	0.680	<0.001*
	LCW DCMAP:						
	Inspiration	15.44 ± 2.97	14.33 ± 2.86	3.694	<0.001*	0.659	<0.001*
	Expiration	18.33 ± 3.20	17.20 ± 2.75	3.850	<0.001*	0.696	<0.001*
Area (ms.mV)	Sternal DCMAP:						
	Inspiration	7.20 ± 2.31	7.35 ± 2.49	0.697	0.488	0.755	<0.001*
	Expiration	8.14 ± 2.35	8.10 ± 2.47	0.180	0.858	0.725	<0.001*
	LCW DCMAP:						
	Inspiration	5.48 ± 2.06	5.71 ± 2.21	1.413	0.163	0.814	<0.001*
	Expiration	6.16 ± 2.11	6.22 ± 2.16	0.374	0.709	0.802	<0.001*

Table (4)

Comparison between DCMAPs recorded in men and women (n=70).

		Men	Women	t-test	P value
	Recording site (Respiratory phase)				
Latency (ms)	Sternal DCMAP:				
	Inspiration	7.21 ± 0.61	6.81 ± 0.59	3.7	<0.001*
	Expiration	7.30 ± 0.62	6.83 ± 0.66	4.12	<0.001*
	LCW DCMAP:				
	Inspiration	7.49 ± 0.59	6.87 ± 0.63	5.761	<0.001*
	Expiration	7.54 ± 0.59	6.95 ± 0.59	5.608	<0.001*
Amplitude (mV)	Sternal DCMAP:				
	Inspiration	1.56 ± 0.37	1.07 ± 0.22	8.227	<0.001*
	Expiration	1.28 ± 0.34	0.91 ± 0.19	6.847	<0.001*
	LCW DCMAP:				
	Inspiration	1.11 ± 0.32	0.84 ± 0.31	4.782	<0.001*
	Expiration	0.97 ± 0.28	0.76 ± 0.27	4.123	<0.001*
Duration (ms)	Sternal DCMAP:				
	Inspiration	14.90 ± 3.14	13.83 ± 3.05	1.871	0.063
	Expiration	17.85 ± 3.17	17.53 ± 3.24	0.549	0.587
	LCW DCMAP:				
	Inspiration	15.32 ± 3.09	14.05 ± 2.52	1.859	0.068
	Expiration	18.00 ± 3.20	17.30 ± 2.61	1.252	0.213
Area (ms.mV)	Sternal DCMAP:				
	Inspiration	8.23 ± 2.12	5.41 ± 1.72	7.62*	<0.001*
	Expiration	8.98 ± 2.20	6.44 ± 1.84	6.554*	<0.001*
	LCW DCMAP:				
	Inspiration	6.28 ± 1.82	4.26 ± 2.07	5.696*	<0.001*
	Expiration	6.71 ± 1.98	5.15 ± 2.04	4.155*	<0.001*
*: Statistically significant at $p \leq 0.05$					

Table (5)

Correlation between phrenic nerve conduction parameters and subjects' characteristics

		Age (years)		Height (m)		BMI (kg/m ²)		Chest circumference (cm)	
		r	p	r	p	r	p	r	p
Latency	Sternal DCMAP:								
	Inspiration	0.622	<0.001*	0.335	0.005*	0.046	0.705	0.319	0.007*
	Expiration	0.590	<0.001*	0.357	0.002*	0.042	0.732	0.288	0.016*
	LCW DCMAP:								
	Inspiration	0.435	<0.001*	0.436	<0.001*	-0.156	0.197	0.133	0.272
	Expiration	0.413	<0.001*	0.415	<0.001*	-0.113	0.353	0.144	0.236
Amplitude	Sternal DCMAP:								
	Inspiration	0.053	0.666	0.157	0.194	-0.425	<0.001*	-0.146	0.227
	Expiration	0.082	0.498	0.077	0.524	-0.288	0.016*	-0.004	0.972
	LCW DCMAP:								
	Inspiration	-0.017	0.888	0.009	0.944	-0.344	0.004*	-0.239	0.047*
	Expiration	0.060	0.623	-0.048	0.696	-0.272	0.023*	-0.193	0.109
Duration	Sternal DCMAP:								
	Inspiration	-0.004	0.976	0.313	0.011*	-0.323	0.009*	-0.204	0.103
	Expiration	-0.136	0.282	0.201	0.108	-0.394	0.001*	-0.358	0.003*
	LCW DCMAP:								
	Inspiration	-0.079	0.532	0.358*	0.003*	-0.162	0.196	0.066	0.601
	Expiration	-0.079	0.534	0.375*	0.002*	-0.153	0.223	0.030	0.815
Area	Sternal DCMAP:								
	Inspiration	-0.051	0.688	0.319	0.010*	-0.545	<0.001*	-0.310	0.012*
	Expiration	-0.016	0.902	0.270	0.010*	-0.447	<0.001*	-0.210	0.094
	LCW DCMAP:								
	Inspiration	-0.062	0.626	0.179	0.155	-0.365	0.003*	-0.178	0.157
	Expiration	0.002	0.986	0.125	0.322	-0.244	0.050*	-0.144	0.253
r: Pearson coefficient									

*: Statistically significant at $p \leq 0.05$

Table (6)

Multivariate analysis for the parameters affecting DCMAP parameters

				Unstandardized Coefficients		Standardized Coefficients	Test of sig.	P value
		Recording site	Respiratory phase	B	Std. Error	Beta		
Latency	Sternal DCMAP	Inspiration	(Constant)	3.495	1.613		2.167	0.034
			Age	0.029	0.005	0.556	6.048	0.000*
		Expiration	(Constant)	3.473	1.761		1.972	0.053
			Age	0.030	0.005	0.528	5.608	0.000*
	LCW DCMAP	Inspiration	(Constant)	4.390	1.750		2.508	0.015
			Age	0.022	0.005	0.384	3.973	0.000*
			Gender	-0.419	0.166	0.305	2.530	0.014*
		Expiration	(Constant)	4.922	1.716		2.869	0.006
			Age	0.020	0.005	0.365	3.685	0.000*
			Gender	0.414	0.163	0.315	2.549	0.013*
Amplitude	Sternal DCMAP	Inspiration	(Constant)	2.329	0.183		12.745	0.000
			Gender	0.432	0.083	0.543	5.227	0.000*
		Expiration	(Constant)	1.732	0.174		9.967	0.000
			Gender	0.351	0.079	0.511	4.470	0.000*
	LCW DCMAP	Inspiration	(Constant)	2.819	0.538		5.239	0.000
			Gender	0.354	0.101	0.524	3.507	0.001*
			Chest circumference	-0.017	0.007	-0.406	-2.335	0.023*
		Expiration	(Constant)	1.322	0.162		8.166	0.000
Gender	-0.167		0.073	0.287	2.276	0.026*		
Duration	LCW DCMAP	Inspiration	(Constant)	4.339	6.328		0.686	0.495
			Height	11.415	3.753	0.358	3.042	0.003*
		Expiration	(Constant)	3.008	6.484		0.464	0.644
			Height	12.334	3.845	0.375	3.208	0.002*
Area	Sternal DCMAP	Inspiration	(Constant)	22.798	5.887		3.873	0.000
			Gender	3.119	0.621	0.661	5.019	0.000*
			Chest circumference	-0.115	0.045	-0.402	-2.570	0.013*
		Expiration	(Constant)	19.644	6.555		2.997	0.004
			Gender	2.301	0.627	0.490	3.671	0.001*

		BMI	-0.120	0.050	-0.276	-2.400	0.019*
LCW DCMAP	Inspiration	(Constant)	9.803	1.146		8.556	0.000
		Gender	-1.651	0.518	-0.388	3.186	0.002*
	Expiration	(Constant)	9.068	1.225		7.404	0.000
		Gender	-1.344	0.554	-0.317	-2.425	0.018*

Figures

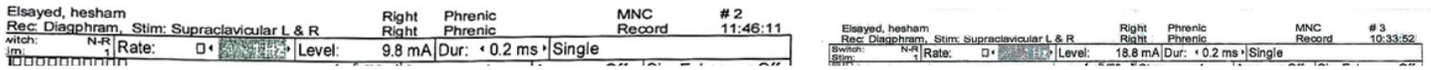


Figure 1

DCMAPs recorded from over the sternum and LCW in one of the studied subjects showing shorter latency and duration, higher amplitude, and area in sternal compared to LCW recordings