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### Melatonin treatment and light therapy for chronic sleep onset insomnia in children

*Effects on sleep, cognition, health, and psychosocial functioning*

van Maanen, A.

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## CHAPTER 4

# Classical conditioning for preserving effects of short term melatonin treatment in children with chronic sleep onset insomnia

*A pilot study*

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### **THIS CHAPTER IS BASED ON:**

Van Maanen, A., Meijer, A. M., Smits, M. G., & Oort, F. J. (in press). Classical conditioning for preserving effects of short melatonin treatment in children with delayed sleep: A pilot study.

**Nature and Science of Sleep**

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## ABSTRACT

*Melatonin treatment is effective in treating sleep onset problems in children with chronic sleep onset insomnia, but effects usually disappear when treatment is discontinued. In this pilot study, we investigated whether classical conditioning might help preserving treatment effects of melatonin in children with sleep onset problems, with and without comorbid ADHD or autism.*

*After a baseline week, 16 children (mean age 9.78 years, 31% ADHD / autism) received melatonin treatment for three weeks, and then gradually discontinued treatment. Classical conditioning was applied by having children drink biological lemonade while taking melatonin, and by using a lamp that was turned on when children went to bed. Results were compared with a group of 41 children (mean age 9.43 years, 34% ADHD / autism) who received melatonin without classical conditioning.*

*Melatonin treatment was effective in advancing Dim Light Melatonin Onset and reducing sleep onset problems, and positive effects were found on health and behaviour problems. After stopping melatonin, sleep returned to baseline levels. We found that for children without comorbidity in the experimental group sleep latency and sleep start delayed less in the stop week, which suggests an effect of classical conditioning. However, classical conditioning seems counterproductive in children with ADHD or autism. Further research is needed to establish these results.*

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## INTRODUCTION

**S**leep problems are common among children, with reported prevalence rates as high as 30-40%.<sup>1,2</sup> Considering these high prevalence rates and the negative consequences of sleep problems on mood, behaviour and school performance,<sup>3,4</sup> it is important to treat these problems. Moreover, sleep problems in childhood predict later behavioural and emotional problems,<sup>5,6</sup> which stresses the importance of early intervention.

Sleep problems can have different manifestations and causes requiring different types of treatment. In the present study, we focus on children with chronic sleep onset problems related to a delayed melatonin onset. Melatonin is a hormone produced by the pineal gland in the brain. Melatonin secretion starts to rise in the evening and increases sleep propensity.<sup>7</sup> In some children, melatonin secretion is delayed, which is associated with falling asleep late and difficulty awakening in the morning. Exogenous melatonin, if well-timed and given in the appropriate dose, may advance sleep onset and offset.<sup>8</sup>

Several studies indeed found positive effects of melatonin treatment on sleep onset problems in children,<sup>9-11</sup> and also on health,<sup>10</sup> behaviour problems and parenting stress.<sup>12</sup> This was also found for children with ADHD and autism,<sup>13,14</sup> disorders with a high comorbidity with sleep problems.<sup>15-18</sup> However, previous research showed that the positive effects of melatonin disappeared after discontinuing short-term treatment,<sup>12</sup> indicating that treatment should be continued for a longer period of time. Yet unnecessary long-term treatment is not desirable considering that there is a (theoretical) risk of delayed puberty onset associated with long term melatonin use, although several longitudinal studies did not find any deviations when comparing children who used melatonin for a long time compared to the general population.<sup>19-21</sup>

The question then arises how the positive effects of melatonin treatment can be preserved when treatment is discontinued after a short period. A suggestion comes from research on classical conditioning. Classical conditioning is a learning principle originally described by Pavlov.<sup>22</sup> He showed that, after coupling a neutral stimulus (e.g., a tone) to a stimulus that naturally evokes a specific response (stimulus: food, response: salivation), the former neutral stimulus can in itself evoke that response (tone evokes salivation). In the classical conditioning theory, the stimulus that automatically evokes a certain response is called an Unconditioned Stimulus (US). The response that is evoked by the US is called the Unconditioned Response (UR). The former neutral stimulus that leads to the same

response after the learning process, is called the Conditioned Stimulus (CS). The response that is now evoked by the CS is called the Conditioned Response (CR).

This principle has been applied to patients with insomnia.<sup>23,24</sup> Evans and Bond treated a 45-year old man with insomnia with methohexital sodium (US) while asking him to count (CS). The methohexital sodium made the man fall asleep (UR), and after some trials without using methohexital sodium but still counting (CS), the man reported that his sleep had improved, suggesting that the counting had become a conditioned stimulus resulting in better sleep (CR). A similar result was found by Poser et al.,<sup>24</sup> who treated a man with insomnia with methohexital injections (US) while listening to the ticking of a metronome (CS). After a number of trials, the man reported clinical improvement when he kept using the metronome. In addition to the studies described above, a recent study<sup>25</sup> discussed the literature on classical conditioning in relation to placebo response. It was found that conditioning, as a 'learned placebo response', can influence peripheral immune functions as well as disease symptoms and disease progress. In the aforementioned studies forward conditioning was used, i.e. the onset of the CS precedes the onset of the US to signal that the US will follow. Generally, the UR starts immediately or shortly after the US.

The results described above suggest that classical conditioning might be successful in preserving the effects of short term melatonin treatment. To investigate this novel idea, we conducted a pilot study with a sample of children with chronic sleep onset insomnia, associated with a delayed melatonin onset. Their sleep behaviour was assessed on 41 consecutive days: before melatonin treatment (1 week), during treatment and conditioning (3 weeks), during half dose treatment (1 week), and after stopping treatment (1 week) (see Figure 1). After baseline, classical conditioning was applied. As it takes some time to become sleepy after melatonin intake, we used two conditioning procedures. Children drank biological lemonade while taking melatonin, and a dim light red lamp was turned on when children went to bed. The daily assessments were compared to the daily assessments of children in a previous study who underwent the same procedure but without the conditioning of melatonin treatment.<sup>12</sup> In both groups, a substantial proportion of children also had ADHD or autism (31% in the experimental group and 34% in the comparison group). In this way we aimed to test the hypothesis that classical conditioning could help to preserve the effects of short-term melatonin treatment in children with chronic sleep onset insomnia related to a delayed melatonin onset, and we tested whether effects were different for children with and without ADHD or autism.

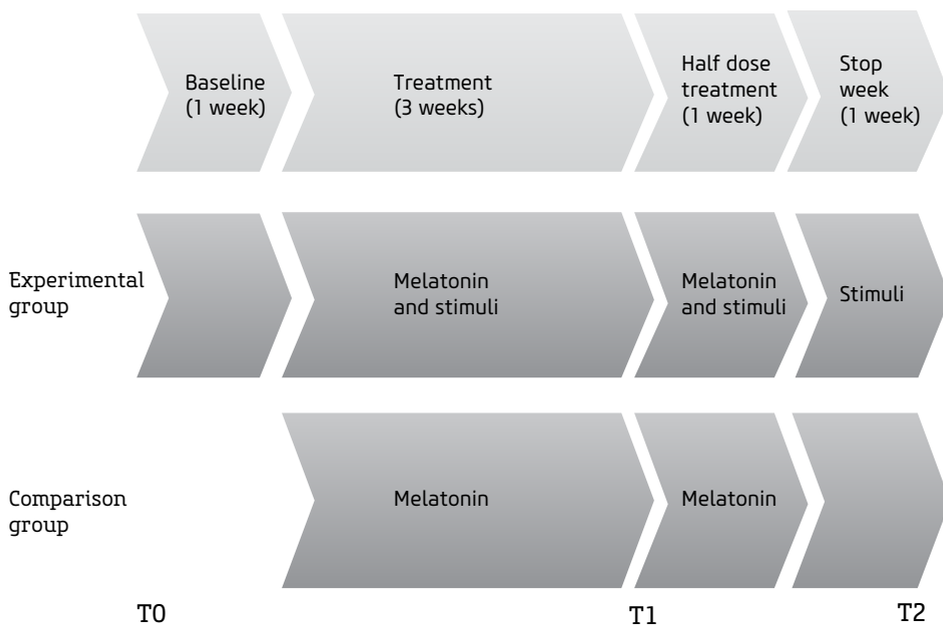


Figure 1. Study design

## METHODS

### Participants

The study was conducted in the Centre for Sleep–Wake Disorders and Chronobiology in the hospital. Children were referred to the hospital by their general practitioners because of their chronic sleep onset problems. Inclusion criteria for participation were (1) age between 5 and 12 years old; (2) the child has chronic sleep onset problems defined as (a) complaints of sleep-onset problems expressed by parents and/or child, (b) occurrence on at least 4 days/week for longer than 1 year, (c) average sleep onset later than 20:15 h for children at age 5 years and for older children 15 min later per year, and (d) average sleep latency exceeding 30 min;<sup>9,10,14</sup> (3) Dim Light Melatonin Onset (DLMO) later than 20:00 h; (4) the child attends a regular school (IQ is in the normal range); and (5) parents of the child have sufficient command of the Dutch language to understand the treatment and to complete the questionnaires. Children were not eligible for participation if (1) the child had a diagnosis of another sleep disorder (e.g., restless legs syndrome, narcolepsy, obstructive sleep apnoea syndrome), and/or (2) the sleep onset problems were caused by physical problems (e.g., pain). See Table 1 for a description of the experimental and comparison group.

Table 1. Descriptive statistics of the experimental and comparison group

	<b>EXPERIMENTAL GROUP (N = 16)</b>	<b>COMPARISON GROUP (N = 41)</b>
	Mean (SD)	Mean (SD)
Age (years)	9.78 (1.67)	9.43 (2.05)
	Frequency (%)	Frequency (%)
Boys	6 (37.50)	24 (58.54)
ADHD	4 (25.00)	8 (19.51)
Autism spectrum disorder (ASD)	0 (0.00)	5 (12.20)
ADHD and ASD	1 (6.25)	1 (2.44)
ADHD medication	4 (25.00)	4 (9.76)
Enuresis	2 (12.50)	0 (0.00)
Complaints of headache	3 (18.75)	0 (0.00)

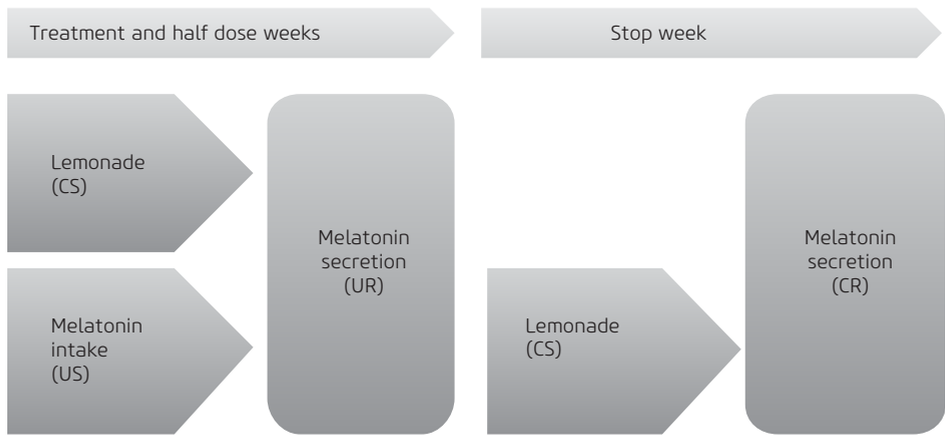
### **Procedure**

The study was approved by the Ethical Committee of the Research Institute of Child Development and Education of the University of Amsterdam. Parents gave active informed consent for participation. Before their first appointment in the hospital, Dim Light Melatonin Onset, the clock time at which the endogenous melatonin secretion reaches a certain threshold, was measured in saliva. Parents were asked to instruct their children to chew on cotton plugs according to a predetermined schedule for one evening.<sup>10,14</sup> In order to make sure that all participants had similar knowledge about sleep hygiene practices, we discussed this topic during the first appointment in the hospital, and we provided written instructions for sleep hygiene. The researcher also explained the procedures of the study, and provided the stimuli for the classical conditioning (see Figure 2).

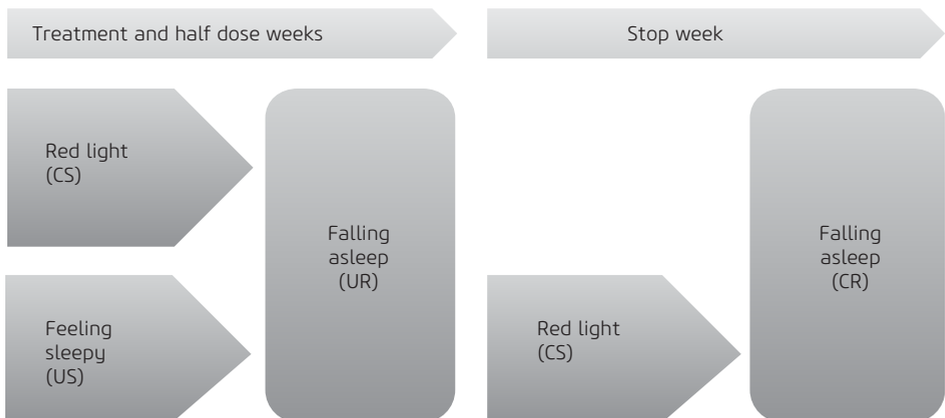
### **Design**

The study started on the first Sunday after the appointment in the hospital, with a baseline week in which parents completed questionnaires and daily sleep diaries, and children wore actometers. After a week, treatment started. All children started with a dose of 1 mg, fast-release melatonin. If parents did not see any effect of melatonin use after four days, they were allowed to increase the dose to 2 mg. If this still had no effect, the dose could be further increased every

## BEFORE BEDTIME



## IN BED



**Figure 2. Classical conditioning procedure**

*Conditioned stimuli were a biological lemonade with peppermint taste and a dim light red-coloured lamp with curly cord. Children were instructed to drink the lemonade (CS) at the moment that they took the melatonin tablet (US) to couple the lemonade to the melatonin secretion (UR) and making the lemonade a CS for melatonin secretion (CR). The red light (CS) was turned on when the child went to bed and felt sleepy (US) to couple the red light to falling asleep (UR) and making it a CS for falling asleep (CR). Parents were instructed to switch the red light off when the child got up in the morning.*

four days, until sleep onset advanced. Maximum dose was 5 mg. The procedure to increase the dose gradually in case of no effect was in accordance with recent clinical recommendations for prescribing melatonin in children, although they advise to start with an even lower dose (0.2-0.5 mg).<sup>8</sup> However, this was not possible in our study, as all children had to respond to treatment within three weeks. Melatonin was administered three hours before DLMO, but not earlier than 19:00 h, according to Keijzer et al.<sup>26</sup>

After three weeks of melatonin treatment, parents filled in the questionnaires and children chewed on the cotton plugs. Then treatment was discontinued by first taking a half dose for one week (hereafter called the “half dose week”) and then stopping completely for another week (hereafter called the “stop week”). After this stop week we repeated the assessments. Parents filled in sleep diaries and children wore actometers during this whole period of six weeks. The stimuli were used during the three treatment weeks, the half dose week and the stop week. Families received reminders through text messages or e-mails on days that questionnaires had to be filled in, children had to chew on cotton plugs, melatonin dose had to be halved or treatment had to be discontinued. In addition, all families were contacted by telephone twice during this study to discuss their experiences: once in the first week of treatment and once after the end of the stop week. Children were allowed to recommence with melatonin after the last day of the stop week.

The study consisted of three measurements (Figure 1): baseline (in the week before the start of treatment; T0), directly after 3 weeks treatment (T1), and at the end of the stop week (T2). At these measurement occasions DLMO was determined in saliva and questionnaires were filled in.

Participants in the previous study<sup>12</sup> served as comparison group. They went through the same procedures, with the same measurements, except for the baseline sleep measurements (sleep diaries and actigraphy) and the use of stimuli for classical conditioning.

## **Measures**

The same variables were measured in both groups, but different questionnaires were used to measure behaviour problems in the experimental and comparison groups.

## **Sleep**

Parents filled in sleep diaries daily via internet. The sleep diary consisted of questions concerning bed time, lights off time, sleep onset time, and questions on

whether the child woke up during the night. Sleep latency (time children spent in bed before falling asleep) and sleep start were used as sleep variables in the analyses.

In addition to sleep diaries sleep was measured with AW4 acti-watches (Cambridge Neurotechnology Ltd, Cambridge, UK). Children were instructed to wear the actigraph on their non-dominant wrist when they went to bed and to remove it in the morning when they got out of bed. The actometers were used to obtain objective information about sleep latency, sleep start, actual sleep time (time children actually slept during the night), and sleep efficiency (actual sleep time/time in bed). Nocturnal activity data were logged at 1-min epochs and scored with the Actiwatch Sleep Analysis 7 (Cambridge Neurotechnology Ltd, Cambridge, UK) software.

### **Dim Light Melatonin Onset**

Dim Light Melatonin Onset, which can be used as an indication of the timing of the biological clock, was measured in saliva using Bühlmann RIA kits (Bühlmann, Schönenbuch, Switzerland). Children chewed on cotton plugs hourly from 19:00 to 23:00 h in the evening at dim light.<sup>10,14</sup> Children were not allowed to use melatonin in the evenings DLMO was measured.

### **Health**

The health status of the children was measured with the first part of the Functional Status II (FSII).<sup>27,28</sup> This first part consists of 14 items concerning activities and behaviours in the past 2 weeks. An example of an item is: "Did your child seem to feel sick and tired?" Parents had to indicate how often these behaviours or activities occurred on a three-point scale varying from (0) "never or rarely" to (1) "some of the time" and (2) "almost always". A higher score indicates a better functional status and consequently a better health. Cronbach's alpha varied between .80 and .93 at the different measurement occasions.

### **Parenting stress**

Parenting stress was measured with the Nijmegen Parental Stress Index short version (NOSIK).<sup>29</sup> The NOSIK is a questionnaire with 17 items that measures to what extent parents experience stress in parenting their child. An example of an item is: "My child often demands more attention than I can give". Parents answered the items on a 4-point scale, ranging from (1) "strongly disagree" to (4) "strongly agree". A higher score indicated more parenting stress. Reliability varied from .93 to .96.

## **Behaviour problems**

Behaviour problems were operationalised differently in the experimental and comparison group, and could thus not be compared across groups.

In the experimental group, behaviour problems in children were measured with the Strength and Difficulties Questionnaire (SDQ).<sup>30,31</sup> This questionnaire has 25 items asking about various child behaviours, for example “Often loses temper”. Response options are (1) “not true”, (2) “somewhat true”, (3) “certainly true”. The following subscales are distinguished: emotional symptoms, conduct problems, hyperactivity/ inattention, peer relationship problems, and prosocial behaviour. The first four scales together yield a total problem behaviour score. Cronbach’s alpha varied between .88 and .91 at the different measurement occasions.

In the comparison group, behaviour problems were measured with the Child Behavior Checklist (CBCL).<sup>32,33</sup> The CBCL is a comprehensive (112 items) questionnaire. An example of an item is: “Temper tantrums or hot temper”. The response scale ranged from (0) “not true” to (2) “very true or often true”, with a higher score indicating more behaviour problems. The reliability varied from .95 to .96 at the different measurement occasions.

## **Statistical analysis**

Data were analysed using linear mixed models in SPSS, treating the repeated observations as nested within children. In this way all available data were used to answer the research questions, including data from children with missing observations. For the sleep variables, it was first determined which longitudinal structure best described the variances and covariances of the 41 daily assessments.<sup>34</sup> For all sleep variables a first-order autoregressive covariance structure was chosen. For the questionnaire data that were collected at three measurement occasions, free covariance structures were used.

Outcome variables were sleep variables measured during the different phases of the treatment (baseline week, treatment weeks, half dose week, stop week), and DLMO and sum scores for the different questionnaires measured at T0 through T2. In the sleep data analyses, treatment days for which parents reported that no melatonin was taken (forgotten or on purpose) were omitted from the analyses. In addition, in several cases children used melatonin before the end of the stop week. These days were also omitted as to prevent distortion of the effects.

Changes in sleep were tested by including main effects of treatment phases as explanatory variables. To investigate whether treatment effects were different for the experimental versus comparison group, we tested interaction effects between treatment phases and group status. A significant interaction

effect of half dose or stop week with group, would indicate an effect of the classical conditioning. We also added binary indicators for group, ADHD / autism and weekend (as a control variable). We included interaction effects between treatment phases and ADHD / autism, group and ADHD / autism, and higher order interaction effects between ADHD / autism, group and treatment phases. A significant higher order interaction would mean that classical conditioning has differential effects for children with or without ADHD / autism.

For DLMO and questionnaire data we included main effects of measurement occasions, group status, and ADHD / autism. We also included interaction effects between measurement occasions, group and ADHD / autism, and higher order interaction effects between ADHD / autism, group and measurement occasions. A significant interaction effect of stop week and group would indicate an effect of classical conditioning, and a significant higher order interaction would indicate that there are different effects of classical conditioning for children with ADHD and / or autism.

## RESULTS

### **Dose of melatonin and time of administration**

All participants took melatonin at 19:00 h. The majority of children took 1 mg melatonin (66.7% in the experimental group and 61% in the comparison group). Mean (SD) dose in the experimental group was 1.6 (1.1) mg, and 1.5 (1.0) mg in the comparison group.

### **Sleep outcomes**

Means and standard deviations of all variables for the different measurement occasions are reported in Table 2. Average sleep scores in different treatment phases by group and ADHD / autism diagnosis are shown in Figure 3.

### **Sleep diary**

Results of the linear mixed models analyses showed that sleep latency reduced during melatonin treatment, as latency was significantly longer during baseline ( $\beta = .45$ ,  $p < .01$ ; see Table 3). In the half dose week, sleep latency increased compared to treatment ( $\beta = .13$ ,  $p = .02$ ), but not for children with ADHD / autism ( $\beta = -.24$ ,  $p = .01$ ). During the stop week sleep latency was longer compared to treatment ( $\beta = .58$ ,  $p < .01$ ), but the increase was less for children in the experimental group ( $\beta = -.25$ ,  $p = .03$ ). The latter result indicates an effect of classical conditioning. However, this was only true for children without ADHD/

autism, as a significant higher order interaction indicated that for children with ADHD / autism in the experimental group sleep latency increased more in the stop week, so for them classical conditioning was counterproductive ( $\beta = .62, p < .01$ ). In addition, we found a main effect of weekend ( $\beta = -.09, p < .01$ ), indicating that sleep latency was shorter during weekends. A significant interaction effect between group and ADHD / autism ( $\beta = .62, p = .01$ ) indicated that children with ADHD / autism in the experimental group had longer sleep latencies during treatment. See Figure 3a for an illustration of the effects.

Sleep start was significantly earlier during melatonin treatment compared to baseline. During the half dose week and stop week, sleep start was later than during melatonin treatment, although for children with ADHD / autism in the comparison group sleep start did not increase during half dose week and the increase in the stop week was less strong. There was an effect of classical conditioning, as the interaction effect of group and stop week showed that for the experimental group sleep start delayed less in the stop week. However, a higher order interaction indicated that for children with ADHD / autism in the experimental group sleep start increased in the stop week, which suggests that for these children classical conditioning was counterproductive (see Table 3 and Figure 3b). In addition, there was a main effect of weekend, indicating that sleep start was later during weekends.

### **Actigraphy**

Sleep latency was significantly reduced during melatonin treatment compared to baseline, and this was still true during the half dose week. During the stop week, sleep latency was significantly longer than during melatonin treatment. There were no significant interaction effects between group and treatment phases, indicating that the treatment effects were not different for the two groups and there was no effect of classical conditioning. However, there was a significant higher order interaction effect, indicating that for children with ADHD / autism in the experimental group sleep latency increased more in the stop week (see Figure 3c).

Sleep start was significantly earlier during melatonin treatment compared to baseline. During the half dose week and the stop week, sleep start was significantly later than during melatonin treatment. For the experimental group sleep start delayed less in the stop week (which shows an effect of classical conditioning), but for children with ADHD / autism in the experimental group sleep start increased more in the stop week (see Figure 3d). There were main effects for ADHD / autism and weekend, indicating that in the comparison group children with ADHD / autism had an earlier sleep start during treatment, and that sleep

start was later during weekends. Also, children with ADHD / autism in the experimental group fell asleep later during treatment.

Actual sleep time was longer during melatonin treatment compared to baseline. During the stop week actual sleep time was shorter than during melatonin treatment. For actual sleep time, no effect of classical conditioning was found (see Figure 3e).

Sleep efficiency was higher during melatonin treatment compared to baseline. During the stop week, sleep efficiency was lower than during melatonin treatment, but for children with ADHD / autism in the comparison group the decrease in sleep efficiency in the stop week was less strong. No effect of classical conditioning was found, but the significant higher order interaction showed that for children with ADHD / autism in the experimental group sleep efficiency decreased more in the stop week (see Table 3 and Figure 3f). Sleep efficiency generally was lower during the weekends.

### **Other variables**

DLMO significantly advanced during melatonin treatment, but this was no longer the case after the stop week. There were no significant interaction effects of group with measurement occasions. Health improved after melatonin treatment, but this positive effect disappeared after the stop week. A significant interaction effect indicated that children with ADHD / autism in the experimental group had worse health scores. All participants scored higher on health after treatment, especially the experimental group. Parenting stress did not significantly change, but was higher for parents of a child with ADHD / autism.

Behaviour problems decreased after melatonin treatment in the experimental group, and this was still true after the stop week. Children with ADHD / autism scored higher on behaviour problems. As different questionnaires were used in the experimental and comparison group, the results could not be compared between the two groups, but in the comparison group the reduction in behaviour problems after treatment also remained after the stop week.<sup>12</sup> See Table 4 for results.

Table 2. Means for outcome variables at the different phases of treatment

Sleep variables	<b>EXPERIMENTAL GROUP (WITH CLASSICAL CONDITIONING, N = 16)</b>			
	Baseline M (SD)	Treatment M (SD)	Half dose M (SD)	Stop week M (SD)
<b>Sleep diary</b>				
Sleep latency <sup>a</sup> (in minutes)	68.17 (51.50)	41.98 (37.47)	48.39 (41.14)	66.13 (54.57)
Sleep start (clock time)	22:21 (1:05)	21:35 (0:58)	21:49 (1:08)	22:18 (1:02)
<b>Actometer</b>				
Sleep latency <sup>a</sup> (in minutes)	61.70 (53.47)	28.69 (26.44)	34.33 (35.22)	67.65 (54.83)
Sleep start (clock time)	22:13 (1:11)	21:26 (0:58)	21:44 (1:08)	22:20 (1:11)
Actual sleep time (in hours)	8.11 (0.88)	8.42 (1.00)	8.42 (1.01)	8.00 (0.93)
Sleep efficiency	75.87 (7.21)	78.28 (7.27)	77.64 (7.50)	74.76 (7.37)
<b>Other variables</b>	<b>T0 (baseline) M (SD)</b>	<b>T1 (end of treatment) M (SD)</b>	<b>T2 (end of stop week) M (SD)</b>	
Dim Light	20:39	20:03	20:43	
Melatonin Onset (clock time)	(1:09)	(1:28)	(1:20)	
Health	17.50 (4.76)	23.29 (4.27)	18.00 (7.19)	
Parenting stress	35.00 (14.40)	29.50 (12.16)	29.92 (11.36)	
Behaviour problems <sup>b</sup>	12.56 (7.90)	9.14 (6.97)	8.85 (7.56)	

<sup>a</sup> Sleep latency: difference between bed time and sleep start. <sup>b</sup> Behaviour problems were measured with different questionnaires in the experimental and comparison groups, and can therefore not be compared.

<b>COMPARISON GROUP</b> <b>(N = 41)</b>		
Treatment M (SD)	Half dose M (SD)	Stop week M (SD)
37.27 (28.47)	38.77 (34.36)	69.39 (46.77)
21:00 (0:59)	21:11 (1:14)	21:43 (1:06)
28.37 (26.91)	28.42 (27.29)	63.65 (43.65)
21:07 (1:07)	21:29 (1:15)	22:08 (1:16)
8.62 (1.06)	8.49 (1.02)	8.13 (1.04)
75.72 (7.31)	75.57 (7.19)	72.34 (7.13)
T0 (baseline) M (SD)	T1 (end of treatment) M (SD)	T2 (end of stop week) M (SD)
20:58 (0:50)	19:31 (0:52)	20:55 (1:09)
20.89 (3.14)	22.50 (3.98)	20.64 (3.98)
31.99 (11.14)	29.65 (9.78)	29.58 (9.61)
40.28 (23.37)		28.91 (21.09)

**Table 3. Differential effects of classical conditioning on sleep variables during different treatment phases**

	<b>SLEEP DIARY</b>					
	Sleep latency			Sleep start		
	$\beta$	se	p	$\beta$	se	p
<b>Treatment phases (within effects)</b>						
Baseline (vs. treatment)	.45	.08	<.01	.79	.11	<.01
Half dose (vs. treatment)	.13	.06	.02	.32	.07	<.01
Stop week (vs. treatment)	.58	.06	<.01	1.00	.08	<.01
Weekend (vs. week days)	-.09	.03	<.01	.45	.04	<.01
<b>Group effects (between effects)</b>						
Group (conditioning vs. comparison)	-.13	.13	.32	.23	.28	.41
ADHD/autism	-.11	.12	.36	-.51	.26	.06
Group $\times$ ADHD/autism	.62	.24	.01	.92	.51	.07
<b>Differential effects of Conditioning and ADHD/autism during treatment phases</b>						
Half dose $\times$ Conditioning	.08	.10	.42	.09	.13	.50
Stop week $\times$ Conditioning	-.25	.11	.03	-.38	.15	.01
Baseline $\times$ ADHD/autism	-.03	.15	.86	.05	.20	.79
Half dose $\times$ ADHD/autism	-.24	.09	.01	-.39	.12	<.01
Stop week $\times$ ADHD/autism	-.13	.10	.21	-.45	.13	<.01
Half dose $\times$ Conditioning $\times$ ADHD/autism	-.09	.19	.65	.08	.25	.75
Stop week $\times$ Conditioning $\times$ ADHD/autism	.62	.22	<.01	1.06	.29	<.01

Notes:  $N = 57$ ;  $\beta$  = regression coefficient, se = standard error,  $p$  =  $p$ -value. Significant effects are indicated in bold.

Regression effects are additive. That is, treatment phase effects are valid for the comparison group without ADHD/autism. The effects of conditioning and ADHD/autism are valid for the treatment phase. In order to obtain the total effects of conditioning and ADHD/autism in the various treatment phases, the interaction effects of the other treatment phases (Baseline, Half dose, Stop week) with conditioning and ADHD/autism have to be added to the aforementioned effects (cf. Figure 3).

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**ACTIGRAPHY**

Sleep latency			Sleep start			Actual sleep time			Sleep efficiency		
$\beta$	se	p	$\beta$	se	p	$\beta$	se	p	$\beta$	se	p
.53	.07	<.01	.83	.13	<.01	-.37	.12	<.01	-2.13	.86	.01
.05	.05	.31	.37	.09	<.01	-.14	.09	.10	-.61	.60	.31
.64	.06	<.01	1.13	.10	<.01	-.55	.10	<.01	-4.43	.67	<.01
-.02	.03	.57	.51	.05	<.01	.10	.05	.06	-.78	.35	.02
-.09	.09	.33	.01	.28	.98	.02	.20	.93	2.56	1.66	.13
.03	.09	.73	-.63	.26	.02	.32	.19	.09	-1.62	1.53	.29
.29	.16	.08	1.01	.50	.05	-.70	.35	.05	.28	2.90	.93
.08	.09	.41	-.03	.16	.87	.04	.15	.80	.10	1.08	.92
-.19	.11	.08	-.37	.18	.04	.13	.18	.47	1.71	1.23	.17
.10	.14	.45	-.09	.23	.69	.16	.22	.48	-.83	1.56	.60
-.12	.09	.16	.15	.14	.29	-.01	.14	.97	1.05	.98	.29
-.10	.09	.30	-.03	.16	.86	.18	.15	.25	2.11	1.07	.05
.02	.16	.88	-.27	.28	.32	.24	.27	.37	-1.45	1.87	.44
.71	.20	<.01	.71	.33	.03	-.17	.33	.60	-6.09	2.27	<.01

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**Table 4. Treatment effects on Dim Light Melatonin Onset (DLMO), sleep quality, health, behaviour problems and parenting stress**

	<b>DLMO</b>		
	$\beta$	SE	p
<b>Measurement occasions (within effects)</b>			
Treatment vs. baseline (T1)	-1.58	.23	<.01
Stop week vs. baseline (T2)	-.19	.23	.42
<b>Group effects (between effects)</b>			
Group (conditioning vs. comparison)	-.28	.32	.39
ADHD / autism	-.43	.29	.14
Group x ADHD / autism	-.17	.59	.78
<b>Differential effects of conditioning and ADHD/autism during treatment phases</b>			
Treatment x Conditioning	.60	.44	.18
Stop week x Conditioning	-.23	.49	.64
Treatment x ADHD / autism	.32	.41	.44
Stop week x ADHD /autism	.22	.41	.59
Treatment x group x ADHD / autism	.80	.83	.34
Stop week x group x ADHD / autism	1.21	.92	.20

*Note:  $\beta$  = regression coefficient, se = standard error, p = p-value. Significant effects are indicated in bold. \* For behaviour problems only results for the experimental group are reported.*

HEALTH			PARENTING STRESS			BEHAVIOUR PROBLEMS*		
$\beta$	SE	$p$	$\beta$	SE	$p$	$\beta$	SE	$p$
1.80	.81	.03	-1.57	1.26	.22	-2.63	1.18	.04
.16	.98	.87	-1.76	1.29	.18	-2.98	.84	<.01
-1.94	1.23	.12	2.36	3.81	.54			
.38	1.13	.74	10.32	3.51	<.01	10.53	3.18	<.01
-4.60	2.17	.04	3.06	6.74	.65			
3.64	1.51	.02	-1.20	2.36	.61			
.27	1.80	.88	-1.54	2.40	.52			
-.58	1.35	.67	-2.18	2.12	.31	-.69	2.18	.76
-1.41	1.62	.39	-2.74	2.16	.21	1.58	1.62	.34
.84	2.72	.76	-.18	4.27	.97			
-.57	3.49	.87	5.52	4.52	.23			

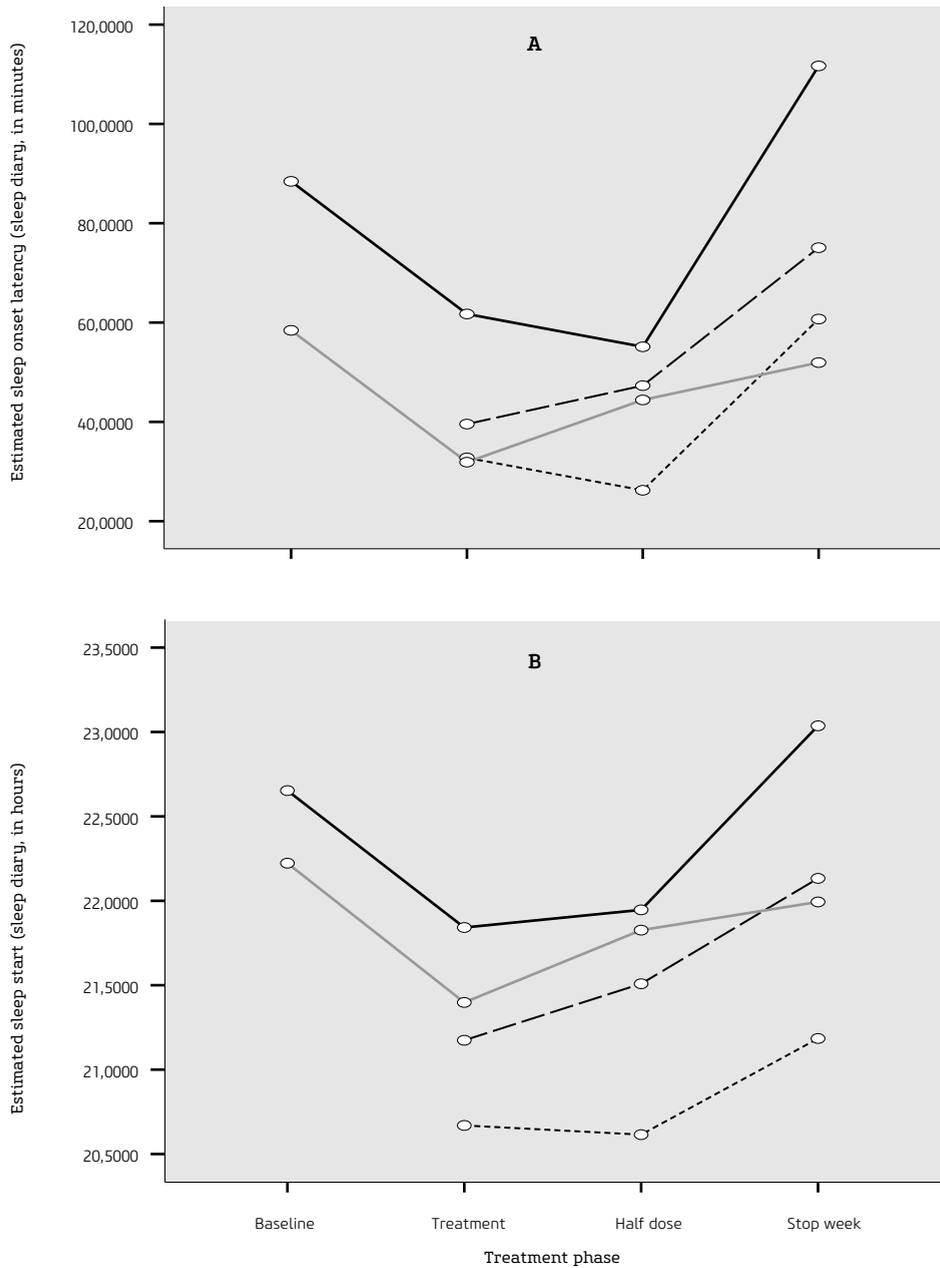
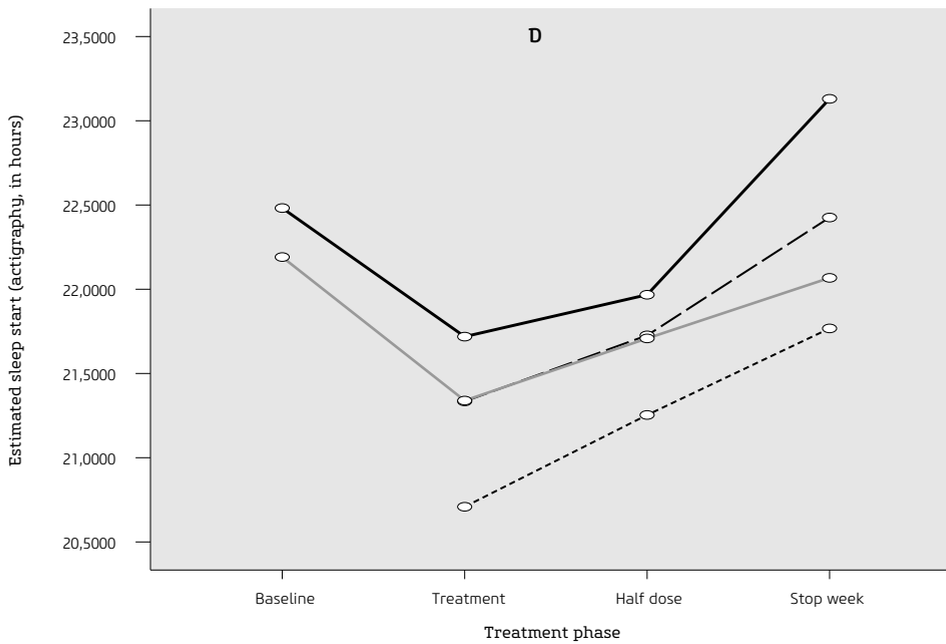
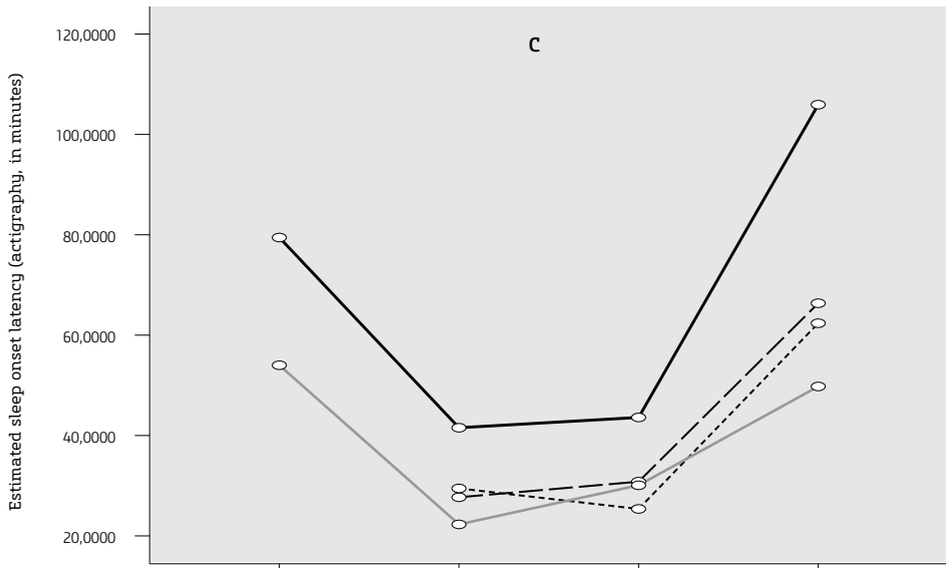
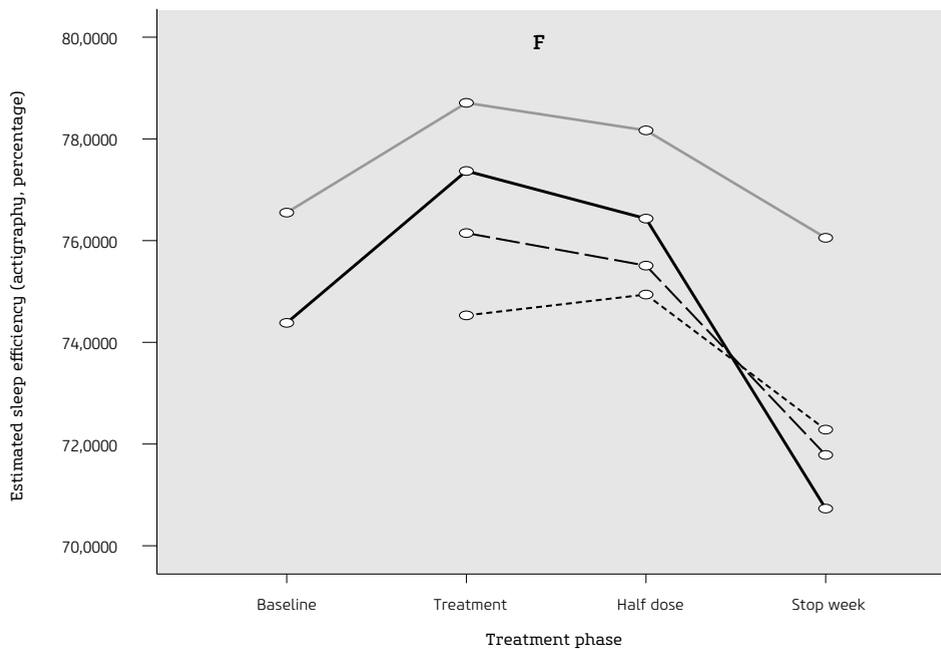
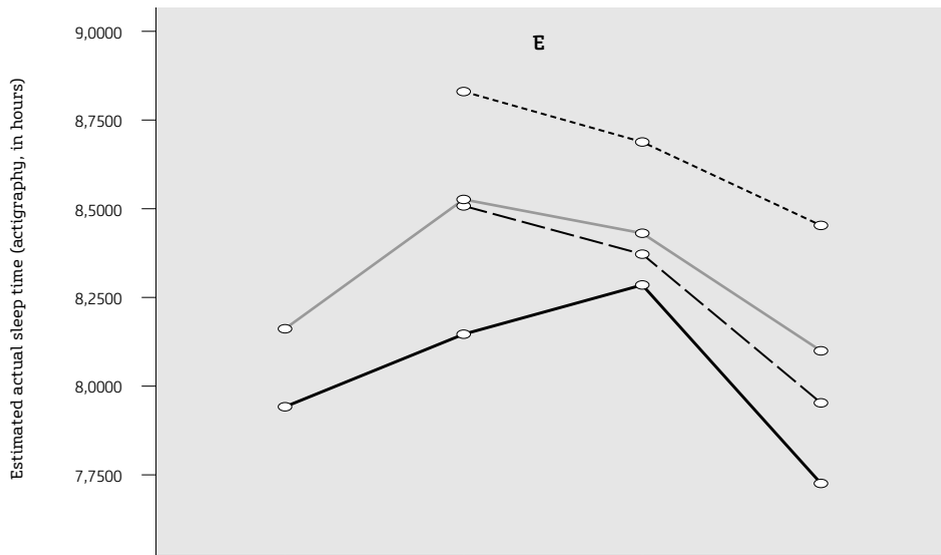


Figure 3a-f. Average sleep scores in different treatment phases by group and ADHD / autism diagnosis

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GROUP: — Comparison, no ADHD/autism    — Experimental, no ADHD/autism    - - - Comparison, ADHD/autism    — Experimental, ADHD/autism



GROUP: — Comparison, no ADHD/autism    — Experimental, no ADHD/autism    - - - Comparison, ADHD/autism    — Experimental, ADHD/autism

## DISCUSSION

This pilot study shows that melatonin treatment advanced Dim Light Melatonin Onset (DLMO) and sleep onset, improved parent-reported health, and reduced behaviour problems in children with chronic sleep onset problems related to a delayed DLMO. However, most treatment effects disappeared after melatonin treatment was discontinued. Only for sleep latency (according to parent report) and sleep start (according to both parent report and actigraphy), we found that sleep deterioration in the stop week was less strong for the experimental group, which suggests an effect of classical conditioning.

When controlling for ADHD / autism, we found that sleep deterioration in the stop week in many cases was stronger for children with ADHD / autism in the experimental group. This suggests that classical conditioning is counterproductive in children with ADHD / autism. A possible explanation for the stronger deterioration of children with ADHD/autism in the experimental group is that proportionally more children with ADHD /autism (4 of 5) in the experimental group than in the comparison group (4 of 14) received methylphenidate, which is found to negatively impact sleep.<sup>35</sup>

In our analyses, we operationalised comorbidity with a control variable for ADHD and autism combined, as the small sample size did not permit to include two separate variables. However, we further investigated the idea that comorbidity in general, or medication use in children with ADHD specifically, might deteriorate the effect of classical conditioning by repeating our analyses in two ways. Firstly after excluding children diagnosed with autism, and secondly after also excluding children diagnosed with ADHD but not using medication, but this did not change our results.

Remarkably, children with ADHD / autism in the experimental group had a later sleep start and shorter actual sleep time than children with ADHD / autism in the comparison group, which is probably due to the two-year age difference between these two groups (10.6 versus 8.7 years respectively). However, it should also be noted that due to the small group sizes individual children had a large impact on the results. For example, one of the children with ADHD in the experimental group slept extremely bad without melatonin, which had a strong impact on the average scores of this small group.

Interestingly, we found that an improvement of children's sleep coincides with an improvement in their health and behaviour. The effects on behaviour problems remained after the stop week. As the stop week was only one week, it is

possible that the positive effects will disappear when treatment is discontinued for a longer time. Although it would be interesting to investigate this hypothesis in future research, it might be difficult to gain the cooperation of parents and children, because they often want to continue melatonin treatment as soon as possible when the positive effects on sleep disappear.

There are several possible explanations for the fact that we did not find larger effects of classical conditioning in children without comorbidity. One explanation could be that the period of conditioning in our study (four weeks) was too short. Another explanation is that the timing of the stimuli that we used was not suitable. The drink was taken at the same time the melatonin was taken, because we aimed to condition the drink to the physiological reaction that the intake of melatonin induces (i.e. rise in melatonin secretion), following the studies of Evans and Bond<sup>23</sup> and Poser et al.<sup>24</sup> However, in these studies methohexitone was used, which almost immediately induced sleep. Melatonin does not work that fast. At the time children took the melatonin, they were still feeling awake. It might be that herewith the drink was coupled to the feeling of being awake, instead of to the physiological response as we intended. The timing of the lamp was probably better as children felt sleepy when they went to bed, but as we do not know any studies that used a lamp as conditioned stimulus, we do not have any evidence for the effectiveness of this kind of stimulus. A lamp is very different from counting or listening to the ticking of a metronome as was used as stimulus in the previous studies,<sup>23,24</sup> which demands a more attentive role of the person. Perhaps a stimulus requiring involvement of the person would have a better effect.

This study has several strengths, such as daily assessments of sleep in a time-series design with multiple cases and a comparison group, the use of objective sleep measurements, and a naturalistic setting. However, there are also some important limitations inherent to the fact that this was a pilot study. First, we should note that the sample in the experimental study was small, and that some of the children resumed melatonin treatment during the stop week already. However, due to the large number of observations per child, the effective sample size was still sufficient to find significant effects.

We should also note that this study did not include a real control group, but used a comparison group that participated in a previous study. Moreover, the experimental group was studied in spring, and the comparison group in the fall. Although melatonin was found to be effective in both groups, we cannot ignore

the possibility that longer light exposure also had an adverse effect on sleep after discontinuation of treatment in the experimental group. We did not check the effect of the sleep hygiene instructions. However, sleep hygiene instructions provided were the same in the experimental and comparison group, so we do not have any indication to suspect that effect of sleep hygiene instructions would differ between the groups.

## **CONCLUSIONS**

The present pilot study is the first study investigating the effects of classical conditioning in preserving effects of melatonin treatment. The results of this study indicate that there might be some effects of classical conditioning of melatonin treatment in children with sleep onset problems related to delayed melatonin onset, but only for children without comorbid ADHD or autism. We therefore believe it is important to further investigate the effectiveness of classical conditioning of melatonin use in children without comorbid ADHD or autism. For future research we would recommend to study larger, more homogeneous samples, to use other stimuli, preferably more specifically directed at the time of falling asleep, and to vary the duration of both melatonin treatment and the period of classical conditioning.

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### **Ethical standards**

**Ethical approval:** All procedures performed in studies involving human participants were in accordance with the Declaration of Helsinki (1964). The study was approved by the Ethical Committee of the Research Institute of Child Development and Education, University of Amsterdam.

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

**Conflict of interest:** The authors declare that there is no conflict of interest regarding the publication of this paper.

## REFERENCES

1. Fricke-Oerkermann L, Plücker J, Schredl M, et al. Prevalence and course of sleep problems in childhood. *Sleep*. 2007;30:1371-1377.
2. Spruyt K, O'Brien LM, Cluydts R, Verleye GB, Ferri R. Odds, prevalence and predictors of sleep problems in school-age normal children. *J Sleep Res*. 2005;14:163-176.
3. Dewald JF, Meijer AM, Oort FJ, Kerkhof GA, Bögels SM. The influence of sleep quality, sleep duration and sleepiness on school performance in children and adolescents: a meta-analytic review. *Sleep Med Rev*. 2010;14:179-189.
4. Fallone G, Owens JA, Deane J. Sleepiness in children and adolescents: clinical implications. *Sleep Med Rev*. 2002;6:287-306.
5. Gregory AM, O'Connor TG. Sleep problems in childhood: a longitudinal study of developmental change and association with behavioral problems. *J Am Acad Child Adolesc Psychiatry*. 2002;41:964-971.
6. Gregory AM, Caspi A, Eley TC, Moffitt TE, O'Connor TG, Poulton R. Prospective longitudinal associations between persistent sleep problems in childhood and anxiety and depression disorders in adulthood. *J Abnorm Child Psychol*. 2005;33:157-163.
7. Brzezinski A. Melatonin in humans. *N Engl J Med*. 1997;336:186-195.
8. Bruni O, Alonso-Alconada D, Besag F, et al. Current role of melatonin in pediatric neurology: Clinical recommendations. *Eur J Paediatr Neurol*. 2015;19:122-133.
9. Smits MG, Nagtegaal EE, van der Heijden J, Coenen AM, Kerkhof GA. Melatonin for chronic sleep onset insomnia in children: a randomized placebo-controlled trial. *J Child Neurol*. 2001;16:86-92.
10. Smits MG, van Stel HF, van der Heijden KB, Meijer AM, Coenen AML, Kerkhof GA. Melatonin improves health status and sleep in children with idiopathic chronic sleep-onset insomnia: A randomized placebo-controlled trial. *J Am Acad Child Adolesc Psychiatry*. 2003;42:1286-1293.
11. Van Geijlswijk IM, Korzilius HPLM, Smits MG. The use of exogenous melatonin in Delayed Sleep Phase Disorder: A meta-analysis. *Sleep*. 2010;33:1605-1614.
12. Van Maanen A, Meijer AM, Smits MG, Oort FJ. Termination of short term melatonin treatment in children with delayed Dim Light Melatonin Onset: Effects on sleep, health, behavior problems, and parenting stress. *Sleep Med*. 2011;12:875-879.
13. Rossignol DA, Frye RE. Melatonin in autism spectrum disorders: a systematic review and meta-analysis. *Dev Med Child Neurol*. 2011;53:783-792.
14. Van der Heijden KB, Smits MG, van Someren EJW, Ridderinkhof R, Gunning B. Effect of melatonin on sleep, behavior, and cognition in ADHD and chronic sleep-onset insomnia. *J Am Acad Child Adolesc Psychiatry*. 2007;46:233-241.
15. Johnson KP, Giannotti F, Cortesi F. Sleep patterns in autism spectrum disorders. *Child Adolesc Psychiatr Clin North Am*. 2009;18:917-928.
16. Stein MA. Unravelling sleep problems in treated and untreated children with ADHD. *J Child Adolesc Psychopharmacol*.

- 1999;9:157-168.
17. Sung V, Hiscock H, Sciberras E, Efron D. Sleep problems in children with attention-deficit/hyperactivity disorder: prevalence and the effect on the child and family. *Arch Pediatr Adolesc Med.* 2008;162:336-342.
  18. Williams PG, Sears LL, Allard A. Sleep problems in children with autism. *J Sleep Res.* 2004;13:265-268.
  19. Carr R, Wasdell MB, Hamilton D, et al. Long-term effectiveness outcome of melatonin therapy in children with treatment-resistant circadian rhythm sleep disorders. *J Pineal Res.* 2007;43:351-359.
  20. Hoebert M, van der Heijden KB, van Geijlswijk IM, Smits MG. Long-term follow-up of melatonin treatment in children with ADHD and chronic sleep onset insomnia. *J Pineal Res.* 2009;47:1-7.
  21. Van Geijlswijk IM, Mol RH, Egberts TCG, Smits MG. Evaluation of sleep, puberty and mental health in children with long-term melatonin treatment for chronic idiopathic childhood sleep onset insomnia. *Psychopharmacology.* 2011;216:111-120.
  22. Pavlov IP. *Conditioned reflexes.* Oxford: Oxford University Press, 1927.
  23. Evans DR, Bond IK. Reciprocal inhibition therapy and classical conditioning in the treatment of insomnia. *Behav Res Ther.* 1969;7:323-325.
  24. Poser EG, Fenton GW, Scotton L. The classical conditioning of sleep and wakefulness. *Behav Res Ther.* 1965;3:259-264.
  25. Vits S, Cesko E, Enck P, Hillen U, Schadendorf D, Schedlowski M. Behavioral conditioning as the mediator of placebo responses in the immune system. *Philos Trans R Soc B Biol Sci.* 2011;366:1799-1807.
  26. Keijzer H, Smits MG, Duffy JF, Curfs LMG. Why the dim light melatonin onset (DLMO) should be measured before treatment of patients with circadian-rhythm sleep disorders. *Sleep Med Rev.* 2014;18:333-339.
  27. Post MWM, Kuyvenhoven MM, Verheij TJM, de Melker RA, Hoes AW. De Nederlandse 'Functional Status II (R)': een vragenlijst voor het meten van de functionele gezondheidstoestand van kinderen. *Ned Tijdschr Geneeskd.* 1998;142:2675-2679.
  28. Stein REK, Jessop DJ. Functional status II(R): a measure of child health status. *Med Care.* 1990;28:1041-1055.
  29. De Brock AJLL, Vermulst AA, Gerris JRM, Abidin RR. Nijmeegse ouderlijke stress index: meetinstrument voor de vaststelling van stress bij opvoeders: een uitgebreide versie (NOSI) voor psychodiagnostische doeleinden en een verkorte versie (NOSIK) voor signaleringsdoeleinden. Lisse, The Netherlands: Swets & Zeitlinger, 1992.
  30. Goodman R. The Strengths and Difficulties Questionnaire: A research note. *J Child Psychol. Psychiatry.* 1997;38:581-586.
  31. Van Widenfelt BM, Goedhart AW, Trefers PDA, Goodman R. Dutch version of the Strengths and Difficulties Questionnaire (SDQ). *Eur Child Adolesc Psychiatry.* 2003;12:281-289.
  32. Achenbach TM *Manual for the Child Behavior Checklist and 1991 Profile.* Burlington: University Associates in Psychiatry, 1991.
  33. Verhulst FC, van der Ende J, Koot HM. *Handleiding voor de CBCL/ 4-18.* Erasmus University Rotterdam, The Netherlands, 1996.

34. Snijders TAB, Bosker RJ. Multilevel analysis: an introduction to basic and advanced multilevel modeling. London: Sage, 1999.
35. Galland BC, Tripp E, Taylor BJ. The sleep of children with attention deficit hyperactivity disorder on and off methylphenidate: a matched case-control study. *J Sleep Res.* 2010;19:366-373.