Size and strength of the respiratory and quadriceps muscles in patients with chronic asthma

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ABSTRACT: There have been few studies of respiratory and limb muscle size and function in middle-aged patients with asthma and persistent airways obstruction.

We have compared the forces generated by the respiratory and thigh muscles with their dimensions assessed by ultrasound in nine middle-aged patients with chronic asthma (mean age 56 (so 8) yrs; functional residual capacity/total lung capacity ratio (FRC/TLC) 60 (10) %), and in nine normal subjects (aged 53 (7) yrs; FRC/TLC 55 (5) %). Diaphragm thickness was measured at the zone of apposition by B-mode ultrasound during relaxation (DiTrelax) and during a maximum-effort inspiratory manoeuvre (DiTPLmax) at FRC. Cross-sectional area of the relaxed rectus femoris muscle (Aur) was determined by ultrasound at mid-thigh level. Isometric strength of the right quadriceps muscle group was measured during maximum voluntary contraction.

Asthmatic patients had preserved quadriceps strength and Aur but moderately impaired maximum inspiratory pressure (PImax) (-52 (18) cmH2O) and thicker DiTrelax (2.2 (0.4) mm), compared to normal subjects (-73 (21) cmH2O and 1.7 (0.3) mm, respectively).

Middle-aged patients with chronic asthma and a small increase in functional residual capacity/total lung capacity ratio have preserved limb muscle force and dimensions, modestly impaired inspiratory muscle strength, and slightly increased thickness of the costal diaphragm. Future studies of respiratory muscle function in asthma should be aided by measurement of diaphragm thickness and of limb muscle strength and size. Such studies are required particularly in older patients with severe hyperinflation who are most likely to have impairment of muscle function.

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Many studies have been made of the morphometric [1, 2] and functional [3–7] characteristics of the respiratory muscles in emphysema and relatively advanced chronic obstructive pulmonary disease. In such patients, the major factors impairing force generation by the respiratory muscles appear to be hyperinflation, loss of body weight and, by implication, loss of respiratory muscle mass [8]. Hyperinflation would affect force generated by the inspiratory muscles selectively; generalized loss of weight and skeletal muscle mass might reduce the force generated both by inspiratory and expiratory muscles, as well as by limb muscles. Despite the suspicion that fatigue of the inspiratory muscles may be a critical factor in the most severe attacks of asthma, few studies of the respiratory muscles have been performed in patients with asthma [5, 9–13]. In particular, older subjects with persistent airflow obstruction due to chronic asthma have been studied little and there is no information on respiratory muscle mass. Because skeletal muscle strength is known to decline with age [14], any effects of treatment and disease per se are more likely to be found in this subgroup.

Ultrasound has proved to be useful for the study of anatomical characteristics of many muscle groups, including the quadriceps muscle [15]. Recently, high-resolution ultrasound has been used to assess diaphragmatic changes during tidal breathing and during relaxation at different lung volumes in normal subjects [16, 17]. Using B-mode ultrasound, we have shown that it is possible to image the costal portion of the diaphragm, both at rest and during maximum voluntary contractions [18]. This technique has not been applied to patients. In this study, we have compared the forces generated by the inspiratory and thigh muscles with their dimensions assessed by ultrasound in nine middle-aged patients with chronic asthma and in nine age-matched normal subjects.

Methods

Subjects

Nine patients (one current smoker, five ex-smokers, and three lifetime nonsmokers) with asthma of more than 10 yrs duration and persistent airway obstruction despite treatment took part in the study. In three patients, onset of asthma had been in childhood; in the remaining six patients asthma had developed between the ages of 21 and 59 yrs. Six patients, including the three with
childhood onset, had positive skin tests to common aero-
allergens. The diagnosis of asthma was based on evidence of
more than 30% variation in spirometry (spontaneous or after bronchodilator or corticosteroid treatment), a nor-
mal chest radiograph, and normal carbon monoxide trans-
ferr coefficient (mean (sd) 122 (29) % of predicted values
[19]). Although six of the subjects were current or ex-
smokers, this history did not appear relevant to the aeti-
ology of the airways obstruction. Thus, the one current
smoker was an atopic female whose symptoms com-
menced at 21 yrs of age. Symptoms commenced in early
childhood in one of the ex-smokers, and the remaining
four ex-smokers had given up in middle-age 3–15 years
before the onset of an initially highly variable airways
obstruction.

All were regularly using inhaled glucocorticosteroids
and beta-agonists, and had had previous short courses
of oral steroids. They all maintained moderate habitual
activity without any specific training programmes, and
were considered by their physicians to have been in a
stable condition in the previous 3 months. Nine non-
smoking normal subjects were studied as controls. They
were also moderately active but none participated in ath-
etic activities; they had no respiratory symptoms and
no history of cardiovascular or neuromuscular disease.

Methods

Height and weight were recorded and ideal body weight
was estimated from the updated Metropolitan Life In-
surance Tables [20]. Spirometry was recorded using a
Fukuda Sangyo spiro-analysrer (ST250). A constant-vol-
ume body plethysmograph was used to measure total
lung capacity (TLC), functional residual capacity (FRC)
and residual volume (RV) [21]. Predicted values were
derived from Cotes [22].

Respiratory muscle assessment. Real-time movement of
the diaphragm was recorded by B-mode ultrasonography.
A 7.5 MHz ultrasound linear probe (PLE 7055 Toshiba
Medical System, Tokyo, Japan) was held perpendicular
to the chest wall in the 9th or 10th intercostal space,
between the antero- and mid-axillary lines. If the angle
of incidence was changed significantly by movement of
the transducer head, the returning ultrasound signal of the
diaphragm became distorted or was lost altogether, as
reported previously [16, 18]. The diaphragm was observed
in the zone of apposition: two clear outer parallel lines
corresponded to the pleural and peritoneal membranes,
with an irregular bright layer of connective tissue and
vessels within a relatively echo-free muscle layer (fig.
1). Diaphragm thickness was measured from the middle
of the pleural to the middle of the peritoneal line. The
shortest distance that could be resolved between the two
lines using this system was 0.3 mm, considerably less
than the thickness of the normal relaxed diaphragm. Mea-
surements of thickness were made to the nearest 0.1 mm
with callipers. On each ultrasound image, three measure-
ments of thickness were made at 0.5–2.0 cm below the
costophrenic sinus and the results averaged.

Initially, while the subject was sitting comfortably,
wearing a noseclip and breathing through a conventional
mouthpiece, three separate measurements of resting
diaphragm thickness at FRC were taken and the mean
value (DiTrelax) used for analysis. The subject was then
asked to perform a maximum-effort inspiratory mano-
euvre against a closed airway at FRC for 2–3 s, while di-
aphragm thickness was assessed using ultrasound. The
pressure generated at the mouthpiece was recorded using
a variable inductance differential pressure transducer (HP
267 B, Hewlett Packard), with a small internal volume
and linear response over the range of 200 to 200 cmH₂O.
A small leak was introduced in the system to ensure glu-
tal patency and to avoid the use of cheek muscles. The
procedure was repeated at least three times. Maximum-
effort inspiratory mouth pressure (P₁max) was taken as
the most negative pressure sustained for at least 1 s, and
diaphragm thickness obtained during this manoeuvre
(DiTPLmax) was selected for analysis. Increase in diaph-
granm thickness during this maximum-effort inspiratory
manoeuvre was calculated as the thickening ratio (TRdi):

\[
TRdi = \frac{DiTPL_{max}}{DiT_{relax}}
\]

Maximum-effort expiratory mouth pressure (PEmax) was
measured near TLC, and the best of three technically
satisfactory manoeuvres sustained for at least 1 s was
selected for analysis. Normal values for mouth pressures
were taken from Wilson et al. [23].

Limb muscle assessment. Muscle strength was deter-
mined as the maximum voluntary force that could be
developed by the knee extensor muscles during isometric
contractions in the (dominant) right leg. The apparatus
used was similar to that described by Edwards et al.
[24]. All subjects were studied while seated in an adjust-
able straight-backed chair. The pelvis was secured by a
belt to prevent hip joint extension. The lower leg was
kept dependent and the knee flexed to 90° with its pos-
terior aspect positioned at the front edge of the chair.
Force was measured with a strap looped around the right
leg just proximal to the ankle and connected to a strain
gauge. The amplified output from this system was dis-
played on a pen recorder. At least three maximum vol-
untary contractions were recorded for each subject, and
the best taken for analysis. Before each contraction, the cross-sectional area of the relaxed homolateral rectus femoris muscle (Arf) was determined using B-mode ultrasound. Ultrasonograms were obtained half-way between the major trochanter and the lateral joint-line of the knee using a 5 MHz linear transducer (PLE-505S; Toshiba) placed gently on the anterior aspect of the thigh to avoid distortion of the underlying tissue. The quality of image allowed easy identification of the rectus femoris muscle in all cases (fig 2). The image displayed on a monitor was outlined using a movable cursor, and the area within the line automatically calculated by a planimetric technique. The Arf was taken to be the mean of at least three separate measurements.

Written informed consent was obtained in all cases, and the protocol was approved by the local Research Ethics Committee.

**Statistical analysis**

Spearman rank correlation coefficients were used to define correlations. The significance of differences between means was assessed by two-tailed unpaired t-test or the Mann-Whitney test, as appropriate. Group results are expressed as means with standard deviations in parentheses.

**Results**

The patients with chronic asthma showed considerable reductions in spirometry and an increase in RV/TLC ratio, and a small, nonsignificant increase in FRC/TLC ratio (table 1).

Good quality images of the costal diaphragm were obtained in all the normal subjects and patients. Resting muscle thickness (DiTrelax) was greater in patients with chronic asthma than in normal controls (table 2 and fig. 3) and was significantly correlated to body weight in both groups (r²=0.579; p=0.017; r²=0.485; p=0.037,

| Table 1. – Subject characteristics, spirometry and lung volumes |
|-------------------|-------------------|-----------------|---------|
|                   | Control           | Asthma          | p-value |
| Sex M/F           | 4/5               | 3/6             | 0.376†  |
| Age yrs           | 53 (7)            | 56 (8)          | 0.490†  |
| Body weight kg    | 67.7 (12.0)       | 65.0 (8.5)      | 0.022†  |
| % ideal           | 106 (17)          | 108 (9)         | 0.825†  |
| FEV₁ L            | 3.04 (0.69)       | 1.43 (0.41)     | <0.001† |
| % pred            | 104 (14)          | 58 (21)         | <0.001† |
| FVC L             | 3.73 (0.81)       | 2.75 (0.65)     | 0.012†  |
| % pred            | 106 (11)          | 92 (24)         | 0.102†  |
| FEV₁/FVC %        | 82 (6)            | 53 (12)         | <0.001† |
| RV/TLC %          | 32 (6)            | 47 (13)         | 0.012†  |
| FRC/TLC %         | 55 (5)            | 30 (10)         | 0.171†  |
| TLC L             | 5.51 (1.03)       | 5.31 (0.86)     | 0.661†  |

Values are presented as mean, and so in parenthesis. M/F: male/female; FEV₁: forced expiratory volume in one second; FVC: forced vital capacity; RV: residual volume; TLC: total lung capacity; FRC: functional residual capacity; % pred: percentage of predicted values; †: control vs asthma group, unpaired t-test; ‡: control vs asthma group, Mann-Whitney test.

| Table 2. – Respiratory and limb muscle strength and dimensions assessed by ultrasonography |
|---------------------|-------------------|-----------------|---------|
|                     | Control           | Asthma          | p-value |
| P₁max cmH₂O        | -73 (21)          | -52 (18)        | 0.037†  |
| % pred              | 92 (25.4)         | 73 (28.0)       | 0.085‡  |
| DiTrelax mm         | 1.7 (0.3)         | 2.2 (0.4)       | 0.022‡  |
| DiTmax mm           | 4.0 (1.2)         | 4.3 (0.9)       | 0.587‡  |
| TRₘ max mm H₂O      | 2.2 (0.6)         | 2.0 (0.3)       | 0.223†  |
| Pₑ₁max cmH₂O       | 90 (25)           | 73 (29)         | 0.209†  |
| % pred              | 81 (16.7)         | 72 (27.9)       | 0.102‡  |
| QF N                | 281 (114)         | 317 (92)        | 0.517†  |
| Arf mm²             | 626 (165)         | 603 (270)       | 0.844‡  |

Values are presented as mean, and so in parenthesis. †: eight subjects only; ‡: seven subjects only; P₁max: maximum-effort inspiratory mouth pressure; DiTrelax: resting diaphragm muscle thickness at functional residual capacity; DiTmax: diaphragm muscle thickness during P₁max manoeuvre; TRₘ: diaphragm thickening ratio; Pₑ₁max: maximum-expiratory pressure; QF: quadriceps muscle strength measured during maximum voluntary contraction; Arf: cross-sectional area of the rectus femoris muscle; % pred: percentage of predicted values; †: control vs asthma group, unpaired t-test; ‡: control vs asthma group, Mann-Whitney test.
respectively). During maximum-effort inspiratory manoeuvres at FRC, average TRdi was 2.2 in normal subjects and 2.0 in asthmatic subjects (table 2). On average, less negative $P_{E,max}$ was generated by the asthmatic than by the normal group (-52 (18) and -73 (21) cmH$_2$O, respectively; p=0.037) (table 2 and fig. 3). A significant correlation between DiTrelax and $P_{E,max}$ was found in normal ($r^2=0.559$; p<0.01) but not in asthmatic subjects. Some reduction in $P_{E,max}$ was found in asthmatic compared to the normal subjects, without statistical significance (table 2). There was no significant difference in the strength of the quadriceps muscle group between asthmatic patients (317 (92) N) and normal controls (281 (114) N) (table 2). Mean ARF during relaxation was similar in control (626 (165) mm$^2$) and asthmatic subjects (603 (270) mm$^2$). ARF showed a strong positive correlation with the strength of the quadriceps muscle group in normal controls ($r^2=0.796$; p<0.003) but not in asthmatic patients.

**Discussion**

These results show that middle-aged patients with chronic, incompletely reversible asthma and minor increases in FRC/TLC% have slightly increased resting diaphragm thickness and a normal thickening ratio during maximum inspiratory efforts despite some impairment of maximum inspiratory pressures. Quadriceps mass and strength were normal.

We are not aware of earlier studies using ultrasound to image the costal diaphragm in patients with asthma. In the present control group, the mean DiTrelax of 1.7 (0.3) mm was identical to that found in our earlier study of 10 seated normal male subjects. TANIGUCHI et al. [17] using B-mode ultrasound observed a mean diaphragm thickness of 2.0 (0.5) mm at FRC in 61 supine normal subjects of both sexes. Diaphragm thickness at FRC was greater in males (2.2 (0.5) mm) than females (1.9 (0.3) mm; p<0.01) and was positively correlated to body weight (r=0.53; p<0.01) and weakly to height (r=0.28; p<0.05). In the present study, average resting diaphragm thickness was slightly increased in asthmatic patients compared to normal controls, although it was very close to the normal values reported previously [16–18].

We also found no abnormality of the size and strength of the quadriceps muscle in asthmatic subjects. The force generated by limb muscles was positively correlated to their cross-sectional area (determined by ultrasound) as originally shown for the arm flexors in young normal subjects [25]. Subsequently, YOUNG et al. [26] studied 25 normal subjects (11 males and 14 females) aged 19–48 yrs, and reported a stronger correlation between cross-sectional area and the torque exerted by the quadriceps muscle (r=0.84) than between body weight and quadriceps torque (r=0.54). They also reported that the reduced isometric strength in elderly females correlated with reduced ultrasonographic dimensions of the quadriceps muscle [15]. Other studies using computed tomography have confirmed a significant positive correlation between quadriceps muscle strength and cross-sectional area [27, 28]. The correlations that we obtained in the normal individuals between isometric strength and $ARF$ were actually stronger than previously reported for the whole quadriceps group [15, 26].

The only widely-used clinical test of respiratory muscle force in asthma has been the measurement of mouth force in asthma has been the measurement of mouth

### Table 3. Previous studies of mouth pressures in adult asthmatic patients

<table>
<thead>
<tr>
<th>First author</th>
<th>[Ref.]</th>
<th>Group</th>
<th>M/F</th>
<th>Age yrs</th>
<th>FEV$_1$ % pred</th>
<th>FRC % pred</th>
<th>FRC TLC</th>
<th>$P_{E,max}$ cmH$_2$O</th>
<th>$P_{E,max}$ cmH$_2$O</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Decramer</td>
<td>[5]</td>
<td>Asthma</td>
<td>7/0</td>
<td>42 (15)</td>
<td>47 (10)</td>
<td>-</td>
<td>61 (10)</td>
<td>-65$^*$</td>
<td>137$^*$</td>
<td>After in-hospital treatment, also studied inspiratory and expiratory muscle endurance</td>
</tr>
<tr>
<td>McKenzie</td>
<td>[9]</td>
<td>Asthma</td>
<td>10/10</td>
<td>34 (15)</td>
<td>76 (15)$^*$</td>
<td>125 (21)$^*$</td>
<td>-</td>
<td>-113$^*$</td>
<td>135$^*$</td>
<td>Acute bronchospasm Stable; slightly more negative $P_{I,max}$ after bronchodilator</td>
</tr>
<tr>
<td>Laviotes</td>
<td>[10]</td>
<td>Asthma</td>
<td>9/11</td>
<td>39 (3)</td>
<td>47 (6)$^*$</td>
<td>121 (10)</td>
<td>-</td>
<td>-97 (8.7)$^*$</td>
<td>-136 (6.8)</td>
<td>Less negative $P_{I,max}$ after inducing increase in FRC</td>
</tr>
<tr>
<td>Weiner</td>
<td>[11]</td>
<td>Asthma</td>
<td>5/5</td>
<td>33 (4)</td>
<td>64 (3)$^*$</td>
<td>123 (5)$^*$</td>
<td>-</td>
<td>-82 (3.7)</td>
<td>-95 (5.9)</td>
<td>Also studied $P_{E,max}$ and inspiratory muscle endurance after induced increase in FRC</td>
</tr>
<tr>
<td>Gorman</td>
<td>[13]</td>
<td>Asthma</td>
<td>4/2</td>
<td>26 (7)</td>
<td>96 (19)</td>
<td>118 (29)</td>
<td>48 (7)</td>
<td>-114 (22)</td>
<td>-</td>
<td>Stable</td>
</tr>
<tr>
<td>De Bruin</td>
<td>(present study)</td>
<td>Asthma</td>
<td>3/6</td>
<td>56 (9)</td>
<td>58 (21)$^*$</td>
<td>113 (26)</td>
<td>60 (10)</td>
<td>-52 (18)$^*$</td>
<td>73 (29)</td>
<td></td>
</tr>
</tbody>
</table>
pressures generated during maximal inspiratory and expiratory efforts against a closed valve [22, 29]. Although \( P_{\text{I}_{\text{max}}} \) assesses the overall force produced by the inspiratory muscles, maximum activation of the diaphragm is usually achieved with this manoeuvre, at least in normal subjects [30, 31]. In the present study, there was a modest reduction in \( P_{\text{I}_{\text{max}}} \) despite the increase in diaphragm thickness. Previous studies of \( P_{\text{I}_{\text{max}}} \) in asthma, in which FRC was measured, are summarized in table 3. All have involved small numbers of subjects and, with the exception of the study by LAVIETES et al. [10], FRC has only been modestly increased. Only one study has examined the effects of inducing acute hyperinflation in asthma [13], although clearly this cannot be assumed to have the same effect as a more chronic increase in FRC. Most earlier studies have been made in adults under 40 yrs of age. Nevertheless, the results are compatible with a trend in asthmatic subjects to less negative values of \( P_{\text{I}_{\text{max}}} \) with increasing age and/or increase in FRC. There appears to be no consistent trend to any alteration in \( P_{\text{E}_{\text{max}}} \).

Clearly, any comprehensive study of the respiratory muscles in asthma has to take account of age, gender, the degree of hyperinflation and corticosteroid treatment. We suggest that the strength and size of thigh muscles and estimates of diaphragmatic mass are also likely to be relevant. Hyperinflation leads to passive shortening of the diaphragm, reducing its capacity to generate inspiratory force due to a shift to a less efficient part of its force-length relationship [32, 33]. In preliminary studies, we have found a tendency for DiTrelax to increase with increasing lung volume [18] presumably due to muscle shortening. While this is of considerable importance in assessing the costal diaphragm in emphysema, there was only a small increase in FRC/TLC in the present asthmatic subjects, so this seems unlikely to account for the increased DiTrelax, which may indicate some muscle hypertrophy. We do not know whether the modest impairment of \( P_{\text{I}_{\text{max}}} \) in the present asthmatic subjects can be attributed to this slight hyperinflation or not; indeed, if there was abdominal muscle contraction at FRC, diaphragm length might be greater than suggested by the FRC/TLC ratio, but chest wall configuration was not measured in these studies. Another possible explanation is reduced activation of the rib cage inspiratory muscles. The patients were trained in the maximum inspiratory manoeuvre but, as in previously reported studies (table 3), we have no proof that the inspiratory muscles were activated maximally.

Alternatively, the reduction in \( P_{\text{I}_{\text{max}}} \) (and trend to lower \( P_{\text{E}_{\text{max}}} \)) might be related to use of corticosteroids. Although all the patients in this study had had booster courses of oral prednisolone within the preceding 12 months, none was on maintenance oral corticosteroids and had had a booster course of oral corticosteroids within the preceding 3 months. A careful study by PICADO et al. [12] found no difference in respiratory and shoulder muscle strength between 34 middle-aged patients with asthma, who were taking an average prednisone dose of 11.9 mg·day\(^{-1}\), and 34 matched control patients, who had never been on continuous steroid treatment. Respiratory muscle strength appeared normal in both groups of patients. In contrast, a recent study of 21 patients with airways obstruction (six of whom had asthma), receiving treatment with corticosteroids and studied on the 10th day after hospital admission [34], found that mean oral corticosteroid dosage during the preceding 6 months was inversely related to \( P_{\text{I}_{\text{max}}} \) and, more strongly, to quadriceps force. Eighteen of these 21 patients received methylprednisolone or triamcinolone, which are believed to have greater deleterious effects on muscle bulk and strength than the prednisolone used for booster courses in the present patients. Since quadriceps strength was normal in the asthmatic patients in the present study, it seems unlikely that their reduction in \( P_{\text{I}_{\text{max}}} \) can be attributed to corticosteroid usage.

In summary, middle-aged patients with chronic asthma have preserved limb muscle force and dimensions but slightly impaired inspiratory muscle pressure generation. Relaxed diaphragm thickness was slightly increased. Further studies of older patients with larger increases in functional residual capacity/total lung capacity ratio are required; these patients are likely to have the most significant chronic functional impairment of inspiratory muscles.

References


13. Gorman RB, McKenzie DK, Gandevia SC, Plassman...


