Decreased sleep quality and increased sleep related movements in patients with Tourette’s syndrome

S Cohrs, T Rasch, S Altmeyer, J Kinkelbur, T Kostanecka, A Rothenberger, E Rüther, G Hajak

Abstract

Objective—Sleep quality and movement patterns across sleep stages in patients with Tourette’s syndrome were examined to determine the influence of syndrome severity on sleep quality and the differential effect of sleep stages on tic and non-tic movements.

Methods—Twenty-five patients with Tourette’s syndrome (mean age 29 (SD 7) years) and 11 control subjects (29 (5) years) were studied by polysomnography and simultaneous split screen video monitoring to record standard sleep variables as well as to evaluate movements to differentiate between tics and regular movements. Severity of Tourette’s syndrome during the day was assessed with the Tourette’s syndrome severity scale.

Results—Sleep was significantly more disturbed in patients with Tourette’s syndrome than in controls, with decreased sleep efficiency and slow wave sleep percentage, increased sleep latency, percentage of stage I, percentage of awakeness, number of awakenings, and sleep stage changes and more overall movements during sleep. Severity of Tourette’s syndrome during the day correlated significantly and positive with number of awakenings and sleep stage changes and negatively with sleep efficiency. In addition to an increased number of regular movements patients had tics in all sleep stages. Tic frequency as well as frequency of regular movements was significantly higher in REM than in non-REM sleep which was also the case for regular movements of the controls. No disturbance of either REM sleep percentage or REM latency was found.

Conclusion—Despite normal total sleep time and unaltered REM sleep variables patients with Tourette’s syndrome have markedly disturbed sleep. Severity of the syndrome during the day is an important predictor of sleep alteration in patients. The increased rate of tics during REM sleep parallels the overall increased movement activity of patients during REM sleep as well as non-REM sleep. The increased motor activity may be attributable to a state of hyperarousal rather than a disturbed cholinergic system.

Keywords: Gilles de la Tourette’s syndrome; polysomnography; sleep; movement analysis

Gilles de la Tourette’s syndrome is a neuropsychiatric disorder that is characterised by multiple motor and vocal tics. It usually starts in childhood and persists in most patients through adulthood with a waxing and waning symptomatology of phenomenologically changing tics of varying severity. Besides other associated symptoms such as coprolalia, attentional deficit disorder, obsessive compulsive traits, echolalia, copropraxia, and echopraxia, up to 60% of the patients with Tourette’s syndrome complain about disturbed sleep. Most often reported are complaints of insomnia such as difficulty falling asleep or early awakening.

Few polysomnographic studies with varying methodological quality have been undertaken to determine the objective sleep quality of patients with Tourette’s syndrome. Conflicting results have been reported concerning an unaffected or decreased sleep latency, unaffected or prolonged sleep latency, increased, decreased or unaffected slow wave sleep, and reduced or unaffected rapid eye movement (REM) sleep. No data exist about the influence of Tourette’s syndrome severity on objective sleep parameters. Furthermore, the origin and pathophysiology of tic related sleep disturbances has not been unequivocally clarified.

Motor phenomena during sleep in patients with Tourette’s syndrome have been interpreted in different ways ranging from REM sleep behaviour disorder to periodic limb movements. Earlier studies reported that tics occur during REM as well as non-REM sleep. However, no data exist on the frequency, intensity, and complexity of tics across sleep stages. Data on tic patterns related to sleep stage are of particular interest to better understand the pathophysiology of Tourette’s syndrome. Different neurotransmitter systems have been discussed as being involved in pathophysiology of the disease. Besides the dopaminergic, serotonergic, opioid, and noradrenergic system it has been postulated that the cholinergic system is involved in the pathogenesis of tics.

Considering the aminergic-cholinergic interaction model of sleep regulation a differential effect of sleep stages on tics and regular (non-tic) movements would be expected. Taking into account that the various neurotransmitters involved in the regulation of sleep have been hypothesised to play a major part in the pathophysiology of tics in patients with Tourette’s syndrome the aim of this study was to (1) objectively identify sleep quality in patients...
with Tourette’s syndrome and (2) to determine the influence of severity of Tourette’s syndrome on standard polysomnographic variables as well as on the frequency and type of movements exhibited in comparison with healthy subjects in REM and non-REM sleep.

**Methods**

**PATIENTS AND CONTROLS**

Twenty-five patients with Tourette’s syndrome (mean age 29.04 (SD 6.8) years, range 16–44; 16 male and nine female) and a group of 14 healthy volunteers (28.7 (SD 5) years, range 23–39; eight male and six female) were studied for comparison of sleep variables. Eleven of these 14 healthy subjects (28.9 (SD 5.2) years, range 24–39 years; eight male and four female) were included in the video analysis of movement variables. Patients were recruited from the outpatient clinic and self-help groups where two of the authors (TR and SA) presented the possibility to the participants of being examined polysomnographically within a scientific study. Control subjects were recruited by word of mouth, mostly being university associated students, employees, or friends of these who heard of the study and were interested in participating. A detailed clinical history was taken to ensure current health of the control subjects. All patients fulfilled the diagnostic criteria for Tourette’s syndrome according to the International Classification of Diseases, 10th edition (ICD-10) and the Diagnostic and Statistical Manual, 4th edition (DSM IV). Demographic data of the patients, current medication, and severity of the disease according to the Tourette’s syndrome severity scale (TSSS) are summarised in Table 1. Severity ranged from 0.5 (mild) to 8.0 (severe) with most of the patients being moderately to severely affected.

The study was approved by the local ethics committee and was conducted according to the declaration of Helsinki. Each subject gave written informed consent after a complete description of the study.

**POLYSOMNOGRAPHY**

All subjects underwent polysomnographic recordings for two consecutive nights according to standard criteria. To exclude first night effects only results of the second night were analysed. Polysomnography included EEG (C3/A2), electro-oculogram, submental EMG, ECG, and EMG of the anterior tibial muscles. Additional superficial EMG leads were placed on the muscle groups exhibiting typical tics during wake time (sternocleidomastoid, trapezius, masseter, and brachioradial muscles). Patients and subjects were allowed to go to bed at their usual bedtime and to get up at their convenience. Sleep was recorded at a paper speed of 10 mm/s on a 21 channel polysomnograph (Nihon Kohden, Japan) and a computerised video monitoring system. Sleep was scored according to the standard criteria of Rechtschaffen and Kales in 30 second epochs. Evaluation of EEG arousals followed criteria of the American Sleep Disorders Association.

**MOVEMENT ANALYSIS**

In addition to polysomnographic recordings subjects’ movements were monitored with an infrared camera. Illumination was provided by two infrared lighting systems, allowing excellent video recordings in a dark room. Sound was recorded from a microphone located 91 cm (3 feet) above the subject’s bed. Split screen video monitoring was used to have the visual image as well as polysomnographic data stored on the same tape. With this technique simultaneous analysis of polysomnographic variables, visual analysis of movements, and evaluation of audio signals was possible.

Patients and subjects spent 2 days in the sleep laboratory. Daytime observation gave an opportunity to the scorers (TR and SA) to become familiar with the individual tics of the patients. Tics were classified using a rating system which followed the DSM-IV definition of a tic as a sudden, rapid, recurrent, non-rhythmic stereotyped movement or vocalisation. Inconclusive scoring of a particular movement was followed by evaluation of this movement by the second scorer and final classification was done after consensus conference. Tic intensity was classified into three levels as: mild (1), moderate (2), and severe (3). Key aspects of tic intensity were strength, impulse, and range of the tic movement. Tic complexity was divided into two categories according to Leckman et al: Simple tics consist of purposeless abrupt movements such as grimacing, eye blinking, shoulder shrugging etc. Complex tics were slower movements that look as if they had a purpose such as touching, biting, head banging, itching etc.

Tics and regular movements during sleep of patients and control subjects were scored by analysis of the split screen video recordings. Because of the brief and abrupt nature of tics it was possible to identify and count individual tics; whereas it was often impossible to determine the end of a regular movement and the beginning of the next regular movement. For determination of reliability of movement scoring the concordance rate of the two raters...
Table 2 Sleep variables of patients with Tourette’s syndrome (n=25) compared with healthy controls (n=4)

<table>
<thead>
<tr>
<th>Sleep variables</th>
<th>Tourette patients (n=25)</th>
<th>Controls (n=14)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>TIB (min)</td>
<td>519 (81)</td>
<td>486 (53)</td>
<td>NS</td>
</tr>
<tr>
<td>SPT (min)</td>
<td>474 (87)</td>
<td>464 (56)</td>
<td>NS</td>
</tr>
<tr>
<td>TST (min)</td>
<td>428 (93)</td>
<td>446 (60)</td>
<td>NS</td>
</tr>
<tr>
<td>SE (TST in %TIB)</td>
<td>82 (10)</td>
<td>92 (6)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Sleep latency (min)</td>
<td>40 (49)</td>
<td>17 (9)</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>REM latency (min)</td>
<td>97 (62)</td>
<td>94 (38)</td>
<td>NS</td>
</tr>
<tr>
<td>Wake% (SPT)</td>
<td>9.6 (5.6)</td>
<td>3.9 (3.7)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>StI% (SPT)</td>
<td>11.9 (4.1)</td>
<td>8.7 (2.4)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>StII% (SPT)</td>
<td>47.1 (9.1)</td>
<td>49.4 (5.8)</td>
<td>NS</td>
</tr>
<tr>
<td>SWS% (SPT)</td>
<td>10.2 (7.0)</td>
<td>14.7 (3.9)</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>REM% (SPT)</td>
<td>20.8 (4.0)</td>
<td>22.8 (4.8)</td>
<td>NS</td>
</tr>
<tr>
<td>StI% (TST)</td>
<td>13.3 (5.1)</td>
<td>9.1 (2.9)</td>
<td>&lt;0.005</td>
</tr>
<tr>
<td>StII% (TST)</td>
<td>51.9 (9.0)</td>
<td>51.3 (5.8)</td>
<td>NS</td>
</tr>
<tr>
<td>SWS% (TST)</td>
<td>11.2 (7.8)</td>
<td>15.2 (3.9)</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>REM% (TST)</td>
<td>23.4 (4.5)</td>
<td>23.7 (4.9)</td>
<td>NS</td>
</tr>
<tr>
<td>Awakenings/h SPT</td>
<td>2.8 (1.6)</td>
<td>1.5 (0.7)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Sleep stage changes/h SPT</td>
<td>18.6 (5.5)</td>
<td>15.8 (2.4)</td>
<td>&lt;0.05</td>
</tr>
</tbody>
</table>

Values are means (SD), t test (two tailed); TIB=time in bed; SPT=sleep period time; TST=total sleep time; SE=sleep efficiency= TST/TIB; Sl= sleep latency; REM-latency= latency to REM sleep from sleep onset; W%, StI%, StII%, SWS%, REM% (SPT)/(TST)=percentage of wake, sleep time (% REM), and REM latency (time from lights out as well as from sleep onset to first epoch of REM sleep) of the patients with Tourette’s syndrome were free of medication for at least 4 weeks. The other 12 patients were taking different medication including neuroleptic and antidepressant drugs (table 1). One patient was included into the group of medicated patients because he medicated himself by frequent application of cannabis.

None of the sleep variables showed a significant difference between medicated and unmedicated patients, which might be due to the fact that treated and untreated patients had tics and associated symptomatology to a comparable degree. Therefore, data for the patients were pooled for further analysis. Time in bed, sleep period time (time from first sleep onset to awakening), total sleep time (sleep period time= time awake), percentage sleep stage II of total sleep time, percentage REM sleep of total sleep time (% REM), and REM latency (time from lights out as well as from sleep onset to first epoch of REM sleep) of the patients with Tourette’s syndrome were not significantly different from the control group. However, in comparison with the control subjects (table 2) patients with Tourette’s syndrome showed significantly reduced sleep efficiency (total sleep time/time in bed) and percentage of slow wave sleep, significantly prolonged sleep latency, significantly increased percentage of sleep stage I, percentage of time awake, and increased number of awakenings and sleep stage changes/hour sleep period time.

A significant correlation between Tourette’s syndrome severity score and number of awakenings/hour, number of sleep stage changes during sleep period time, and sleep efficiency was found (fig 1).
There was no significant differential effect of tics versus non-tics in movements in consecutive epochs of wake after a tic or non-tic movement. However, epochs with only tics showed significantly less often EEG arousals than epochs with non-tics movements (55.4% (17.1%) vs 73.8% (18.3%); p<0.001). If both tics and non-tics movements occurred in one epoch there were significantly more often EEG arousals in comparison with either only tics or only non-tics movements (88.0% (19.8%) vs 55.4% (17.1%); p<0.01; 73.8% (18.3%); p<0.01 respectively).

**Discussion**

**SLEEP VARIABLES**

Patients with Tourette’s syndrome showed a markedly altered sleep quality with difficulties initiating and maintaining sleep. They had significantly longer sleep latency, reduced sleep efficiency, more percentage stage I, higher frequency of sleep stage changes as well as awakenings along with a reduced slow wave sleep percentage. The comparison with data from other studies is only partially possible because most of the authors have not provided essential information about basic polygraphic variables including total sleep time. 8 17–19 The studies providing this polygraphic information examined only very small groups of patients17 18 (two to seven) and lacked details about the time of determining the end of the polygraphic registration. This might be of relevance because of the circadian regulation of REM sleep propensity with more REM sleep in the morning hours.20 Patients and subjects in our laboratory were allowed to sleep at the time of their convenience including spontaneous awakenings in the morning. Thereby the patients with Tourette’s syndrome might have been able to compensate for their lost sleep during the night, which would explain the unaltered total sleep time in comparison with the healthy controls in our study. The unchanged REM percentage together with the unaltered REM latency support the hypothesis of a stable and unaffected cholinergic system in Tourette’s syndrome.

The two aminergic transmitters serotonin and noradrenaline (norepinephrine) have been found to be involved in sleep regulation as well as pathophysiology of Tourette’s syndrome.19–21 An unaltered serotonergic firing has been shown to be a prerequisite for sleep maintenance.22 Our finding of a significant positive correlation between Tourette’s syndrome severity and features of disturbed sleep quality is compatible with the inverse correlation between tic severity and CSF concentrations of tryptophan, a precursor substance of serotonin.20 Both tics and insomnia of patients with Tourette’s syndrome might be related to reduced serotonergic activity. Increased noradrenaline in patients with insomnia on the other hand is considered as a sign of hyperarousal23 that has been found in Tourette’s syndrome24 as well. Together with increased CSF corticotropin releasing hormone concentrations in patients with Tourette’s syndrome24
it may reflect similar psychophysiological hyperactivity in Tourette’s syndrome.

In analogy to the reciprocal interaction of reduced sleep quality and increased daytime functioning in patients with primary insomnia\(^2\), we hypothesise a reciprocal interaction of reduced sleep quality and tic severity during the daytime in patients with Tourette’s syndrome. This is consistent with the positive correlation between Tourette’s syndrome severity and variables of sleep disturbance found in our study—that is, an increased Tourette’s syndrome severity during the daytime being associated with a decreased sleep quality. Sleep with an increased number of awakenings as a sign of reduced sleep continuity is known to be less restorative.\(^2\) As with patients with insomnia\(^2\), patients with Tourette’s syndrome might be more irritable during the awake hours because of their reduced sleep quality. Triggered by this irritability they are prone to exhibit more tics in distressful situations.\(^2\) The additional psychological strain induced by the need for coping with tics might maintain the hyperarousal. The hyperarousal in turn may affect the ability to initiate and maintain sleep making it less restorative, thereby closing the vicious circle of daytime tic intensity and sleep disturbance.

Because of the possible negative effects of insomnia on Tourette’s syndrome symptomatology a disturbed sleep through tics should be taken into consideration in the planning of treatment strategies.

**MOVEMENT ANALYSIS**

The markedly disturbed sleep of patients with Tourette’s syndrome was accompanied by increased motor activity in REM as well as non-REM sleep in comparison with healthy controls. This increased frequency of movements was not only due to the additional tics in patients with Tourette’s syndrome but was also caused by an equally increased frequency of non-tic movements in patients with Tourette’s syndrome. This can be interpreted as a sign of hyperarousal and is concordant with a reduced intracortical inhibition of motor pathways in Tourette’s syndrome.\(^10\)

The higher incidence of tics during REM sleep is paralleled by a similar increase in non-tic movements of patients with Tourette’s syndrome in this sleep stage. A comparable relation between movement frequency during REM sleep and non-REM sleep was found in our control subjects. A nearly 50% increase in movement frequency in REM sleep in comparison with non-REM sleep had been reported for healthy subjects in earlier work.\(^31\)–\(^34\)

Therefore similar mechanisms controlling the occurrence of tic and non-tic movements during this sleep stage are likely. In fact, the activity of all motor pathways is greatly enhanced in an apparently random manner during REM sleep\(^10\) and might account for the higher movement frequency despite the markedly reduced muscle tone during this sleep stage.

In addition to the increased movement frequency the overall increased activity of motor pathways might be the source of the slight but significant increase in tic intensity during REM sleep in comparison with non-REM sleep. The absolute level of tic intensity, however, remains the same both in REM as well as non-REM sleep. Most of the tics exhibited during REM and non-REM sleep were simple tics by contrast with the frequent occurrence of complex tics during wakefulness. No difference in complexity of tics between REM and non-REM sleep was found. Unlike movement frequency tic intensity and tic complexity seemed to be largely un influenced by the type of sleep.

Often, movements during sleep are accompanied by short arousals. These EEG arousals during sleep reflect a brief cortical activation. Patients with Tourette’s syndrome showed EEG arousals in conjunction with a movement during sleep significantly less often than healthy controls. This was mainly due to a reduced EEG arousal frequency in the temporal vicinity of tics. Therefore, tics seem to be accompanied by less cortical activity than regular movements. A corresponding aspect of the reduced cortical activity accompanying tics is the absence of a normal premovement EEG potential in awake patients with Tourette’s syndrome before involuntary tics.\(^36\) These findings are consistent with the hypothesis that subcortical parts of the cortico-striato-thalamocortical circuits play a major part in the pathophysiology of tics.\(^26\)

Within the search for important factors in the pathophysiology of Tourette’s syndrome there is a debate about the voluntary versus involuntary nature of tics. Sleep can be considered as a stage of extremely reduced or nearly absent voluntary control of cognition, emotion, and movement. Together with the occurrence of tics in all sleep stages the reduced EEG arousal frequency of tic movements is consistent with the hypothesis that involuntary aspects play a major part in the pathophysiology of tics.

---