Clinical course of H1N1-vaccine-related narcolepsy

Sarkanen, Tomi

2016-03


http://hdl.handle.net/10138/224040
https://doi.org/10.1016/j.sleep.2015.11.005

Downloaded from Helda, University of Helsinki institutional repository.

This is an electronic reprint of the original article.
This reprint may differ from the original in pagination and typographic detail.
Please cite the original version.
Clinical course of H1N1-vaccine-related narcolepsy

Tomi Sarkanan b,c,*, Anniina Alakuijala b,d, Markku Partinen b,c

a Clinic of Neurology, Central Finland Central Hospital, Jyväskylä, Finland
b Helsinki Sleep Clinic, Vitalmed Research Centre, Helsinki, Finland
c Department of Neurological Sciences, University of Helsinki, Helsinki, Finland
d Department of Clinical Neurophysiology, HUS Medical Imaging Center, Helsinki University Central Hospital, Finland

ABSTRACT

Objective: To follow and analyze the clinical course and quality of life of Pandemrix H1N1-vaccine-related narcolepsy (pNT1).

Methods: Twenty-six drug-naïve confirmed pNT1 subjects completed Epworth Sleepiness Scale (ESS), Ullanlinna Narcolepsy Scale (UNS), Swiss Narcolepsy Scale (SNS), Rimón’s Brief Depression scale (RDS), and WHO-5 Well-being index questionnaires near the disease onset and in a follow-up a minimum of two years later. The number of cataplexies and body mass index (BMI) were recorded. The effects of hypocretin-1 levels and sleep recording results were analyzed. The findings at the follow-up visit were compared with 25 non-vaccine-related type 1 narcolepsy (NT1) subjects.

Results: In pNT1, RDS score decreased significantly (mean 10.2, SD 4.7 vs mean 6.7, SD 4.5, p < 0.001). There were no significant differences in other sleep scores. However, deviation and range in questionnaire scores at the follow-up were wide. Subjects with very low or undetectable hypocretin-1 levels had worse scores in UNS (mean 26.4, SD 6.95 vs mean 19.1, SD 3.83, p = 0.006) and ESS (mean 17.9, SD = 4.29 vs mean 14.1, SD = 3.70, p = 0.047) than those with hypocretin-1 levels of 20–110 pg/mL. Most disabling symptoms were excessive daytime sleepiness and disturbed sleep. There were no significant differences between the scores in pNT1 and NT1.

Conclusions: Clinical course of pNT1 is heterogeneous but the evolution of pNT1 seems similar to NT1. Lower hypocretin levels in pNT1 are associated with a more severe phenotype.

© 2015 Elsevier B.V. All rights reserved.

1. Introduction

Narcolepsy type 1 is a central disorder of hypersomnolence characterized by a loss of hypocretin-1 (orexin-) producing neurons in the hypothalamus. According to the current hypothesis, narcolepsy type 1 is an autoimmune disease, although the specific mechanism remains elusive [1–3]. The main symptoms of narcolepsy are excessive daytime sleepiness, disturbed nocturnal sleep, cataplexy, sleep paralyses, and hallucinations during sleep–wake transitions. Other common associated features are increased weight, precocious puberty, behavioral problems, and psychiatric comorbidity [4–6]. There is a great heterogeneity in symptoms as some subjects are severely affected and disabled, whereas others manage without medication.

The incidence of narcolepsy increased significantly in countries where AS03-adjuvanted Pandemrix H1N1 vaccination was used widely in 2009–2010 [5,7–11]. The increase was first noted in children and adolescents and later, to a lesser degree, in adults. However, as previously reported, diagnostic delay in narcolepsy has been rather long, over 10 years in many cases, although a shorter delay, especially in younger age groups, has also been observed [12,13]. In Pandemrix H1N1-vaccine-related narcolepsy type 1 (pNT1) cases, the diagnostic delay has been much shorter. Reasons for this remain somewhat unclear. A more sudden onset and severe symptoms in pNT1 than in sporadic narcolepsy type 1 (NT1) could be an explaining factor [5,6]. On the other hand, a comparison between Finnish children with pNT1 and Italian children with NT1 appears to show similar clinical pictures, although subjects with pNT1 often had increased nocturnal sleep disturbance [14]. In a recent study on polysomnographic and actigraphic characteristics of pNT1 and NT1, we found that NT1 subjects had delayed sleep–wake rhythms compared to pNT1, but otherwise we did not see dramatic differences [15].

As pNT1 is considered a drug-adverse effect, the countries where Pandemrix was used are now paying monetary compensations to the affected patients. Compensation policy varies. Considering the heterogeneous disease course as described above, determining the long-term (or even short-term) handicap caused by narcolepsy is...
challenging, and the suitable monetary compensation is difficult to determine if it is based on individual consideration. Nonetheless, narcolepsy causes marked social and economical burden for patients and their close relatives, in addition to deleterious effects which may occur from health-related reductions in quality of life [16,17].

The loss of hypocretin might not be the only biological factor contributing to the symptoms of narcolepsy. Different compensatory mechanisms for the loss of function of hypocretin – such as increase in histamine neurons – are probably also involved [18–21]. Furthermore, we have recently found that narcolepsy patients have autoantibodies that stain distinct cell populations in rat brain, namely melanin-concentrating hormone (MCH), pro-opiomelanocortin (POMC), and neuropeptide glutamic acid-isoleucine/alpha-melanocyte-stimulating hormone (NEI/oMSh) [22]. In addition, MCH/POMC injections in rat brains were found to be associated with disturbed sleep patterns. Recently, Heier et al. also reported changes in nerve cell biomarkers in cerebrospinal fluid (CSF) of narcoleptic patients [23]. These biological alterations, shortened diagnostic delay, abrupt onset, and social and economical factors make it important to follow and analyze the clinical course of pNT1.

2. Methods

2.1. Subjects

Twenty-six pNT1 subjects (15 male, 11 female) from Helsinki Sleep Clinic, Vitalmed Research Centre were recruited for this study. They completed a self-administered modified Basic Nordic Sleep Questionnaire (mBNSQ) near the onset of the disease and at the follow-up visit at least two years later. Follow-up results were also compared with 25 (10 male, 15 female) NT1 subjects. All subjects had confirmed type 1 narcolepsy. Hypocretin-1 was measured in 21 subjects in pNT1 group and in 18 subjects in NT1 and was below 110 pg/mL in all cases. When hypocretin-1 was not measured, the subjects had unambiguous cataplexy, were HLA DQB1*06:02 positive, and had a positive multiple sleep latency test (MSLT) (sleep latency less than eight minutes and two or more sleep onset REM sleep periods [SOREMPs]). Hypocretin levels were measured in Rinnekoti Research Laboratory using Human orexin-A RIA Kit (Phoenix Pharmaceutical, Inc., Belmont, CA) with Stanford reference sample. Characteristics of study subjects are listed in Table 1. Classification as pNT1 was done if the subject had had confirmed Pandemrix vaccination during winter 2009–2010 and symptoms of narcolepsy started within 2 years after vaccination. Most sporadic narcolepsy type 1 subjects had their disease onset before a(H1N1)pdm09 pandemic. Four subjects had their onset during 2009 or 2010. None of them were vaccinated before symptom onset or had influenza like illness (ILI). Ten were vaccinated with Pandemrix after the disease onset. Vaccination data was missing for three subjects. They all had the disease onset years before vaccination.

Subjects were drug-naive at their first visit, ie, they did not have any narcolepsy medication including sodium oxybate, stimulants, selective serotonin, or serotonin and norepinephrine reuptake inhibitors. The treating neurologist/sleep expert made following treatment decisions independently of this study.

2.2. Questionnaires and measures

mBNSQ includes multiple sub-scales and standardized tools for assessment of sleep-related symptoms [24]. Questionnaires used in this study were Epworth Sleepiness Scale (ESS), Ullanlinna Narcolepsy Scale (UNS), Swiss Narcolepsy Scale (SNS), WHO-5 Well-Being Scale (WHOS), and Rimon’s Brief Depression Scale (RDS) [24–27]. A separate questionnaire for disability caused by the disease was also used. This questionnaire included main symptoms of narcolepsy and disability caused by them on scale zero (subject does not have this symptom) to four (severe disability). Weight and height were asked to calculate body mass index (BMI). Number of cataplectic attacks per week was asked (CPL/week). If questionnaires were partially completed, one follow-up phone call was made or a new questionnaire was sent.

2.3. Response rate

Initial complete response rate for all questionnaires combined was 61% and partial response rate was 89% (each measure or questionnaire analyzed separately). After follow-up phone calls or new questionnaires, response rates were 81% and 96%, respectively.

2.4. Statistical analyses

Statistical analyses were performed using SPSS (IBM SPSS® Statistics 19.0, Armonk, NY, USA) and STATA version 13.1 (Stata Corporation, TX). Graphs were made using R software (R development core team, 2014) and ggplot2 package (H. Wickham. ggplot2: elegant graphics for data analysis, Springer New York, 2009) or Excel. For statistical comparisons, parametric and non-parametric methods were used according to the normality of distributions, which were analyzed using Kolmogorov–Smirnov and Shapiro–Wilk tests. Mean (M) and standard deviation (SD) are reported for variables following normal distribution and median (Mdn) and range for non-parametrically distributed variables.

<table>
<thead>
<tr>
<th>Table 1 Characteristics of study subjects.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>pNT1</strong></td>
</tr>
<tr>
<td>Female</td>
</tr>
<tr>
<td>Age at onset, years, median (range)</td>
</tr>
<tr>
<td>Age at diagnosis, years, median (range)</td>
</tr>
<tr>
<td>Diagnostic delay, years, mean (SD)</td>
</tr>
<tr>
<td>Age at follow-up visit, mean (SD)</td>
</tr>
<tr>
<td>Disease duration at follow-up visit, years, median, (range)</td>
</tr>
<tr>
<td>Vaccination to disease onset, days, median, (range)</td>
</tr>
<tr>
<td>Hypocretin-1, pg/mL, median (range)</td>
</tr>
<tr>
<td>MSLT sleep latency, minutes, median (range)</td>
</tr>
<tr>
<td>SOREMPs, median (range)</td>
</tr>
</tbody>
</table>

Abbreviations: pNT1, Pandemrix related type 1 narcolepsy; NT1 sporadic type 1 narcolepsy; MSLT, Multiple Sleep Latency Test; SOREMPs, Sleep Onset REM-sleep Periods in MSLT; NA, Not applicable; Hypocretin-1 measured in 19/24 Pandemrix H1N1-vaccine-related subjects and in 15/23 sporadic cases.

* Mann-Whitney U, p < 0.05; ** Mann-Whitney U, p < 0.005; † Independent samples t-test p < 0.005.
2.5. Ethics

The study was approved by the Helsinki and Uusimaa hospital district ethical committee. Written informed consent was obtained from adult subjects and in children and adolescents from themselves and their parents.

3. Results

Age at onset of narcolepsy was similar in pNT1 and NT1 (Table 1). Diagnostic delay was significantly shorter in pNT1. The follow-up visit was done earlier in the disease course in pNT1 than in NT1 and, therefore, pNT1 subjects were younger at the follow-up. There were no significant differences in CSF hypocretin-1 levels or median sleep latencies in MSLT between the groups.

3.1. Follow-up of pNT1

A paired-samples t-test was conducted to compare questionnaire results between the two visits in pNT1 (Table 2). In general, mean differences in scores were rather low: ESS −0.63 points, UNS −1.68 points, SNS 7.88 points, and WHO5 2.54 points. Mean difference in RDS was an exception as the reduction was 3.44 points and was the only statistically significant change, \( t(24) = 0.381, p = 0.003 \), \( d = 0.67 \). However, standard deviations in changes in scores were rather wide which implies that there were patients who were clearly worse or had their symptoms markedly ameliorated on follow-up visit (Fig. 1). Wilcoxon Signed-ranks test was used to compare differences in BMI and CPL/week. Median of BMI increased 2.3 kg m\(^{-2}\), \( Z = 3.8, p < 0.001 \). Amount of weekly cataplexies did not change significantly.

3.2. Effect of CSF hypocretin levels and MSLT

Linear regression analysis was conducted to find out whether CSF hypocretin levels, sleep latency in MSLT, or number of SOREMPs in MSLT explained changes in sleep questionnaire scores. We did not find any model based on these variables that would explain the changes (data not shown). We then divided study subjects into two subgroups based on hypocretin levels: those with very low (<20 pg/mL) or undetectable hypocretin levels (\( N = 13 \) in pNT1 and \( N = 9 \) in NT1) and those with hypocretin levels of 20–110 pg/mL (\( N = 8 \) in pNT1 and \( N = 9 \) in NT1). A cut-off limit of 20 was used because the assay is not absolutely accurate below 20 pg/mL, although we had specific numeric values also for these results. Here, in pNT1 group, subjects who had very low hypocretin levels had higher UNS scores (\( M = 24.4, SD = 6.64 \)) than subjects with higher hypocretin levels (\( M = 18.8, SD = 4.46 \), \( t(19) = -3.1, p = 0.049 \) (Fig. 2)). They also had higher ESS scores (\( M = 17.2, SD = 4.64 \) vs \( M = 13.1, SD = 1.97 \). \( t(18) = -2.1, p = 0.040 \) (Fig. 3)). Furthermore, diagnostic delay was shorter in patients with very low hypocretin (\( M = 207 \) days, SD = 161) compared with hypocretin levels of 20–110 pg/mL (\( M = 803 \) days, SD = 257), \( t(17) = -6.2, p < 0.005 \). We did not see differences in scores at the first visit or mean differences between visits by hypocretin levels. Neither did we find these differences in NT1 (data not shown).

We also analyzed the relationship between separate UNS items and hypocretin status. Those pNT1 subjects who had hypocretin-1 below 20 pg/mL experienced head nods more often than those with higher levels (\( \text{Md} = 4 \) [daily or almost daily], \( M = 3.18, 95\% \text{ CI 2.24–4.12 vs Md} = 1.5 \) [monthly to less than monthly], \( M = 1.63, 95\% \text{ CI 0.29–2.96, } p = 0.033 \) on the follow-up visit. There were no differences in other UNS cataplexy items (knee sagging, mouth opening or falling to the ground).

Table 2

| Symptoms of Pandemrix H1N1-vaccine-related narcolepsy at first and follow-up visits. |
|----------------------------------------|-----------------|------------------|------------------|------------------|
| First visit                           | Second visit    | 95% CI for Mean difference | \( p \)       |
| ESS, mean (SD)                        | 15.8 (6.2)      | 14.9 (5.0)       | −3.3, 2.0        | 0.631            |
| UNS, mean (SD)                        | 23.3 (7.3)      | 21.1 (6.9)       | −5.3, 1.9        | 0.352            |
| SNS, mean (SD)                        | −40.4 (38.3)    | −32.0 (29.7)     | −9.8, 25.6       | 0.368            |
| RDS, mean (SD)                        | 10.2 (4.7)      | 6.7 (4.5)        | −5.5, −1.3       | 0.003            |
| WHO5, mean (SD)                       | 45.5 (24.8)     | 48.0 (19.3)      | −10.4, 15.4      | 0.008            |
| CPL / week, median (range)            | 15 (0–210)      | 10.5 (0–210)     | NA               | 0.281            |
| BMI, kgm−2, median (range)            | 20.8 (14.4–35.1)| 23.4 (16.6–38.3) | NA               | <0.001           |

Mean (SD) and 95% CI for Mean Difference is reported for parametric and median (range) for nonparametric variables. Paired samples t-test was used to compare parametric and Wilcoxon signed rank test to compare nonparametric variables between different visits in pNT1.

Abbreviations: ESS, Epworth Sleepiness Scale; UNS, Ullanlinna Narcolepsy Scale; SNS, SNS Narcolepsy Scale; RDS, Rimon’s Brief Depression Scale; WHO5, WHO-5 Well Being Scale; BMI, Body Mass Index; CPL/week, cataplexies per week; NA, Not applicable (nonparametric variables). Statistically significant differences are written in bold.

---

Fig. 1. Changes in Ullanlinna Narcolepsy Scale (a) and WHO-5 Well-Being index (b) between visits in Pandemrix related narcolepsy. Dotted line, Ullanlinna Narcolepsy Scale cut point of 14 points; solid line with dots, worse or no change; dotted line with triangles: better.
Higher UNS and ESS scores on the follow-up visit in subjects with unetectable or very low hypocretin levels compared to those who had a higher amount of hypocretin in CSF is intriguing. This implies that severity of hypocretin deficiency can have an effect on the course of type 1 post-Pandemrix narcolepsy is very variable. Our study is the first longitudinal study of patients with NT1 according to the new diagnostic classification. Our results show that the clinical course of type 1 post-Pandemrix narcolepsy is very variable. Symptoms of most of our patients had ameliorated at least a little at the follow-up visit, but there were few patients who were clearly worse. This seems to be in line with previous reports in NT1 [28].

We did not see any differences in mean sleep scale scores between pNT1 a few years after the disease onset and NT1 who were, however, examined later. This finding suggests that the phenotype of these diseases and their clinical course is similar. Therefore, it might be possible to estimate prognosis of pNT1 based on prior knowledge on evolution of NT1. The finding is important especially in pNT1 compensation issues. For example, a clinician can be inquired about the employment possibilities and disability caused by narcolepsy when an adolescent narcoleptic patient turns 18 years old, or an adult patient seeks disability pension or reimbursement from vaccination adverse effect.

The classic tetrad of narcolepsy symptoms includes excessive daytime sleepiness, cataplexy, hypnagogic and hypnopompic hallucinations, and sleep paralysis. As showed in our study, disturbed nocturnal sleep is also a common and clearly disabling symptom. It is known that treatment of disturbed nocturnal sleep (by, eg, sodium oxybate or melatonin when appropriate) may also improve daytime alertness and other daytime symptoms [29].

Higher UNS and ESS scores on the follow-up visit in subjects with undetectable or very low hypocretin levels compared to those who had a higher amount of hypocretin in CSF is intriguing. This implies that severity of hypocretin deficiency can have an effect on the final diagnosis of the case...
disease course. Previously, Baumann and coworkers did find more severe MSLT results in patients with undetectable hypocretin levels compared to patients with low, intermediate, or normal hypocretin levels [30]. Conversely, they did not find differences in the same sleep questionnaire scores (UNS, SNS, ESS) that we had used, but their hypocretin detection limit was 60 pg/mL while in our study it was 20 pg/mL. It is worth noting that our study demonstrated that there was no strict direct relationship between clinical course or symptom severity and CSF hypocretin levels or MSLT findings. For example, we had patients who had very mild symptoms but still undetectable hypocretin levels. Furthermore, those patients whose hypocretin levels were near 110 pg/mL could have severe symptoms as well.

Lack of direct correlation of hypocretin levels on symptoms implicates also that other neural networks than solely hypocretin are contributing to symptom severity in narcolepsy. Possibilities are, eg, NEI/αMSH and histamine [18,20–22]. It has been shown that there is an inverse relationship between dynamics in histamine and hypocretin levels [19]. Conversely, hypocretin neurons in the perifornical hypothalamus of rodents strongly excite histaminergic tuberomamillary nucleus neurons through hypocretin-2 receptors. In patients with narcolepsy, histamine may oppose the effects of hypocretin loss. Accordingly, one promising group of medications for narcolepsy are histamine H3 receptor inverse agonists such as pitolisant [31]. The patients with the worst symptoms may be those whose histamine system, in addition to the hypocretin system, is also defective or not compensating sufficiently.

It has been previously reported that obesity is very common in narcolepsy [4]. BMI increase was clear also in our study while the majority of the subjects still stayed within the normal range. BMI was even higher in NT1 which may also be explained by greater age and longer disease duration of NT1 subjects. Still, reasons for the weight increase remain unknown. It may be due to changes in calorie uptake or expenditure, medication side-effects, or altered metabolic status. As some narcolepsy medications are weight increasing (eg, venlafaxine, clomipramine) and some weight decreasing (sodium oxybate), this is an important aspect to take into account, when considering treatment options.

Somewhat surprisingly, we did not see differences in questionnaire scores between adults and younger subjects. Some previous reports have suggested that childhood narcolepsy presents a distinct clinical picture and with specific symptoms such as facial or generalized hypotonia and increased total sleep time, that gradually ameliorate during the following years [28]. Our questionnaire might not be sensitive enough to separate these symptoms as they are not designed and validated for such purpose. Our adult cases might also have more severe disease than previous sporadic narcolepsy patients in general, since their diagnostic delay has been

---

**Table 3**

<table>
<thead>
<tr>
<th>Medication used on follow-up visit.</th>
<th>No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Modafinil</td>
<td>16</td>
</tr>
<tr>
<td>Sodium oxybate</td>
<td>12</td>
</tr>
<tr>
<td>Methylphenidate</td>
<td>3</td>
</tr>
<tr>
<td>Venlafaxine</td>
<td>3</td>
</tr>
<tr>
<td>Clomipramine</td>
<td>2</td>
</tr>
<tr>
<td>MAO-B-inhibitors</td>
<td>1</td>
</tr>
<tr>
<td>Other narcolepsy medications (eg, SSRI)</td>
<td>3</td>
</tr>
<tr>
<td>Other medications (incl. melatonin)</td>
<td>8</td>
</tr>
<tr>
<td>No medication</td>
<td>2</td>
</tr>
</tbody>
</table>

Abbreviations: MAO-B, monoamineoxidase; SSRI, selective serotonin reuptake inhibitor.
rather short. Previously, diagnostic delay of more than 10 years has been seen. Type 2 error is also possible due to our relatively small sample size.

Our study has some other limitations as well. The study was conducted in a tertiary sleep clinic, which may cause selection bias. Our study population might be more severely affected than most narcoleptic patients as some patients with mild symptoms may have visited our clinic only for diagnostic reasons and the follow-up might have been done in secondary or primary healthcare institutions. Frequent complaints about medication in our study could support this hypothesis. However, we also had some patients without medication on the follow-up visit. Despite these limitations, we are confident that these results can be generalized to pNT1.

Unfortunately, we were not able to perform hypocretin measurements for all the patients, but their diagnosis was still solid and they were most likely hypocretin deficient. They were all HLA-DQB1*06:02 positive and they all had unambiguous cataplexy and two or more SOREMPs and sleep latency of less than 8 min in MSLT. Theoretically, it could have been interesting to also follow hypocretin levels on second lumbar puncture, but as it would not have had any effect on treatment, we decided not to do it for ethical reasons. Sleep recording follow-up would have been interesting but was not financially possible and not real-life practice either. Moreover, patient-reported outcomes are most important in following the treatment effect and symptoms of narcolepsy in real-life settings [32].

In summary, the clinical course of post-Pandemrix narcolepsy is very heterogeneous. Most patients get at least somewhat better at two years’ follow up. Improvement in depression scale points could reflect adaptation to disease symptoms. Weight gain is common also in Pandemrix H1N1-vaccine-related narcolepsy. MSLT findings do not correlate very well with symptom severity but lower hypocretin levels are associated with a more severe phenotype on follow-up visit. Therefore, the severity of hypocretin deficiency can have some effect on clinical course of narcolepsy. There is no marked difference in symptoms between sporadic or Pandemrix-associated narcolepsy after two or more years after disease onset.

Conflict of interest

Dr Sarkanen reports grants from Academy of Finland, during the conduct of the study; non-financial support from UCB Pharma, non-financial support from Bioprojet, outside the submitted work. Dr Partinen reports grants from Academy of Finland, during the conduct of the study; personal fees from UCB Pharma, personal fees from Leiras Takeda, personal fees from MSD, personal fees from Cephalon, outside the submitted work. Dr Alakuijala reports grants from Academy of Finland, during the conduct of the study.

The ICMJE Uniform Disclosure Form for Potential Conflicts of Interest associated with this article can be viewed by clicking on the following link: http://dx.doi.org/10.1016/j.sleep.2015.11.005.

Acknowledgements

We are grateful to all of our patients and their parents who participated in the study.

We thank Anne Haustioniemi, Raili Reemets and all study personnel for their valuable help in organizing and collecting data. We thank Eric Burns for language editing. This study is supported by Academy of Finland grant 260603 (NarpaNord).

References

[18] Conflict of interest

Dr Sarkanen reports grants from Academy of Finland, during the conduct of the study; non-financial support from UCB Pharma, non-financial support from Bioprojet, outside the submitted work. Dr Partinen reports grants from Academy of Finland, during the conduct of the study; personal fees from UCB Pharma, personal fees from Leiras Takeda, personal fees from MSD, personal fees from Cephalon, outside the submitted work. Dr Alakuijala reports grants from Academy of Finland, during the conduct of the study.

The ICMJE Uniform Disclosure Form for Potential Conflicts of Interest associated with this article can be viewed by clicking on the following link: http://dx.doi.org/10.1016/j.sleep.2015.11.005.

Acknowledgements

We are grateful to all of our patients and their parents who participated in the study.

We thank Anne Haustioniemi, Raili Reemets and all study personnel for their valuable help in organizing and collecting data. We thank Eric Burns for language editing. This study is supported by Academy of Finland grant 260603 (NarpaNord).

References

[18] Conflict of interest

Dr Sarkanen reports grants from Academy of Finland, during the conduct of the study; non-financial support from UCB Pharma, non-financial support from Bioprojet, outside the submitted work. Dr Partinen reports grants from Academy of Finland, during the conduct of the study; personal fees from UCB Pharma, personal fees from Leiras Takeda, personal fees from MSD, personal fees from Cephalon, outside the submitted work. Dr Alakuijala reports grants from Academy of Finland, during the conduct of the study.

The ICMJE Uniform Disclosure Form for Potential Conflicts of Interest associated with this article can be viewed by clicking on the following link: http://dx.doi.org/10.1016/j.sleep.2015.11.005.

Acknowledgements

We are grateful to all of our patients and their parents who participated in the study.

We thank Anne Haustioniemi, Raili Reemets and all study personnel for their valuable help in organizing and collecting data. We thank Eric Burns for language editing. This study is supported by Academy of Finland grant 260603 (NarpaNord).