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Effect of experimental temporomandibular disorder pain on sleep bruxism: a pilot study in males

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Abstract

Objectives Sleep bruxism (SB) is considered to play an important role in the provocation of temporomandibular disorder (TMD) pain. However, clinical studies investigating this relation yielded contradictory results. These contradictory results can, at least in part, be explained by a possible influence of TMD pain on SB activity. The aim of this experimental study was to assess the effect of TMD pain on SB.

Materials and methods Nine male participants with clinical signs of SB underwent two subsequent baseline ambulatory polysomnographic (PSG) recordings before undergoing an experimental pain provocation protocol. Thirty-two hours after the pain provocation part a third ambulatory PSG recording was obtained to study the effect of pain on SB.

Results Decrease for all bruxism parameters was found between the recording after the provocation part and the second baseline recording.

Conclusions Experimentally induced TMD pain causes a reduction in SB activity in healthy individuals.

Clinical relevance A reduction in sleep bruxism activity was recorded in all participants who experienced jaw-muscle pain. This is in line with the pain adaptation model. It supports the negative association between sleep bruxism and jaw muscle pain reported by numerous polysomnographic studies.

Keywords Bruxism · Temporomandibular disorder · Pain provocation experiment

Introduction

Bruxism is defined as a repetitive jaw-muscle activity characterized by clenching or grinding of the teeth and/or by bracing or thrusting of the mandible [1]. It has two circadian manifestations occurring either during wakefulness (wake bruxism) or sleep (sleep bruxism). Among other oral parafunctions, sleep bruxism (SB) is considered very common in the general population, with a self-reported prevalence of up to 41% [2].

In the everyday dental practice, SB has been for many years commonly considered as a major risk factor for temporomandibular disorders (TMDs) [3]. TMD is a collective term embracing a number of clinical problems of the musculoskeletal structures of the masticatory system [4]. The most frequently

reported symptom of TMD is pain, which usually aggravates during masticatory function [5]. The origin of this pain most often lies to the masticatory muscles [6].

The assumption that SB is an important contributor for developing chronic jaw-muscle pain was tested in a number of studies that investigated a one-way relation between the oral parafunction and the TMD condition. Most of these studies had a cross-sectional design and led to contradictory conclusions regarding the association between SB and TMD pain [7, 8]. Based on self-reports, bruxism has been found to be strongly associated with TMD pain.

In contrast, studies that used tooth wear as a manifestation of bruxism failed to prove an association with TMD pain [9, 10].

Currently, polysomnography (PSG) with simultaneous audio/video recording is considered the most accurate method to diagnose SB [11, 12]. Studies that used PSG for setting the diagnosis of SB yielded contradictory results regarding its association with TMD pain: some found a positive association [13], while others detected either no association [14] or even a negative one [15, 16].

These diverging results from studies on the SB-TMD association can, at least in part, be explained by the fact that

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previous studies dealt with patients with SB who suffered from TMD pain at the same time. Such a long-standing musculoskeletal pain, which fluctuates over time in presence and intensity, may have varying effects on jaw-muscle activity. There are two major but conflicting theories aiming to explain the association: the vicious cycle theory [17] and the pain adaptation model [18]. Nowadays, the pain adaptation model is considered to explain such effects best, suggesting that muscle pain leads to a decrease in muscle activity. However, there is still no common agreement on the effects of pain on muscle activity, leading to theories aiming to improve the extrapolation of these models (integrated pain adaptation model; [19]).

The aim of the present study was to assess how experimentally provoked TMD pain, caused by masticatory muscles' overloading, affects SB. We hypothesized that experimentally induced jaw-muscle pain would lead to decreasing jaw-muscle activity during sleep.

Materials and methods

Participants

The following inclusion criteria were used to recruit study participants: self-report of presence of SB and presence of clinical signs of SB viz., muscle hypertrophy, hyperkeratosis of the tongue and cheek mucosa, tongue and lip impressions, tooth attrition, and/or matching wear facets on the teeth.

Participants were excluded when a shortened dental arch was present, because this would hinder wearing a protection splint during the pain provocation part of the protocol, when sleep disorders other than SB were present, when they used on a regular basis painkillers or medications that have a known influence on sleep structure or SB (e.g., selective serotonin reuptake inhibitors, anti-Parkinson medications), when TMD pain was present, when smoking more than five cigarettes per day or drinking more than two glasses of wine or equivalent alcoholic drinks per day, and/or when using a dental splint at night [20, 21].

Participants were also excluded from the study in case they had less than 4 bruxism episodes per hour of sleep [11]. This exclusion parameter was assessed after the second ambulatory polysomnographic recording (see below). Patients were also excluded if no jaw-muscle pain was present 32 h after the experimental jaw-muscle pain provocation exercises.

To calculate sample size, "G power" software was used [22]. The following values to calculate sample size were used: Standard deviation for bruxism episodes per hour of sleep was 2 [23], the significance level was 0.05, the power was 0.8, and the clinically relevant difference was 3 episodes per hour of sleep. This resulted in a sample size of 10.

In total, 23 male participants who self-reported sleep bruxism were screened. Twelve of them demonstrated clinical

signs of sleep bruxism and agreed to participate in the study. The main reason for drop out at this stage was unwillingness to continue participation due to the complicated and time-consuming design of the study.

Further, one participant was excluded after the second PSG recording due to too few (1.9) SB episodes per hour of sleep, and another two were excluded 32 h after the pain provocation test because no jaw muscle pain could be recorded.

Data from nine male volunteers (mean age \pm SD 29.2 ± 5.7 ; range 21–42 years) who fulfilled all inclusion criteria were used for the analysis. All participants were in good health. For all of them, the diagnosis of SB, graded as "probable," according to the diagnostic grading system for SB [1] was established. During the intake procedure, all participants were examined according to DC/TMD guidelines and no TMD pain diagnosis was established. The DC/TMD protocol was supplemented by dynamic and static pain tests [24]. These tests investigate possible pain from the masticatory muscles and/or temporomandibular joints by performing mandibular movements and static muscle efforts. During dynamic tests, the joint moves over its full range of motion against a slight manual resistance. During static tests, mandibular joints are held in place while the muscle exerts a high force. At baseline, all dynamic and static tests were negative.

The DC/TMD examination was performed by the same examiner (KM) for all participants. The examiner was extensively trained for DC/TMD protocol by a calibrated examiner (MK).

The study protocol was approved by the Medical Ethical Committee of the VU Medical Center (file number 2014/179). All subjects signed an informed consent form and received a non-pecuniary compensation (viz., an appointment for professional mouth cleaning and protection splint in case it was indicated) for their participation.

Procedure

At intake, an oral history was taken and a set of questionnaires was administered, including among others a standard Dutch medical questionnaire and the Dutch version of the Sleep Disorders Questionnaire (SDQ). In addition, a clinical examination was performed according to the DC/TMD protocol [25] and supplemented by dynamic and static pain provocation tests [24].

Following the intake procedure, the selected bruxers were invited to sleep with an ambulatory polysomnographic (PSG) device for two consecutive nights. Those of them who had at least 4 SB episodes per hour of sleep based on PSG analysis, underwent an experimental provocation protocol for the jaw muscles. This protocol has been proven to cause delayed-onset muscle soreness (DOMS) with signs and symptoms leading to a TMD-pain diagnosis [26]. Thirty-two hours after the provocation part, participants were again examined

clinically and a third PSG recording was made. These procedures are described below in detail.

Polysomnography

Ambulatory polysomnography (PSG) was performed using an Embla Titanium unit (Embla, Ontario, Canada) in the home environment of the participants for a total of three nights. The first two consecutive recordings were used to confirm the presence of SB and to document the characteristics of SB before the experimental TMD-pain provocation part (baseline). The third recording was performed 32 h after the provocation part to measure the SB when TMD-like pain was present.

The entire montage was performed in each participant's home. Participants were connected to the PSG device around 20:00 time. The recordings were made from approximately 22:30 to 7:00. Time of the recordings was closely adjusted to the participant's usual sleep time.

The electrodes for the PSG recordings were placed according to the protocol recommended by the American Academy of Sleep Medicine [27]. The montage protocol consisted of the following recordings:

1. Electroencephalography (EEG; F3, C4, O1, M1, M2)
2. Electro-oculography (EOG; right and left)
3. Electromyography (EMG; right masseter muscle, submental area)

The right masseter EMG signal was recorded at 512 Hz and filtered (Embla Titanium; 50 Hz notch; 3 Hz high pass; 100 Hz low pass). No audio/video recordings were obtained.

Experimental TMD-pain provocation part

The provocation of experimental TMD pain was performed 32 h before the third PSG registration using the protocol described extensively by Türker et al. [28]. In short, a “custom-made” pain-provocation apparatus was used. Participants were sitting upright in a chair with an adjustable height, so that they could comfortably bite with their central incisors on the biting plates of the apparatus. During the provocation part, series of subsequent eccentric (opening while contracting the jaw closers) and concentric (“regular” closing) contractions of the jaw-closing muscles were performed in 12 sets of exercises, each lasting 5 min and with 1 min of rest in between. During the provocation part, the researcher repeatedly released the compression force of the apparatus, thus allowing the jaw-closing muscles to contract eccentrically.

During the provocation part, participants were constantly biting at a level of 25% of the baseline maximum voluntary contraction level of their right masseter muscle. The values of the EMG activity were displayed to the participant through

visual feedback by the use of a voltmeter, with bipolar electrodes placed over the belly of the right masseter.

In order to protect the teeth from possible damage, participants were wearing soft acrylic mouth guards (Bioplast, 125 mm, clear; Scheu Dental Technology, Iserlohn, Germany), made on individual plaster casts, on both dental arches throughout the entire provocation part.

Pain measurement

Participants were asked to rate the amount of jaw-muscle pain intensity on a questionnaire with a Numeric Rating Scale (NRS) every evening before the PSG recordings. Thus, a total of three NRS scores were obtained. Moreover, 32 h after the provocation part, a standardized clinical examination of the masticatory muscles and joints, according to the DC/TMD criteria supplemented by dynamic and static pain test, was performed in order to evaluate the presence of jaw-muscle pain (see above: procedure).

Data analysis

Before the PSG analysis, all registrations were coded to ensure that the examiner (KM) was blinded for the specific PSG recording under analysis.

Subsequently, all PSG recordings were analyzed with the use of RemLogic (Embla, Ontario, Canada) and Bruxism Detector (ACTA, Amsterdam, The Netherlands; custom-made) software. The analyses consisted of two parts: a sleep analysis and a SB analysis. The sleep analysis was performed to exclude sleep disorders other than SB in addition to the SDQ (see above), to determine any abnormalities in sleep structure, and to enable the sole inclusion of masticatory muscle activities during actual sleep in the analysis of SB. Using 30-s epochs, all sleep analyses were carried out automatically and checked manually according to the criteria described in The American Academy of Sleep Medicine Manual for the Scoring of Sleep [27]. Total sleep time and percentage of time spent in each sleep stage were calculated.

As a first step in the analysis of SB, the sleep stages and the EMG signal from the right masseter were transferred from RemLogic to Bruxism Detector software, using European Data Format (EDF). Bruxism Detector is a custom-made software that uses clinical research diagnostic criteria for SB (RDC/SB) proposed by Lavigne et al. [11] to quantify rhythmic masticatory muscle activity (RMMA) episodes. The EMG signal was low-pass and high-pass filtered (100 Hz and 5 Hz, respectively). During the next step, periods of increased EMG activity were detected using a threshold of three times above the noise level. During the last step, SB outcome variables, presented as the number of bruxism episodes per hour of sleep (Epi h⁻¹), the number of bursts per hour of sleep (Bur h⁻¹), and the bruxism time index (BTI: the percentage of

total sleep time spent bruxing) were calculated, and only the outcome variables detected during sleeping periods were used in the analyses. All analyses were manually checked.

Statistics

The non-parametric Friedman test was used to analyze the data. First, it was verified whether there were natural fluctuations in jaw muscle activity across the nights, using the SB data of the first two PSG recordings. Subsequently, it was tested whether there was a significant difference in SB activity between the night before and after the provocation part, comparing the data of the second and third PSG recordings. Statistical analysis was performed using IBM SPSS Statistics 23 software (IBM Corp., Armonk, NY, USA).

Results

Thirty-two hours after the DOMS provocation test, five participants reported pain graded as 1 on the 11 points NRS scale, three participants graded pain as 2, and one graded his pain as 8. For all of them, the diagnosis of myalgia could be established based on the DC/TMD protocol. They also responded positively with pain during both the dynamic and static tests. All nine participants described the pain during the test as mild and familiar to the pain they experienced after the provocation.

Based on the PSG recordings, all hypnograms (two baseline recordings and the recording 32 h after the jaw-muscle pain provocation) had a normal structure. Table 1 shows the mean values of the sleep variables.

The SB variables are presented in Table 2. The Friedman tests showed that there was no difference between the two PSG recordings made at baseline (first and second PSG recordings; $p = 0.26–0.74$). In contrast, after the pain provocation test (third PSG recording), a decrease for all SB parameters was recorded compared with the measurements of the second baseline PSG recordings ($p = 0.01–0.03$).

The decrease in SB activity varied among participants: for three of the participants, there was < 15% decrease in $Epi\ h^{-1}$,

for four participants, there was a decrease between 15% and 50%, while for two participants, there was > 50% decrease (Figs. 1 and 5).

Discussion

The aim of the present study was to assess how experimental TMD pain, caused by masticatory muscles' overloading, affects SB. We hypothesized that experimentally induced jaw muscle pain would lead to decreasing jaw muscle activity during sleep. To that end, we provoked TMD-pain using a previously published protocol in pain-free bruxers and recorded the SB activity in the absence (before provocation) and presence (after provocation) of pain. Our results confirmed our research hypothesis, because a decline in SB activity was found for all nine participants who experienced jaw muscle pain after the pain provocation tests.

The effect of pain on jaw muscle activity has been a subject of interest in previous experimental studies as well. The results of these studies were contradictory: some showed an increase in muscle activity in response to pain [29], while others detected either no [30] or a negative effect of pain on jaw muscle activity [31]. These previous studies either used intramuscular injections (e.g., with capsaicin) to provoke the pain in humans [30] or were performed in animals [29]. To the best of our knowledge, this is the first study in humans on the effects of experimentally provoked TMD pain, due to muscle overloading, on SB.

The provocation protocol used in the present study has been previously described in detail and tested in a group of 40 healthy participants. It was shown that a state of DOMS can be provoked in healthy individuals, with signs and symptoms that lead to the diagnosis of TMD pain [26]. The advantage of this protocol, in contrast to the experimental models provoking short-lasting acute pain like the injection of painful substances into the muscles, is the fact that the pain is longer lasting and is present for several hours after the provocation part. Moreover, this type of experimental pain mimics the signs and symptoms of clinical TMD pain better, as compared to pain caused

Table 1 Descriptive statistics (mean \pm SEM) of the standard sleep variables of the two baseline nights and the night 36 h after experimental jaw-muscle pain provocation

Sleep variable	Night 1	Night 2	Night 3	P (N1-N2)	P (N2-N3)
Total sleep time (min)	456 \pm 25.7	473 \pm 29.4	443 \pm 21.2	0.37	0.21
Proportion of sleep (%)					
Awake	5.3 \pm 1.2	3.3 \pm 1.9	4.2 \pm 2.8	0.58	0.44
Stage 1	3.0 \pm 1.1	5.9 \pm 3.1	1.7 \pm 1.5	0.61	0.93
Stage 2	53.8 \pm 6.2	56.1 \pm 4.9	61.6 \pm 9.4	0.18	0.59
Stage 3	20.9 \pm 4.5	18.5 \pm 1.4	17.8 \pm 3.6	0.32	0.27
REM	17.1 \pm 2.1	16.1 \pm 4.4	14.8 \pm 3.9	0.76	0.82

Table 2 Sleep bruxism variables (median; first and third quartiles, *p* values based on Friedman test) of the two baseline nights and the night 36 h after experimental jaw muscle pain provocation

Bruxism	Night 1	Night 2	Night 3	P (N1-N2)	P (N2-N3)
Epi h ⁻¹	9.2 (7.8–12.7)	9.7 (7.3–13.5)	7.02 (5.7–10.2)	0.74	0.03
Bur h ⁻¹	66.7 (47.7–79.9)	75.2 (45.1–99.5)	51.1 (36.5–70.1)	0.32	0.01
BTI	2.2 (1.6–2.4)	2.0 (1.4–2.7)	1.7 (0.9–2.1)	0.48	0.01

Epi h⁻¹ episodes per hour of sleep, *Bur h⁻¹* bursts per hour of sleep, *BTI* Bruxism time index

by other experimental protocols, like intramuscular injections. That is because the study protocol involves actual tissue damage due to overloading.

The way pain and muscle activity are related has been a matter of debate for several decades, leading to the publication of two major theories in order to explain this relation: the vicious cycle theory [17] and the pain adaptation model [18]. These two theories contradict each other. The vicious cycle theory suggests that an initiating factor, which could be SB for example, causes pain that reflexively leads to muscle spasm. This spasm leads to further pain and dysfunction, thus completing and perpetuating the loop. However, the evidence that supports the vicious cycle theory is until now limited [32, 33].

The pain adaptation model, on the other hand, suggests that muscle pain leads to a reduction in muscle activity of the painful muscles, aiming to protect the muscle system from further injury and, therefore, promote healing [34]. This model is commonly considered to explain most appropriately the effects of pain on muscle performance [19]. The reduced SB activity found after the experimental TMD pain was provoked and is in accordance with the Pain Adaptation Model. The pain causes a decrease in the muscle activity most probably in order to protect the muscles from further damage. This is in agreement with previous studies that found that in patients with TMD pain, the PSG-recorded bruxism is decreased compared with healthy controls [14].

The following possible limitations of this study should be kept in mind. First, even though we used a provocation protocol inducing longer-lasting TMD pain compared with other pain-provocation modalities, this pain still has an acute nature because it disappears after 1 week. TMD pain though is a chronic condition. It is likely that the relationship between chronic TMD pain and SB is more complex than just a linear association, and also that other factors, such as general health, genetics, and psychological status, may play an important role on the association under study [35].

Second, TMD pain has a higher prevalence in women [36]. In this study, we provoked jaw muscle pain in male volunteers in order to exclude the possible effects of cycle-dependent hormonal fluctuations on pain perception [37]. This limits the generalizability of our findings to the

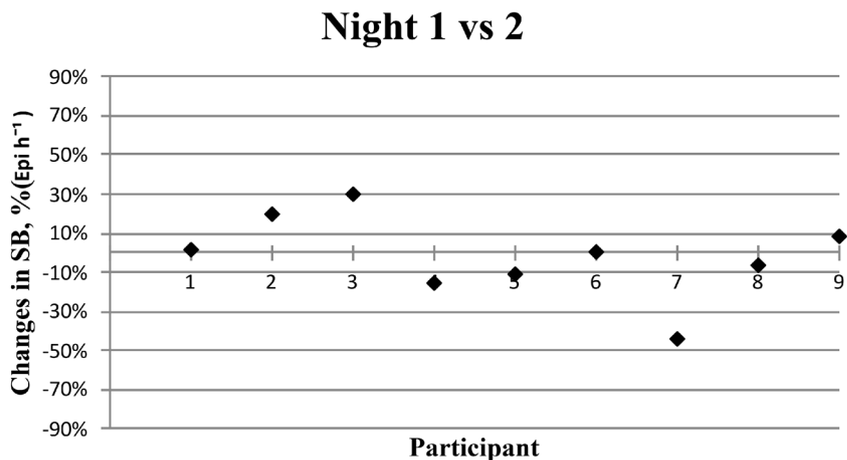
general TMD-pain population. It could also be interesting to investigate the possible gender differences in the association between TMD pain and SB, and whether hormones or other gender-related factors play a role in this association.

Third, even though we found a significant difference in SB activity before and after the provocation of experimental TMD pain on the group level, the individual participants reacted differently to the presence of pain. One participant showed a decrease of about 90% in SB activity, while another one demonstrated almost no changes in SB characteristics over the course of the three nights (see “Results” and Figs. 1 and 5). Considering an unusually low number of SB activity recorded for participant no. 3 (see Fig. 1, 2, 3, 4, and 5) during the third night registration, we checked whether his exclusion would provide a different study result. The difference between second- and third-night registrations still remained significant.

Hence, due to the high variation of the individual reactions to jaw muscle pain and the small size of the group, the extrapolation of our results to the population should be done with caution. Moreover, this variation could be an explanation why some patients develop chronic pain while others do not. One can speculate that a decrease of 90% in SB activity as a response to muscle damage leads to adequate protection from further injury and promotes healing. In contrast, it is unclear whether a decrease in muscle activity of, e.g., 15%, is enough to protect damaged muscle tissue. It can be further speculated that some patients failed to “adjust” their jaw muscle activity to the amount that is needed for the recovery, and therefore damage the muscles. This phenomenon could be similar to the overtraining syndrome that is known to occur in limb muscles after excessive training.

Continuous low intensity load on the damaged limb muscle, such as walking, can cause a repetitive tissue trauma and lead to the development of chronic pain [38]. In our case, all participants were contacted 1 week after the provocation, and none of them reported any pain complaints. We also checked whether the variation in the decreased SB activity could be explained by the amount of pain that participants reported on NRS scale. The

Fig. 1 Difference in SB activity measured by Epi h⁻¹ between first and second nights (%). *Epi h⁻¹—episodes per hour of sleep



statistical analysis did not yield any significant results regarding a dose-response association ($r = -.03$, $p = .92$, based on Spearman’s correlation test).

Further, we recorded unusually high rates of SB episodes in our study (e.g., the average for the first two nights was nine SB episodes per hour of sleep). Previous studies that used sleep-laboratory environment with simultaneous audio/video registration demonstrated that the amount of sleep bruxism events per night is relatively low. For example, a previous study [39] showed that out of 146 participant who underwent two nights laboratory-based polysomnography registration, only 76 participants (52%) had more than 1 SB episode per hour of sleep, and only 18 (13% of the studied sample) had more than four SB episodes per hour of sleep, fulfilling the criteria for the high SB intensity group. Studies that compared the outcome of ambulatory polysomnography registrations with laboratory-based registrations have showed that the first one tends to overestimate SB [40]. With the absence of audio/video recordings, it is difficult to distinguish SB from other types of

oromotor activities such as swallowing, lip sucking, chewing-like movements, etc. To counter the above-mentioned disadvantage of ambulatory polysomnography recording, an additional registration channel can be used. For example, it has been documented that an increase in cardiac sympathetic activity ([41]) and an increase in breathing amplitude [42] precede SB. In future studies, registration of, e.g., heart rate, may improve the accuracy of scoring SB and may reduce false-positive outcomes of ambulatory PSG recordings. The diagnostic validity of these additional methods to diagnose SB has however yet to be verified.

Last but not least, the way the threshold for SB is defined could introduce a potential bias when patients with and without jaw muscle pain are compared. Previous study of Raphael et al. [43] suggested that the levels of EMG activity occurring outside SB and other nocturnal motor events are significantly higher in patients experienced TMD pain compared to pain-free controls.

In fact, this background EMG activity is routinely considered as the threshold to define SB episodes. It

Fig. 2 Difference in SB activity measured by Epi h⁻¹ between second and third nights (%). *Epi h⁻¹—episodes per hour of sleep

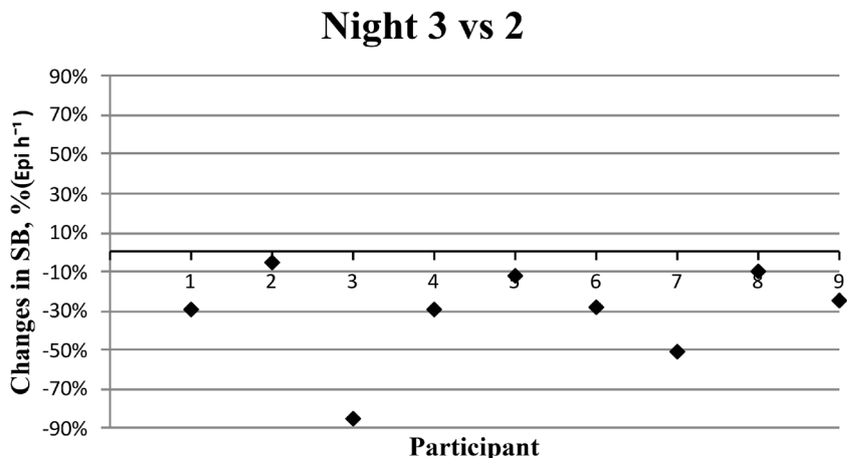
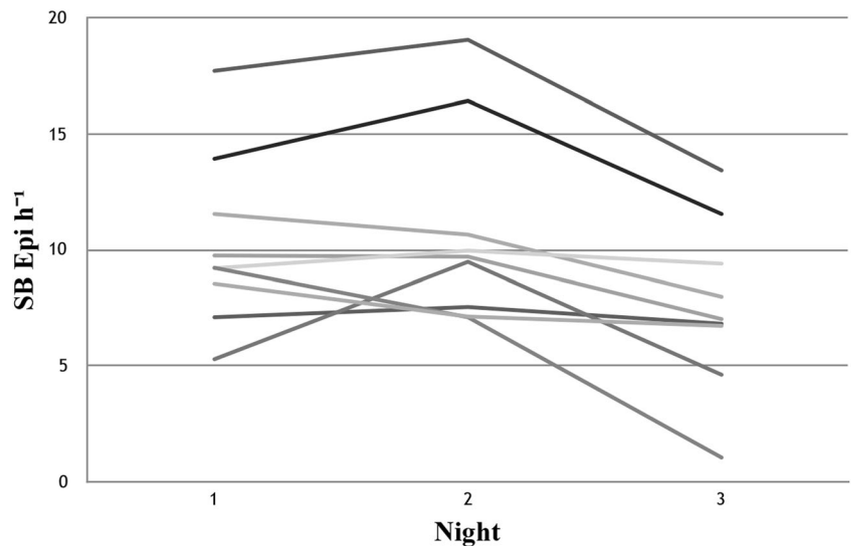


Fig. 3 Individual difference in SB episodes between first, second and third nights. *Epi h⁻¹—episodes per hour of sleep



can be therefore speculated that the thresholds used to identify SB activity in this study were significantly higher after the pain provocation exercise when compared to two nights baseline registrations. These higher thresholds recorded in the presence of pain would univocally lead to inclusion less SB episodes during the third night registration when compared to two baseline registrations in the absence of jaw muscle pain. The alternative threshold could be the one based on percentages from the maximum voluntary contractions (MVCs), as proposed by Lavigne et al. [11].

However, using this threshold in the presence of jaw muscle pain could also introduce a bias. Participants could try to avoid pain during function and therefore do not express maximum bite force during the MVC registration at third night

PSG recording when jaw muscle pain was present. Therefore, there is a need for further investigation of the most reliable threshold for studies which aimed to compare the jaw muscle activity in TMD pain patients and pain-free individuals.

In conclusion, our study suggests that experimentally induced TMD pain reduces SB activity, which is in line with the pain adaptation model. It can be therefore expected that patients who experience TMD pain have less SB activity than pain-free individuals. Moreover, it can be assumed that eliminating jaw muscle pain during treatment could increase SB activity. Further research should take into account that the interaction between TMD pain and SB is complex, and caution in interpreting the results is required.

Fig. 4 Individual difference in SB bursts between first, second, and third night. *Bur h⁻¹—bursts per hour of sleep

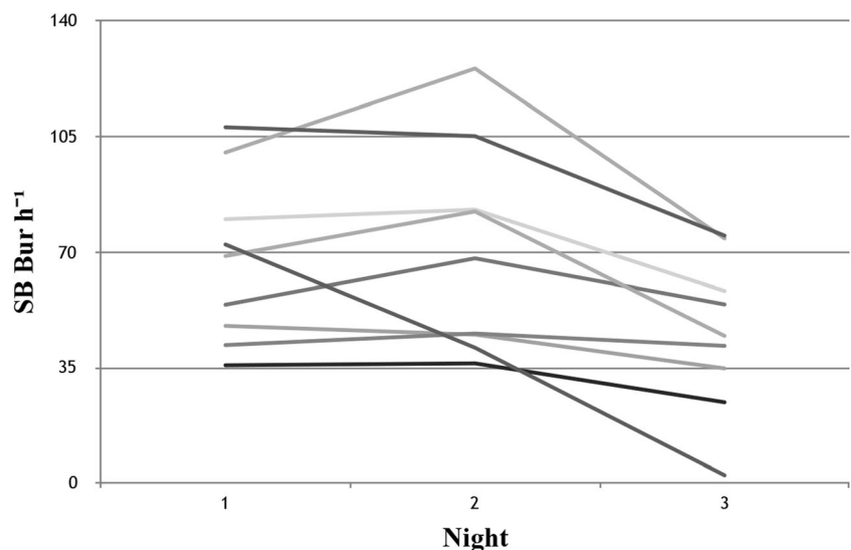
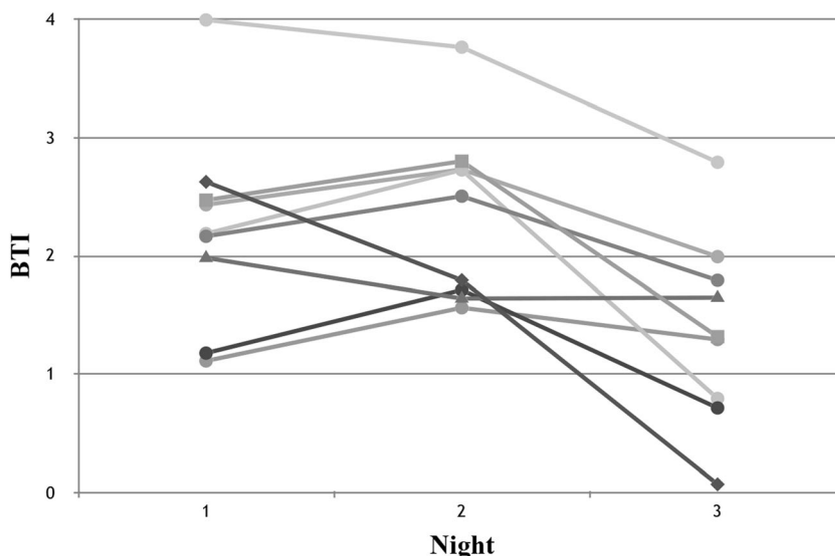


Fig. 5 Individual difference in SB time index between first, second, and third nights. *BTI—Bruxism time index



Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Ethical approval All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Informed consent Informed consent was obtained from all individual participants included in the study.

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