Required levels of nasal continuous positive airway pressure during treatment of obstructive sleep apnoea

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ABSTRACT: The improvement in the severity of obstructive sleep-related breathing disorders during nasal continuous positive airway pressure (NCPAP) therapy can account for the decrease in the required NCPAP level with time. The aim of this study was to prospectively quantify the changes in the required NCPAP level over time of use in sleep apnoea-hypopnoea syndrome (SAHS).

Forty sleep apnoea-hypopnoea patients were evaluated before and during the time course of NCPAP therapy. The effective NCPAP level was defined as the positive pressure level that abolished apnoeic and hypopnoeic events and snoring in all sleep stages and sleep positions. This pressure level was determined within 2 weeks after baseline diagnostic sleep study. Sleep studies with NCPAP and NCPAP titration were performed after 2 (n=40), 8 (n=40), and 20 (n=24) months of NCPAP therapy.

The initial effective NCPAP level was 9.6±0.4 cmH₂O. It progressively decreased to 8.8±0.4, 7.9±0.4 and 7.7±0.5 after 2, 8 and 20 months, respectively; the difference being significant between the first three NCPAP nights. There was a weak negative relationship between the changes in the effective NCPAP and changes in weight recorded at the different visits. There was a weak negative relationship between the changes in NCPAP and the previous NCPAP level. In 13 patients, the apnoea-hypopnoea index (AHI) remained >10 n·h⁻¹ at the first NCPAP trial because the effective NCPAP level was not tolerated. Despite a suboptimal NCPAP level, their sleep architecture improved, and they all reported a subjective improvement in diurnal hypersomnolence. After 2 months of NCPAP therapy, the AHI was <10 n·h⁻¹ in 11 of these suboptimally treated patients.

We conclude that the required NCPAP level progressively decreases with use. The changes in body weight may play a minor role in the change in the effective NCPAP level.

index decreases with NCPAP but who cannot tolerate the optimal NCPAP level. If the improvement of nocturnal breathing disorders induced by NCPAP is of clinical significance, NCPAP use may be accompanied by a progressive decrease in the required level of NCPAP.

The aims of this study were to prospectively determine the changes in the required NCPAP level in SAHS over time of use, and to evaluate the changes in persisting sleep-related breathing disorders with the use of suboptimal NCPAP therapy.

Methods

Subjects

Forty five SAHS patients were included in the study. Each patient was evaluated by one of the authors and had conventional polysomnography for clinical suspicion of SAHS. None had undergone previous surgical treatment for sleep-related breathing disorders and none was taking antidepressants, hypnotics or sedatives. The different treatment alternatives were explained to the patients. Those who chose NCPAP therapy after the first sleep study with NCPAP were asked to participate in this study, and all agreed to do so. This first NCPAP trial was performed during a complete sleep recording within 2 weeks after the baseline sleep study, with determination of the effective NCPAP level (cf. infra). Patients were instructed on NCPAP use and installation principles by the sleep technician. The complete instructions on the use of NCPAP machines and nasal mask installation were given at home by the home care company representative. They were treated with the same type of NCPAP apparatus that was used in the sleep laboratory, adjusted at the effective pressure level that was determined during the first NCPAP sleep trial. For a patient to be included in the study, NCPAP had to be applied at least 4 h·night⁻¹ for ≥5 nights-week⁻¹, as estimated by regular visits to the patients by a home care company and the records from the meters of the NCAP machines. Five patients who were initially enrolled in the study were eliminated due to failure to respect these criteria.

Determination of the NCPAP effective level

Determination of the NCPAP effective level was achieved during full night polysomnographic studies, that included the determination of sleep stages (electroen-cephalogram C₃A₁, C₄A₂; electro-oculogram; submental electromyogram), nasal and mouth airflow with thermocouples (Grass Instruments, Quincy, MA, USA), arterial oxygen saturation (SaO₂) with a Criticare 504 ear oximeter (CSI, Waukesha, WI, USA), electrocardiogram and thoracoabdominal movements by respiratory inductive plethysmography (Respitrace, Ambulatory Monitoring, Ardsley, NY, USA) calibrated by the isovolume method [14]. The snoring sound pressure level (SPL) was displayed after the signal of two microphones (Shure SM 58, Evanston, IL, USA) had been preamplified, mixed, equalized (DOD R 831, Salt Lake City, Utah, USA) and analysed with a spectrum analyser (RTA SA 3050, Audio Control, Mountlake Terrace, WA, USA) [15]. All parameters were recorded on a 16-channel polygraph (Model 78; Grass Instruments, Quincy, MA, USA) at a paper speed of 10 mm·s⁻¹.

Positive pressure level was progressively increased in 1 cmH₂O steps, from 3 cmH₂O until apnoeic and hypopnoeic events, as measured by the apnoea-hypopnoea index (AHI), and snoring were abolished in all sleep stages and all sleep positions, or at the maximal pressure level that was tolerated by the subjects. Polysomnographic recordings were analysed manually in 30 s epochs, and sleep stages, arousals and respiratory events were defined by standard criteria [16–18]. Snoring was defined as a spike in breathing sound intensity greater than 60 dB SPL. Apnoea, hypopnoea, arousal and snoring indices represent the frequency of these events per hour (n·h⁻¹) of sleep.

Protocol

Sleep studies with NCPAP and NCPAP titration were performed after 2 (n=40), 8 (n=40), and 20 (n=24) months of NCPAP therapy. The number of patients had decreased at the last visit because some of them had not reached the 20 months visit. Subjects were asked to contact the sleep laboratory if they had any trouble with the apparatus. Subjects were asked to avoid consuming alcohol for a minimum of 12 h preceding each recording session.

Statistical analysis

Since our data were normally distributed, parametric tests were used. Data expressed as percentages were transformed to obtain an underlying normal distribution [19]. Since our patients were evaluated at different time intervals, the assumption of an equal correlation between the results obtained at the different visits was verified with a sphericity test on orthogonal components [20]. Afterwards, we performed a complete randomized block design, followed by Tukey’s test for multiple comparison for each variable. The influence of weight changes and NCPAP values on the changes in the effective NCPAP level at the different visits were analysed by the least square method. Differences were considered significant at a value of p<0.05.

Results

We analysed the results of forty subjects (34 males and 6 females; age range 35–63 yrs; body mass index 36.3±1.2 kg·m⁻²; mean±SEM). The results of the sleep studies were typical of SAHS, with sleep fragmentation and disorganization and nocturnal desaturations (table 1).
As expected, these abnormalities improved with NCPAP (Table 1). The effective NCPAP level ranged 5–15 cmH2O. Interestingly, the percentage of total sleep time (TST) spent in the supine position increased at the first NCPAP trial and remained elevated at the remaining visits. In each patient, diurnal hypersomnolence and alertness subjectively improved with NCPAP.

The effective NCPAP level progressively decreased with time (first NCPAP trial: 9.6±0.4 cmH2O; after 2 months of NCPAP 8.8±0.4 cmH2O; after 8 months of NCPAP 7.9±0.4 cmH2O; and after 20 months of NCPAP 7.7±0.5 cmH2O), the difference being significant between the first three visits, with no further decrease after 20 months of NCPAP therapy (Fig. 1). For the whole group, there was no significant change in weight between the different visits. There was a significant relationship between changes in weight and in the effective NCPAP level (r=0.37; p<0.0001) (Fig. 2). There was a tendency for the decrease in NCPAP to be greater when the initial NCPAP was higher, but the relationship between these two variables was weak (r=0.32; p=0.002) (Fig. 3).

Results of the sleep studies realized at the different visits. Data are presented as mean±SEM. $: within 2 weeks of baseline prior to home NCPAP. TST: total sleep time; AHI: apnoea + hypopnoea index; REM: rapid eye movement sleep. Data obtained at the different visits were compared by complete randomized block design, followed by Tukey's test for multiple comparison for each variable. Values superscripted by different letters are significantly different from one another (mean±SEM).

### Table 1. Results of sleep studies with nasal continuous positive airway pressure (NCPAP)

<table>
<thead>
<tr>
<th></th>
<th>Baseline (n=40)</th>
<th>First$ (n=40)</th>
<th>2 months (n=40)</th>
<th>8 months (n=40)</th>
<th>20 months (n=24)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight kg</td>
<td>100.6±2.8</td>
<td>100.7±2.8</td>
<td>100.6±2.8</td>
<td>100.9±2.8</td>
<td>101.9±2.8</td>
</tr>
<tr>
<td>TST h</td>
<td>6.1±0.2</td>
<td>5.9±0.2</td>
<td>5.8±0.2</td>
<td>6.2±0.1</td>
<td>5.8±0.1</td>
</tr>
<tr>
<td>Stage I-II % TST</td>
<td>70.7±1.9a</td>
<td>48.7±2.0b</td>
<td>61.0±1.4c</td>
<td>60.7±0.9c</td>
<td>61.4±1.4c</td>
</tr>
<tr>
<td>Stage III-IV % TST</td>
<td>10.6±1.0b</td>
<td>26.1±1.4b</td>
<td>17.9±0.7c</td>
<td>16.3±0.7c</td>
<td>17.6±1.0c</td>
</tr>
<tr>
<td>Stage REM % TST</td>
<td>19.8±1.9a</td>
<td>25.1±1.6b</td>
<td>21.0±1.1ab</td>
<td>22.8±0.9b</td>
<td>20.9±1.0b</td>
</tr>
<tr>
<td>Arousal index O·h⁻¹</td>
<td>46.9±4.7a</td>
<td>11.2±1.1b</td>
<td>9.0±0.8b</td>
<td>8.5±0.6b</td>
<td>9.3±1.1b</td>
</tr>
<tr>
<td>AH1 n·h⁻¹</td>
<td>48.4±4.7a</td>
<td>9.1±1.4b</td>
<td>3.8±0.6b</td>
<td>3.6±0.6b</td>
<td>4.9±0.8b</td>
</tr>
<tr>
<td>% TST &lt;80% Sao₂</td>
<td>18.9±4.4a</td>
<td>2.5±1.3b</td>
<td>0.9±0.8b</td>
<td>0.1±0.05b</td>
<td>0.1±0.06b</td>
</tr>
<tr>
<td>% TST supine</td>
<td>32.7±4.0b</td>
<td>46.0±4.4b</td>
<td>42.1±3.9ab</td>
<td>39.7±3.8ab</td>
<td>40.7±5.9ab</td>
</tr>
</tbody>
</table>

As expected, these abnormalities improved with NCPAP (Table 1). The effective NCPAP level ranged 5–15 cmH2O. Interestingly, the percentage of total sleep time (TST) spent in the supine position increased at the first NCPAP trial and remained elevated at the remaining visits. In each patient, diurnal hypersomnolence and alertness subjectively improved with NCPAP.

The effective NCPAP level progressively decreased with time (first NCPAP trial: 9.6±0.4 cmH2O; after 2 months of NCPAP 8.8±0.4 cmH2O; after 8 months of NCPAP 7.9±0.4 cmH2O; and after 20 months of NCPAP 7.7±0.5 cmH2O), the difference being significant between the first three visits, with no further decrease after 20 months of NCPAP therapy (Fig. 1). For the whole group, there was no significant change in weight between the different visits. There was a significant relationship between changes in weight and in the effective NCPAP level (r=0.37; p<0.0001).

**Fig. 1.** Means±SEM values of the effective nasal continuous positive airway pressure (NCPAP) level measured at the different visits. This pressure level progressively decreased with time, the difference being significant after 2 and 8 months of NCPAP therapy. Values with different letters are significantly different from one another. Means±SEM. Note that the vertical axis is magnified and cut-off from zero.

**Fig. 2.** Relationship between weight changes (differences in % of baseline values) and changes in the effective nasal continuous positive airway pressure (NCPAP) level for the three NCPAP visits. There was a poor but significant relationship between these two variables (r=0.37; p<0.0001).

**Fig. 3.** Relationship between changes in the nasal continuous positive airway pressure (NCPAP) level (difference in pressure between two successive visits) and the previous visit NCPAP level for the first two NCPAP trials (2 months– 8 months). The higher the pressure level, the more it could be decreased with time. The relationship between these variables was poor (r=0.32; p=0.002).
In 13 patients, AHI remained above 10 n·h⁻¹ at the first NCPAP trial (19.4±2.1 n·h⁻¹; range 12–32 n·h⁻¹) because the effective NCPAP level was not tolerated. The range of NCPAP level applied that was tolerated ranged 7–15 cmH₂O (11.5±0.7 cmH₂O). Body mass index and baseline apnoea-hypopnoea index were among the highest values of the group (40.7±2.7 kg·m⁻² and 63.9±7.8 n·h⁻¹, respectively). However, despite a suboptimal nasal continuous positive airway pressure (NCPAP) level during the first NCPAP trial. In 11 of these patients, the sleep-related breathing disorders returned to the normal range (<10 n·h⁻¹) after 2 months of NCPAP. The index values are given on a logarithmic scale for clarity.

Discussion

This study demonstrates that the required NCPAP level progressively decreases during the first 8 months of use, and that the changes in body weight poorly contribute to the changes in the effective NCPAP level.

Recent reports have shown that NCPAP benefits may be prolonged after its interruption following long-term treatment [8, 9]. The improvement reported by RAUSCHER et al. [8] was more important in patients with the most severe disorders. Similarly, the prolonged beneficial effects observed by LEECH et al. [9] after 6–46 months of NCPAP therapy occurred in patients with a respiratory disturbance index greater than 50 n·h⁻¹. SCHWARTZ et al. [21] reported a small but significant decrease in the closing pressure after NCPAP therapy, and a positive relationship between upper airway collapsibility and the severity of sleep-related breathing abnormalities [21, 22]. We found that the decrease in the NCPAP effective level tended to depend on the previous pressure level, and that the decrease in the effective positive pressure plateaued after 8 months of NCPAP therapy. Therefore, it can be assumed that the amount of NCPAP-induced decrease in upper airways collapsibility depends on its baseline value until the mean effective pressure has been reached, with no further improvement over time.

In previous studies, the delayed effects on NCPAP could not be explained by changes in body weight [8, 9]. Changes in body weight could account for only 13% of the changes in the effective NCPAP level that we observed, and the decrease in the pressure level occurred even when body weight remained unchanged. However, the influence of the changes in body weight on the individual decrease in pressure level may be explained by the importance of this parameter on upper airway collapsibility [21].

Recent reports suggest that the effective compliance to NCPAP is overestimated by questionnaires and that less than 50% were regular users [23]. Therefore, one could question the compliance to the treatment, and whether our results are effectively the consequence of regular NCPAP use. All our patients complained of daytime sleepiness, difficulty in concentration, and automatic behaviour that interfered with their professional activities. All these symptoms subjectively improved with NCPAP. Since these factors are good predictors of regular use [23], we are confident that NCPAP was regularly employed and that our results are the consequence of its regular use as assessed by the records of the meters of the machines. Even if NCPAP use was not objectively measured in all our subjects, it is accepted that its use is associated with a correction of haemodynamic [3], ventilatory [9, 24] and neurophysiological [25] abnormalities, and with changes in upper airway structures [10], and collapsibility [21].

The main mechanisms accountable for the delayed effect of NCPAP are: an improvement in upper airway morphology, a correction of sleep fragmentation, and changes in ventilatory control. An increase in the minimal pharyngeal cross-sectional area is observed after 4–6 weeks of NCPAP that is correlated with the decrease in the mucosal water content [10]. This reduction in upper airway oedema could increase upper airway patency and decrease its collapsibility [26, 27]. Reversible local tissue damages could occur as a consequence of recurrent episodes of upper airway closure and the development of high levels of subatmospheric pressure. These two factors would be suppressed with correction of sleep-related breathing disorders, accounting for a decrease in upper airway collapsibility and of the required NCPAP.
level with time. Improvement in sleep fragmentation by NCPAP could also account for the delayed effects of NCPAP. Since the electromyographic activity of the genioglossus is depressed after sleep deprivation [28], the elimination of sleep fragmentation by NCPAP can be accounted for by an improvement in the neuromuscular control of the pharynx. Improvements in diurnal hypoxaemia and hypercapnia occur after long-term treatment with NCPAP [9, 13]. Changes in ventilatory control responsiveness are observed within a few days of an efficient treatment of apnoeic events [24, 29]. Since upper airway patency and central ventilatory activity are intimately linked [30], it is possible that changes in the sensitivity of the ventilatory control are involved in the delayed effects of NCPAP. However, since only six of our patients had diurnal hypoxaemia and hypercapnia, we do not believe that this mechanism can account for the findings in all of our study population; five of the apnoeic patients with an obesity hypoventilation syndrome had normal diurnal arterial blood gases after 2 months of NCPAP therapy.

The progressive improvement in nocturnal breathing disorders during NCPAP therapy can have important practical repercussions. The first practical consideration concerns the required frequency of control sleep studies in SAHS treated with NCPAP. Our results suggest that a 2–6 month control, and subsequent annual visits are needed to adjust the pressure level. In the experience of ourselves and others, nasal problems such as dryness and congestion are the most frequent complaints during NCPAP therapy [31]. Since these side-effects are related to nasal airflow, and indirectly to NCPAP level, the decrease in the effective pressure level may contribute to improved comfort. The second practical consideration deals with "unsuccessful" first NCPAP trials. We found that in 13 subjects, obstructive sleep-induced respiratory abnormalities persisted during the first NCPAP trial because the effective NCPAP level was not tolerated. Since all these patients reported severe daytime hypersomnolence that improved with NCPAP, they were kept on this treatment. The results obtained at the follow-up visits clearly demonstrate that the majority of these patients improved further after 2 months of NCPAP therapy. Therefore, since the first goal of the initial NCPAP therapy is to improve the consequences of sleep-related obstructive breathing disorders, the persistence of apnoeic and hypopnoeic events does not rule out NCPAP efficiency if the frequency of breathing disorders, nocturnal desaturations, and sleep disruption are improved. In such cases, home therapy with NCPAP can be initiated, recognizing that polysomnographic study should be performed within the following months to document delayed improvement and eventually reduce the pressure level. Other therapeutic methods (e.g., bilevel positive airway pressure (BiPAP)) could be used after this delay, if NCPAP failed to normalize sleep-related breathing disorders at follow-up visits.

We conclude that the effective NCPAP level decreases with time and that patients using suboptimal NCPAP levels may have immediate benefits of this therapy and further improvement over time.

References