Non-invasive proportional assist and pressure support ventilation in patients with cystic fibrosis and chronic respiratory failure

A Serra, G Polese, C Braggion, A Rossi

Background: Patients with advanced cystic fibrosis can benefit from non-invasive positive pressure ventilation (NPPV) for the treatment of acute decompensation as well as for the management of chronic respiratory failure. This study was undertaken to compare the physiological effects of non-invasive proportional assist ventilation (PAV) and pressure support ventilation (PSV) on ventilatory pattern, transcutaneous blood gas tensions, and diaphragmatic effort in stable patients with cystic fibrosis and chronic CO₂ retention.

Methods: In 12 patients two periods of spontaneous breathing were followed randomly by PSV (12 (3) cm H₂O) and PAV (flow assist 4.9 (1.3) cm H₂O/l.s, volume assist 18.9 (5.1) cm H₂O/l) set for the patient’s comfort and administered for 40 minutes with 2 cm H₂O continuous positive airway pressure. Ventilatory pattern, transcutaneous blood gas tensions, and surface diaphragmatic electromyography were measured in the last 10 minutes of each application.

Results: Both PSV and PAV improved ventilation (+30%), tidal volume (+30%), and transcutaneous CO₂, (-7%) while reducing diaphragmatic activity (-30% with PSV, -20% with PAV). Mean inspiratory airway pressure was lower during PAV than during PSV. Mean inspiratory breathing frequency was higher during PAV than during PSV. The mean coefficient of variation of tidal volume was about 20% (range 11–39%) during spontaneous breathing and did not change with either PAV or PSV.

Conclusions: These results show that short term administration of nasal PAV and PSV to patients with stable cystic fibrosis with chronic respiratory insufficiency is well tolerated, improves ventilation and blood gas tensions, and unloads the diaphragm.

Patients with advanced cystic fibrosis (CF) can benefit from non-invasive positive pressure ventilation (NPPV) in the treatment of acute decompensation as well as in the management of chronic respiratory failure. In some studies NPPV has been delivered in the assist/control mode whereas other authors have used pressure support ventilation (PSV). In recent years a new mode of partial ventilatory support—proportional assist ventilation (PAV)—has been developed with the aim of improving the interaction between the patient and the ventilator.

PAV is a patient guided mode of synchronised partial assistance in which the ventilator pressure output is proportional to the instantaneous effort of the patient. In this mode there is therefore automatic synchrony between the patient’s effort and the ventilator cycle. With PAV the level of pressure delivered to the patient increases and decreases according to the demand of the patient, so responsibility for the level and pattern of ventilatory assistance depends entirely upon the patient. A few studies have shown that PAV can produce physiological benefits in patients with acute and chronic respiratory failure.

However, PAV has never been used in patients with CF.

In view of the theoretical advantages of PAV, we have studied the acute physiological effects of PAV in patients with severe CF and chronic ventilatory failure (CVF). Furthermore, because few physiological data are available in patients during NPPV, we have compared the effect of PAV with PSV.

METHODS

The study was approved by the institution ethical committee (Azienda Ospedaliera di Verona, Italy) and was conducted according to the declaration of Helsinki. Informed consent was obtained from the patients before enrolment into the study.

Patients

Twelve patients (eight men) with CVF due to CF were recruited into the study. CVF is defined as a consistent increase in the arterial partial pressure of carbon dioxide (PaCO₂) above 6 kPa (45 mm Hg) during spontaneous breathing of room air. The patients were all in a stable clinical condition when they were recruited into the study, as assessed by stable blood gas tensions and pH, and had been free from exacerbations during the preceding 2 weeks. The characteristics of the patients are shown in table 1.

Measurements

Lung volumes and arterial blood tensions were measured from 1 day to 1 week and from 1–2 days before the study according to standard procedures. Transcutaneous CO₂ and O₂ tensions were measured using TCM3 (Radiometer; Copenhagen, Denmark).

Flow (V) and volume (V) by numerical integration were measured by means of a heated pneumotachograph (3700 Series, Hans Rudolph Inc, Kansas City, MO, USA) connected to...
a pressure transducer (Sefam MV+, INSERM, Nancy, France) inserted between the nasal mask and the “plateau valve” of the NPPV circuit. Pressure at the airway opening (Pao) was measured with a differential pressure transducer (Sefam MV+) connected to one port of the nasal mask. Surface electromyography of the diaphragm (Edi) was measured with an isolated amplifier (Physio-Amp, Francesco Marazza, Monza, Italy). All signals were digitised at a sampling frequency of 1000 Hz and analysed using the software package WINDAQ and ADVANCED CODAS (DATAQ Instruments, Ohio, USA).

### Data analysis

**Breathing pattern and airway pressure**

Tidal volume (Vt), respiratory frequency (f), minute ventilation (V′e), and inspiratory capacity (IC) were computed from the volume signal. Total cycle duration (Ttot), inspiratory time (Tt), expiratory time (Te), and Tt/Ttot were calculated from the flow signal as mean values from 10 minute continuous recordings of flow and volume. The variability in Vt was analysed by calculating the coefficient of variation as the ratio of standard deviation over the mean Vt value. Pao was measured as the peak value (Pao,peak) as well as the pressure time integral over Tt (Pao,Tt) and Ttot (Pao,Ttot), and the resulting area was divided by the duration of Tt and Ttot, respectively.

**Diaphragmatic electromyography**

The diaphragmatic electromyogram (Edi) was recorded and filtered as previously described. From the filtered Edi signal the total duration of the Edi activity (Tt,Edi) was computed as well as the time between the onset of one burst of activity and that of the next Edi burst (Ttot,Edi) to compute the Edi duty cycle (Tt/Edi/Ttot,Edi). The Edi was digitally rectified and processed with the moving mean using a time window of 0.1 seconds. From the moving mean Edi we also measured the peak amplitude in arbitrary units (Edi,peak) expressed as a percentage of the value recorded during spontaneous breathing. The integral of the rectified Edi signal over Tt,Edi was measured and this value was multiplied by the respiratory frequency to obtain the electric power used by the diaphragm over 1 minute (Edi,int).

**Setting of ventilator**

Non-invasive ventilation was delivered through a commercial nasal mask (Respironics, Murrysville, PA, USA) by means of a Vision ventilator (Respironics) set at a continuous positive airway pressure (CPAP) of 2 cm H2O. PSV was set initially at 8 cm H2O in all patients and the pressure was then increased progressively in steps of 1 cm H2O until the patient felt uncomfortable with the level of assistance. Throughout the procedure the last level of PSV at which the patient felt comfortable was used.

### RESULTS

All the patients tolerated both PSV and PAV throughout the procedure. Changes in breathing pattern, transcutaneous blood gas tensions, and differences between PSV and PAV are shown in table 2. No significant difference was observed between the two spontaneous breathing control conditions. PAV and PSV were compared with the immediately preceding period of spontaneous breathing. With both PSV and PAV minute ventilation (+30%) and tidal volume (+30%) were significantly increased while respiratory frequency did not change. Inspiratory capacity remained stable throughout the procedure. TcCO2 decreased with both PSV and PAV while TcO2

---

**Table 1 Mean (SD) demographic, anthropometric, and functional characteristics of study patients (n=12)**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Value (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>M/F</td>
<td>8/4</td>
</tr>
<tr>
<td>Age (years)</td>
<td>28 (6)</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>166 (8)</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>53 (10)</td>
</tr>
<tr>
<td>FEV1 (% predicted)</td>
<td>20 (9)</td>
</tr>
<tr>
<td>FVC (% predicted)</td>
<td>44 (9)</td>
</tr>
<tr>
<td>FEV1/FVC</td>
<td>41 (10)</td>
</tr>
<tr>
<td>RV (% predicted)</td>
<td>249 (69)</td>
</tr>
<tr>
<td>TLC (% predicted)</td>
<td>104 (23)</td>
</tr>
<tr>
<td>pH</td>
<td>7.38 (0.02)</td>
</tr>
<tr>
<td>Pao2 (kPa)</td>
<td>7.1 (0.8)</td>
</tr>
<tr>
<td>Pao2 (kPa)</td>
<td>7.6 (0.9)</td>
</tr>
</tbody>
</table>

PAV is delivered by the ventilator according to the equation of motion generating a pressure in relation to the spontaneous effort of the patient (Pmus):

\[
P_{mus} = E \times V + R \times V' \quad \text{[Equation 1]}
\]

where E and R are the elastance and resistance, respectively. A portion of the total mechanical workload (that is, the elastance and resistance) is taken over according to the level of assistance which has been decided by the caregiver and can specifically unload the resistive burden (flow assist, FA) and the elastic burden (volume assist, VA). Hence equation 1 becomes:

\[
P_{mus} = \text{PAV} = (E - \text{VA}) \times V + (R - \text{FA}) \times V' \quad \text{[Equation 2]}
\]

To set PAV we followed the procedure described in our previous study. Briefly, we started with VA and FA set at the minimum value of 2 cm H2O/l and 1 cm H2O/l.s, respectively, and progressively increased the level of assistance until the patient felt uncomfortable. We then applied the last level of VA and FA at which the patient felt comfortable.
slightly longer than TI, Edi. In view of the normal distribution of VT values we used the coefficient of variation to assess VT under all conditions. During spontaneous breathing TI, Edi, matic frequencies were essentially the same in all patients the flow and on the Edi records. The ventilatory and diaphragm suggesting an increase in VT without a substantial reduc-

Mean values and standard deviations of peak airway 

Figure 1

Table 2 Breathing pattern and transcutaneous blood gas tensions with spontaneous breathing, PSV and PAV

<table>
<thead>
<tr>
<th></th>
<th>SB</th>
<th>PSV</th>
<th>SB</th>
<th>PAV</th>
<th>PSV – PAV</th>
</tr>
</thead>
<tbody>
<tr>
<td>VT (l/m)</td>
<td>10.5 (2.3)</td>
<td>13.9 (8.8)*</td>
<td>10.1 (2.7)</td>
<td>12.6 (4.0)*</td>
<td>+1.34 (1.51)</td>
</tr>
<tr>
<td>Vt (l)</td>
<td>0.5 (0.1)</td>
<td>0.7 (0.2)*</td>
<td>0.5 (0.2)</td>
<td>0.7 (0.2)*</td>
<td>+0.2 (0.2)</td>
</tr>
<tr>
<td>Ti (s)</td>
<td>1.1 (0.3)</td>
<td>1.1 (0.2)</td>
<td>1.1 (0.2)</td>
<td>1.1 (0.3)</td>
<td>+0.0 (0.2)</td>
</tr>
<tr>
<td>Te (s)</td>
<td>1.9 (0.6)</td>
<td>2.0 (0.7)</td>
<td>2.0 (0.6)</td>
<td>2.3 (0.9)</td>
<td>–0.2 (0.7)</td>
</tr>
<tr>
<td>Ttot (s)</td>
<td>3.1 (0.9)</td>
<td>3.1 (0.9)</td>
<td>3.2 (0.8)</td>
<td>3.4 (1.1)</td>
<td>+0.2 (0.7)</td>
</tr>
<tr>
<td>IC (l)</td>
<td>1.2 (0.3)</td>
<td>1.1 (0.3)</td>
<td>1.2 (0.3)</td>
<td>1.2 (0.3)</td>
<td>–0.1 (0.2)</td>
</tr>
<tr>
<td>Tco2 (mm Hg)</td>
<td>52 (6.1)</td>
<td>48 (6.2)*</td>
<td>52 (6.8)</td>
<td>49 (5.1)*</td>
<td>–1 (2.8)</td>
</tr>
<tr>
<td>Pao2 (mm Hg)</td>
<td>62 (7.8)</td>
<td>63 (8.0)</td>
<td>61 (8.6)</td>
<td>65 (7.4)</td>
<td>–1 (6.7)</td>
</tr>
<tr>
<td>Slo2 (%)</td>
<td>90 (3.5)</td>
<td>93 (1.8)*</td>
<td>91 (2.4)</td>
<td>92 (2.4)</td>
<td>1 (2.1)</td>
</tr>
</tbody>
</table>

Values are mean (SD). SB = spontaneous breathing; PSV = pressure support ventilation; PAV = proportional assist ventilation; VT = tidal volume; Ti = inspiratory time; Te = expiratory time; Ttot = total cycle duration; IC = inspiratory capacity; Tco2 = transcutaneous CO2, Pao2 = transcutaneous O2, Slo2 = transcutaneous oxygen saturation; PSV = PAV = difference between PSV and PAV.

did not change. Pulse oximetry slightly increased with PSV. No significant difference was found between PAV and PSV (table 2).

Both Edi,peak and Edi,int were significantly reduced by PSV (–32 (14)% and –44 (21)% and by PAV (–20 (17)% and –34 (25)). As shown in fig 1, Pao, TI, and over total cycle duration (Pao, Ttot) during pressure support ventilation (PSV) and proportional assist ventilation (PAV).

At the end of each step the dyspnea VAS score was measured. During spontaneous breathing the mean score was 17 (5) cm which was reduced slightly, but not significantly, to 15 (5) cm at the end of both PSV and PAV. Three patients reported that they felt better with PSV than with PAV; whereas four other patients reported the opposite effect. Five patients had no preference.

**DISCUSSION**

The results of this study show that short term non-invasive application of PAV and PSV in patients with CVF due to advanced CF can improve the patients’ pathophysiological condition compared with spontaneous breathing. Both PAV and PSV resulted in a higher Vt and lower Tco2, and a smaller inspiratory effort. The patients accepted both modes of mechanical ventilation and claimed that they felt better with ventilatory assistance than with spontaneous unsupported breathing as shown by a reduction in the dyspnea score. The lack of statistical significance was perhaps due to the short ventilation time. However, there was no systematic preference for one mode over the other. A significant subjective improvement in patients with CF receiving NPPV has been reported previously. This reduction in symptoms, which may help the general well being of the patients, may explain why NPPV is well tolerated even in the long term.

NPPV reduces the progressive deterioration of gas exchange and provides support during exacerbations while the patients are waiting for lung transplantation.4,5 PSV improved Vt, Slo2, and reduced Tco2.4 In patients with severe CF who have significant gas exchange abnormalities during sleep but are normocapnic in the daytime, nocturnal (one night) nasal PSV was able to prevent oxygen induced hypercapnia.4 The short term benefits were confirmed in a longer study6 in which home NPPV improved physiological variables and quality of life up to 18 months after initiation of the treatment. Hodson and colleagues emphasised the role of NPPV in patients with CF, defining it as “a potential bridge to transplantation”.1 NPPV has also been proposed in several

**Figure 1** Mean values and standard deviations of peak airway pressure (Pao, peak), mean airway pressure over inspiratory time (Pao, Ti), and over total cycle duration (Pao, Ttot) during pressure support ventilation (PSV) and proportional assist ventilation (PAV). *p<0.05 versus PSV.
centres as a first line intervention in patients with CF who require ventilatory support before transplantation.

Our study provides the first physiological assessment of PAV in patients with CF, as well as the first physiological comparison between PAV and PSV administered non-invasively in patients with CF. The two modes of ventilatory assistance had similar results. However, the physiological benefits of PAV occurred at a lower mean airway pressure than PSV (fig 1). This may be of clinical interest in view of the results obtained by Diaz and colleagues who showed that PSV (mean 12 cm H2O) caused a significant fall in cardiac output (mean of –1 l/min) which they attributed to the effect of airway pressure on venous return, and by Haworth and colleagues who reported three cases of barotrauma in adult patients with CF dependent on NPPV. The authors commented that this risk was not different from the general population of patients with CF. Clearly, it is important to use the lowest possible airway pressure during NPPV to prevent both the risk of barotrauma and the fall in cardiac output. The latter may be relevant in patients whose respiratory muscles are contracting under a significant workload. It is interesting to note that the improvement in VT and the reduction in the diaphragmatic effort observed with PAV in patients with CF is similar to that obtained by PAV in patients with COPD.

In agreement with other studies, our data show that, on average, NPPV increases ventilation and unloads the respiratory muscles. However, when individual patients were analysed (fig 2) we found that the mean changes may not reflect individual behaviour. Four different patterns were observed when ventilatory assistance was offered. At the two extremes the increase in VT (fig 2A) and the reduction in the inspiratory effort (fig 2B) were not associated. They both occurred in some patients (fig 2C) and in a few the reaction was different for different ventilatory modes (fig 2D). In some patients VT was increased, in others the respiratory muscles were unloaded, while in some there was a combination of the two. We did not find any criteria to predict the individual response to the ventilatory assistance and we do not know whether this reflects differences in the central control of breathing, particularly PAV, a ventilatory mode which is driven...
by the patient. Comparison between modes of ventilatory assistance always presents problems and there is no perfect solution. In this study we decided to set both modes at a level of comfort determined by the patients because their cooperation is crucial for the success of NPPV and because it is the usual setting for clinical purposes. As far as we are aware, PAV and PSV have only been compared to date in intubated patients and this is the first comparison of the two modes of ventilation during NPPV.

In conclusion, the results of this study show that short term application of NPPV in patients with CF with chronic hypercapnia, both with PSV and PAV set at a level of comfort determined by the patient, has a positive physiological effect on minute ventilation, blood gas tensions, and the amount of diaphragmatic effort. However, PAV gave similar results to PSV at a lower mean inspiratory pressure.

ACKNOWLEDGMENTS

Dr A Serra is a fellow of the Italian Ministry of University and Scientific Research. We thank Dr Fiona Scandellari for language and editorial assistance. The work has been supported by grants from Respironics Inc, Murrysville, PA, USA and from Fondazione Ricerca Fibrosi Cistica.

Authors’ affiliations

A Serra, C Braggion, Centro Regionale Fibrosi Cistica, Azienda Ospedaliera di Verona, Ospedale Civile Maggiore di Borgo Trento, Verona, Italy
G Polese, A Rossi, Unità Operativa Pneumologia, Ospedali Riuniti di Bergamo, Bergamo, Italy

REFERENCES

Non-invasive proportional assist and pressure support ventilation in patients with cystic fibrosis and chronic respiratory failure
A Serra, G Polese, C Braggion and A Rossi

Thorax 2002 57: 50-54
doi: 10.1136/thorax.57.1.50

Updated information and services can be found at:
http://thorax.bmj.com/content/57/1/50

These include:

References
This article cites 20 articles, 6 of which you can access for free at:
http://thorax.bmj.com/content/57/1/50#BIBL

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Topic Collections
Articles on similar topics can be found in the following collections
Cystic fibrosis (525)
Airway biology (1100)

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/