Clinical and polysomnographic assessment of the BRD Appliance in the treatment of Obstructive Sleep Apnea Syndrome

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Abstract

Objectives: This research was designed to perform a clinical and polysomnographic evaluation of the effect of an oral appliance (OA) for the treatment of Obstructive Sleep Apnea Syndrome (OSAS), developed and tested by two Brazilian federal universities. Methods: The sample consisted of 50 patients (aged between 18 and 65 years, 33 men and 17 women) with initial polysomnographic diagnoses of mild to moderate OSAS. All patients underwent a new polysomnographic assessment during an entire night (using the OA) approximately six months after the first evaluation. Based on the reduction of apnea and hypopnea index (AHI) obtained with the use of the OA, the patients were divided into good responders (reduction of 50% or more of the AHI, remaining below 10 events/hour) and poor responders (AHI remained greater than or equal to 10 events/hour). Results and Conclusions: In 54% of the sample, the AHI decreased to less than five events/hour with the use of the OA; in 38% the reduction in AHI was greater than 50% but remained above the five events/hour; and in 6% of the sample the AHI decreased less than 50%. The good responders accounted for 86% of the studied sample, while the other 14% were poor responders. There were significant improvements in the sleepiness scale, the AHI, the micro-arousals and the minimum oxyhemoglobin saturation with the utilized therapy. A high Body Mass Index seemed to negatively interfere in the performance of the device under study.

Keywords: Obstructive Sleep Apnea. Polysomnography. Respiratory protection devices. Snoring.

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INTRODUCTION AND LITERATURE REVIEW

Obstructive Sleep Apnea Syndrome (OSAS) is a respiratory disorder related to sleep characterized by repeated partial or complete obstructions of the upper airway during sleep.\(^3,4\) It is one of the most common clinical entities among sleep disorders, affecting 3 to 5% of the adult population.\(^30\)

Among the classical clinical findings are heavy snoring, intermittent respiratory pauses during sleep, breathless and recurrent awakenings, non-restorative (fragmented) sleep and excessive daytime sleepiness.\(^3,5\) Snoring broken by respiratory pauses is a typical account of the companions of these patients.\(^5,22\) However, snoring can occur in the absence of OSAS, and as such, it is characterized as primary snoring, which affects about 40% of the adult male population.\(^5\)

Precise diagnosis of sleep-disordered breathing is given by means of polysomnography exam (PSG), which permits the quantification of the events reported above and the temporal relationship of the parameters measured during a full night of sleep. The quantification of respiratory events per hour of sleep, given by the apnea-hypopnea index (AHI), confirms the diagnosis and defines the severity of OSAS: mild = between 5 and 15; moderate = between 15 and 30; and severe = above 30 events.\(^3\) There are still other parameters that can be altered in these patients and viewed in the polysomnography, such as oxyhemoglobin desaturation, alteration in the percentages of sleep stages, reduced sleep efficiency and sleep fragmentation.\(^3,4,7\)

The pathophysiology of OSAS appears to be multifactorial, and there are anatomical, functional and neuromuscular factors involved. The soft tissues, adipose tissue, muscles and craniofacial skeleton directly affect the configuration and dimension of the pharynx. In this manner, we frequently observe patients with OSAS with lingual hypotonia, macroglossia, mandibular and/or maxillary retrognathia, micrognathia, ogival palate, transverse maxillary deficiency and crossbite.\(^10,18,22,25,27\)

OSAS is commonly associated with cognitive and cardiovascular complications as a result of intermittent hypoxia and sleep fragmentation, which significantly increases the morbidity and mortality of this syndrome.\(^21\) Recent studies have established that OSAS is an independent risk factor for systemic arterial hypertension (SAH) and contributes to the onset and progression of other cardiovascular diseases.\(^26\) The cognitive consequences are principally related to attention, memory and executive function.\(^1,13\)

Because this syndrome is a chronic and long lasting condition, an effective approach to treatment is mandatory.\(^15\) OSAS treatments consist of clinical and surgical procedures. The choice of the clinical modality of treatment, using positive pressure devices (CPAP, auto-adjusting CPAP, BiPAP) or oral appliances (OA), is directly related to the severity of the disease.

The nasal CPAP (“continuous positive airway pressure”) device is considered the gold standard in the treatment of OSAS\(^14\) (Fig 1). It consists of a non-invasive method of continuous positive air pressure application in the airway, generating a continuous airflow that, through a flexible tube, reaches a nasal mask that is adjusted on the face with straps. Thus, a pneumatic cushion is created.
in its interior that tends to dislocate the soft palate in the direction of the lingual base and dilate the sectional area of the whole pharynx. While it is an extremely effective treatment, there are problems with the adhesion of positive pressure devices and their acceptance in the long term.

Oral Appliances (OA) are devices used in the oral cavity during sleep, with the objective to prevent the collapse between the tissues of the oropharynx and the lingual base, i.e., the obstruction of the upper airway. OA constitute an effective and well-accepted form of treatment, and they have been an increasing line of treatment of OSAS and snoring for more than 20 years. Today there are more than 80 described types of OA, which specifically fit into the categories of lingual retainers and mandibular repositioners. Only some have been approved by the FDA (Food and Drugs Administration), and there are few controlled studies available. There are different types of OA, differentiated by manufacturing method (prefabricated or made in a laboratory), retention, titration of the mandibular position, anterior vertical opening, freedom of mandibular movement and construction material, among others. The effectiveness of the OA seems to be related to some of these aspects. When the precautions related to these factors are not observed during the planning of the device, there is a greater probability of side effects, including those related to temporomandibular joint (TMJ) and occlusal alterations, as well as a reduction in adherence to the treatment.

Treatment with OA is prescribed in patients with primary snoring and mild OSAS, as well as in cases of moderate or severe OSAS in which there was intolerance or refusal of CPAP use, surgical contraindication or the need for a short-term substitute therapy. The objective of the present study was to evaluate the performance of this new design of OA for OSAS, while attempting also to compare responding and non-responding patients with the use of this mode of treatment.

THE INTRA-ORAL DEVICE

The Brazilian Dental Appliance (BRD) was developed from the experience of orthodontics researchers in the area of sleep medicine at UNIFESP-EPM and UFC who had previously used numerous other devices, mainly North American and Canadian, many of which had important limitations, high costs or a lack of consistent studies endorsing their use in patients with OSAS.

The BRD is an adjustable mandibular repositioning OA (Fig 2, A and B). It is comprised
of retaining clamps on the posterior teeth and two acrylic support bases (one superior and another inferior) covering all teeth externally and internally (anterior and posterior). It also possesses two independent (one right and another left in the posterior palate region of the superior acrylic base) expansion mechanisms (screws located with their long axle in the anterior-posterior direction). Two independent palate rods (one right and another left) protrude from these expansion mechanisms and are inserted inferiorly into two small tubes located in the anterior portion (distal of the inferior canines) of the inferior acrylic support base. This proposed design allows successive advances in the mandibular position without, however, impeding lateral mandibular movements. That is, with the device in position, even when the jaw is in an anterior position, the patient can make lateral movements and a small buccal opening (Fig 3).

**METHODOLOGY**

This prospective study included 50 patients with OSAS prescribed for treatment with an OA, following the study design (Fig 4). All the patients had been previously diagnosed by a standardized polysomnography examination in the Department of Psychobiology at UNIFESP (Universidade Federal de São Paulo) and the Universidade Federal do Ceará.

The inclusion criteria consisted of patients with mild and moderate OSAS, aged between 25 and 70 years, with body mass index (BMI) values less than 30 Kg/m² and with at least 10 teeth in each dental arch. Excluded patients were comprised of those who had undergone previous surgeries for OSAS, CPAP users, those currently using any drug that could interfere with the sleep architecture, shift workers and patients with important anatomical alterations in the superior airway, such as deviated septum, turbinate and/or tonsil hypertrophy.

Volunteers who were diagnosed with extensive periodontal disease, severe temporomandibular joint disorder or dental units with insufficient anatomy for the necessary OA retention were excluded from the study.
Odontological documentation (DOC)

All of the patients underwent a basic orthodontic documentation, consisting of panoramic x-ray, cephalometric analysis, intra and extra-buccal photographs and study models.

Polysomnography (PSG)

All of the patients were subjected to a baseline polysomnography (PSG) for OSAS diagnosis and another one with the device (BRD) in the mandibular position of maximum comfortable protrusion. The average time between the first and the second polysomnographies was six months.

The full night of PSG was performed in a sleep laboratory using a system of 13 channels, which included the electroencephalogram, electrocardiogram, electrooculogram, electromyogram of the submentonian muscles and tibialis anterior, nasal and oral airflow (measured by thermistor and/or nasal cannula), thoracic and abdominal movements, body position (supine) and oxyhemoglobin saturation measured by pulse oximetry.

Clinical and laboratory odontologic procedures

Impressions of the dental arches were made with alginate type II and immediately cast with special plaster type IV. The record of the initial mandibular position was obtained with the “George Gauge” device and condensation silicone (dense). The plaster models and records were then used for the manufacture of an individualized BRD device. All devices in the study were made of thermopolymerizable acrylic, with independent upper and lower parts and retention obtained by interproximal clips placed between all the posterior teeth. The connection between the two arches was made by two expanders positioned at the palate region, just below the cervical region of the superior molars. The purpose of these expanders is to allow the advancement of the mandible to move gradually over 11 mm, in 44 increments of 0.25 mm (Fig 5).

To obtain the mandibular position of maximum comfortable protrusion, progressive adjustments were made in the expanders of the devices during a period of three to four 4 months, according to the presence of snoring and additional symptoms (drowsiness, fatigue, fragmented or non-restorative sleep).

A limit of 70% of maximum mandibular protrusion was observed. At each consultation, the patients were asked about episodes of nasal obstruction and weight change.

Statistical Analysis

The Kolmogorov-Smirnov test for normality was applied and was followed by the paired t-test for comparison between the baseline and OA, in cases when the baseline data displayed a normal distribution, or by the Wilcoxon test when the distribution was not normal.

Subsequently, the patients were divided into two groups: good and poor responders. A patient was considered a good responder when the AHI with the OA was reduced more than 50% from the baseline and was lower than 10 events/hour. The poor responder presented an AHI with the OA of greater than or equal to 10. For this situation, to compare these two groups, we used the Student t-test for independent samples if the distribution was normal or the Mann-Whitney test for non-normal distributions.
RESULTS

The data are presented as means and standard deviations. The sample was comprised of 33 men and 17 women with a mean age of 48.6 ± 12.3 and a mean BMI of 26.0 ± 2.8 Kg/m². The mean of the final protruding mandibular position was 8.8 ± 1.3 mm.

The results of the comparison between the baseline (initial) and the with-OA measurements for all patients are shown in Table 1 and, specifically in relation to the number of apneas and hypopneas per hour of sleep (AHI), are illustrated in Graph 1. Statistically significant improvements were noted in several of the parameters studied, like the reduction in sleepiness measured by the Epworth scale, increased percentage of REM sleep, decrease in AHI, increased minimum oxyhemoglobin saturation as well as the decrease in the number of micro-arousals per hour of sleep (Table 1).

In 28 patients (56% of the sample), the AHI normalized with the use of the OA, i.e., it was less than 5 events/hour (success); in 19 patients (38% of the sample), the AHI was more than 5 events/hour with the use of the OA but decreased over 50% compared to the baseline condition (partial success); and in 3 patients (6% of the sample), the AHI was reduced less than 50% from the baseline or not reduced (failure) (Box 1).

According to the AHI results, the patients were divided into two groups: good and poor responders to the treatment. A patient was considered a good responder when the AHI with OA reduced more than 50% from the baseline and was lower than 10 events/hour (n = 43); otherwise, they were considered as poor responders (n = 7) (Tables 2 and 3). It is shown in Table 2 that there were statistically significant improvements in various polysomnographic parameters evaluated in isolation when studying the group of good responders, but it is also important to note that even the group of poor responders showed significant improvement in sleepiness, the AHI and the number of micro-arousals during sleep (Table 3).

A comparison of the results between good and poor responders showed that those who responded poorly to treatment had a greater body mass index (p < 0.05), including weight gained during treatment (Table 4, Graph 2, A and B). On the other hand, two other parameters showed a
A tendency to differentiate, although not in a statistically significant manner; these were age (greater in poor responders) and the severity of OSAS (more severe in poor responders), the latter being insignificant for AHI and oxyhemoglobin saturation. It is noteworthy that the final position (protrusion) with the OA was not different between responders and non-responders (Table 4).
DISCUSSION

The mandibular repositioners for OSAS and snoring treatment may be of two subtypes: adjustable and non-adjustable. This latter category is on the decline due to difficulty of use and worse patient adherence to use. Of the OA, those that seem to be more effective are those of progressive or adjustable mandibular advancement, showing optimal retention as much on the maxilla as on the mandible, in addition to increasing comfort due to a reduced size.8

The differences in the BRD compared to other devices are that it contains a buccal opening and that there are two joining elements between the arches (two expansion mechanisms) positioned on the internal part of the superior arch. Devices that only possess retention in the maxilla cannot maintain a stable mandibular position,2 allowing buccal opening during sleep and a clockwise turning of the mandible that may adversely influence the pharyngeal dimensions. The fact that there are two independent expanders causes the device to position the lower arch in a more stable manner, allowing an anterior position without permitting much buccal opening. Making these devices with a single expander permits greater opening, which does not allow stable mandibular positioning, letting the tongue obstruct the superior airflow passage. Furthermore, a single expander does not allow asymmetric mandibular advancements. In devices made with two expanders positioned externally, the arches have important limitations to mandibular freedom in the lateral direction and the possibility of injury to the jugal mucosa.

This new design, by incorporating two internal expanders, permits asymmetric activation when necessary without impeding small lateral mandibular movements during sleep. Besides this, it does not cause injuries to the dorsal tongue.

The design of the BRD was developed to address these issues, as the device provides a stable mandibular position both vertically (opening) and in the anterior-posterior direction (mandibular protrusion). It also promotes greater comfort, as it is individualized, measurable and extremely versatile with respect to the possibility of progressive mandibular advancements, making its use more physiological.

The results of 30 different studies in 456 patients with OSAS and 224 with snoring during a period of 10 years (1982-1992) shows that intra-oral devices have the potential for significant airway increases, with a 54.3% reduction in the means of OSAS indices and an improvement of snoring in 87.5% of cases.23 In 1995, Nowara et al selected 21 publications with which they reviewed the results of 320 patients treated for OSAS and snoring with OAs. They observed that snoring was completely eliminated in almost all patients and that others showed at least great improvements. For OSAS, there was an improvement of 60% of the apnea-hypopnea index, AHI, on average (AHI before = 47 and after = 19), and approximately half of patients achieved an AHI <10, which is considered successful by many authors.20 Liu et al observed a mean reduction in AHI from 40.3 to 17.1 and an insignificant mean improvement in SaO₂ from 76% to 80% with the Klearway™ device. However, it is noted that these results are not uniform, with some patients responding very quickly to therapy and others in whom the results are not as encouraging. This can be attributed to the lack of standardization of these studies, mainly in relation to selected samples and the process for evaluating results.

In the present study, there was a careful selection of those individuals who actually had a need for OA therapy, and the monitoring of the individuals followed the updated guidelines of the Brazilian consensus of snoring and apnea.7 For this reason, the evaluation of the results involved polysomnographic aspects in relation to respiratory parameters (like the apnea-hypopnea index) but also in relation to sleep architecture (like percentage
of slow-wave sleep and REM sleep). Valid and extremely important clinical criteria such as the Epworth Sleepiness Scale were also used. Statistically significant improvements were observed in several evaluated parameters, like the reduction of sleepiness via the Epworth scale, increased percentage of REM sleep, decreased apnea-hypopnea index, increased minimum oxyhemoglobin saturation as well as the decrease in the number of micro-arousals per hour of sleep (Table 1). These findings corroborate the research of Bernhold and Bondemark, whom reported an improvement in SaO2 and a decrease in daytime sleepiness in the majority of their patients with OA. There was an observed improvement in the sleep quality of the patients, with increased percentages of REM stage, but little influence on the duration of NREM sleep, as also found by Henke; Frantz and Kuna. However, Rose et al reported that the sleep architecture did not significantly change during their study with the Karwetsky-type activator OA.

With regard to obstructive events per hour of sleep (AHI), the results showed that in 56% of the sample the AHI with OA normalized, i.e., it was less than 5 events/hour (success); in 38% of the sample, the AHI was above 5 events/hour with the OA, but it had reduced more than 50% compared to baseline (partial success); and in 6% of the sample the AHI was reduced less than 50% from baseline or not reduced (failure). Generally, the effectiveness is reported based on an improvement in rates reaching 50% to 80% of those obtained initially, but there is still a big controversy on how to standardize the rates of success or the effectiveness of therapy.

When the patients were separated into good responders (AHI with OA reduced more than 50% and less than 10 events/hour) and poor responders to treatment (AHI with OA greater than 10 events/hour), it was found that forty-three patients fell into the first group and seven fell into the second group. Significant improvements in the majority of the evaluated parameters were detected in the group of good responders (Table 2), but it is worth mentioning here that even the group of poor responders showed significant improvements in drowsiness, the AHI and number of micro-arousals during sleep (Table 3).

Liu et al studied patients with OSAS divided into three groups according to the degree of improvement in AHI: a group with good responses (reductions of AHI > 75%), a group with moderate responses (reductions in AHI from 25 to 75%) and a group with poor responses (reductions in AHI < 25%) in relation to treatment with intra-oral devices. This study showed that there was a significant difference in age between the group with poor response and the other groups. In older patients and those with higher BMI values, the OA was less effective. The study presented here also found a greater body mass index, including weight gain during treatment (Table 4, Graph 2, A and B), in patients who responded poorly to treatment with OA. There also seems to be a trend of greater age among the poor responders.

An important aspect is that the final position (protrusion) with the OA was not different between the responders and non-responders. This presupposes that extreme mandibular advancements, without other criteria, will not improve the effectiveness of intra-oral devices. Exaggerated increases in the amount of mandibular advancement do not seem to have a great influence on the improvement of OSAS.

Intra-oral devices offer the most practical and logical way to start a treatment for the majority of OSAS cases. A well-prescribed device that is properly manufactured and periodically monitored is effective, does not represent major expenses for patients and is easily accepted by most of them, while also being associated with other treatment modalities. The device tested in this study achieved satisfactory results and is an alternative to a mandibular repositioning device for the treatment of patients with obstructive sleep apnea.
CONCLUSION

With the utilized therapy, there were significant improvements in the sleepiness scale, AHI, micro-arousals and minimum oxyhemoglobin saturation. Elevated Body Mass Index appears to unfavorably affect the performance of the device under study, suggesting that most patients who are obese or who gain weight during treatment may become an unsuccessful group for this therapeutic approach to OSAS.

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REFERENCES