

Investigating the best treatment for the Delayed Sleep Phase Syndrome in adults with Attention-Deficit/Hyperactivity Disorder and it's effect on health and appetite

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This study will investigate the best treatment of Delayed Sleep Phase Syndrome (melatonin, placebo or melatonin and light therapy), and will determine whether patients with ADHD and DSPS have blood values that may indicate susceptibility for chronic...

Ethische beoordeling	Positief advies
Status	Werving gestart
Type aandoening	-
Onderzoekstype	Interventie onderzoek

Samenvatting

ID

NL-OMON22013

Bron

NTR

Verkorte titel

FASE

Aandoening

English:

Attention-Deficit/Hyperactivity Disorder (ADHD)

Delayed Sleep Phase Syndrome (DSPS)

Dutch:

ADHD

Verlate slaapfase syndroom

Ondersteuning

Primaire sponsor: Marc Blom, PhD, Director PsyQ The Hague

Overige ondersteuning: Fonds NutsOhra

PsyQ Research Fund

PsyQ Expertise Center Adult ADHD

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

The phase shift of the sleep/wake cycle as evaluated by the time of DLMO directly after treatment.

Toelichting onderzoek

Achtergrond van het onderzoek

About 80% of adults with Attention-Deficit/Hyperactivity Disorder (ADHD) have chronic sleep-onset problems and the majority of them has a Delayed Sleep Phase Syndrome (DSPS). DSPS is characterized by chronic sleep debt, which has negative consequences for physical health and is associated on the long term with weight gain, obesity, diabetes, hypertension, metabolic syndrome, cardiovascular disease, and even cancer. This study investigates the best treatment of DSPS (melatonin, placebo or melatonin and light therapy), and will determine whether patients with ADHD and DSPS have suboptimal biomarkers of chronic diseases like diabetes. Fifty-one adults with ADHD and DSPS will be recruited from the PsyQ Outpatient clinics in the Netherlands.

Doel van het onderzoek

This study will investigate the best treatment of Delayed Sleep Phase Syndrome (melatonin, placebo or melatonin and light therapy), and will determine whether patients with ADHD and DSPS have blood values that may indicate susceptibility for chronic diseases like diabetes.

The results will give insight into the association between DSPS, ADHD, and chronic conditions such as diabetes, and may serve as a model for the increase in obesity and metabolic syndrome in the general population, in which sleep loss is common due to the extended use of artificial light.

Onderzoeksopzet

All outcomes will be measured at baseline, directly after treatment and three weeks after treatment.

Methods of measurement:

Improvement of appetite hormones - blood

Improvement of diabetic parameters -blood

Improvement of biomarker profiles -blood

Improvement of cardiovascular parameters - ECG/ Ambulant Blood Pressure Monitor

Improvement of sleep parameters -actigraphy

Improvement of quality of life -questionnaire

Decrease of ADHD symptoms - questionnaire

Decrease of intake of carbohydrate-rich food -questionnaire

Treatment satisfaction -questionnaire

Onderzoeksproduct en/of interventie

Patients will be randomized for sleep education, plus:

(1) 0.5 mg of melatonin daily,

(2) 0.5 mg of placebo daily, or

(3) 0.5 mg of melatonin daily, plus 30 minutes of light therapy in the morning between 7 and 8 AM.

Each treatment takes three weeks; the Melatonin intake will be three hours before the patient's individually determined Dim-Light Melatonin Onset (DLMO), and will be advanced during the treatment period.

Contactpersonen

Publiek

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Wetenschappelijk

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Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

- Age between 18 and 55 years old
- Diagnosis ADHD
- Diagnosis DSPS

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

- Psychotic illness
- Untreated mood disorder
- Untreated anxiety disorder
- Alcohol intake > 2U/day, or for woman >15U/week, for men >21 U/week
- Use of cannabis or harddrugs within one month prior to study participation

- Suspected dementia, anamnestic disorder or other cognitive disorder
- Mental retardation
- Use of the following medication within one month prior to study participation: psychostimulants, melatonin, mirtazapin, sleep medication, antipsychotics, clonidin, benzodiazepins, bêta-blockers
- Insufficient fluency of the Dutch language
- Evening or night shift work
- Travel over >2 time zones two weeks prior to study participation (because of possible jet lag)
- Pregnancy or breast feeding
- Having young children who may disturb night rest
- Type 2 diabetes mellitus
- Light therapy one month prior to study participation

Onderzoeksopzet

Opzet

Type:	Interventie onderzoek
Onderzoeksmodel:	Parallel
Toewijzing:	Gerandomiseerd
Blinding:	Dubbelblind
Controle:	Placebo

Deelname

Nederland	
Status:	Werving gestart
(Verwachte) startdatum:	16-05-2013
Aantal proefpersonen:	51
Type:	Verwachte startdatum

Ethische beoordeling

Positief advies

Datum: 17-05-2013

Soort: Eerste indiening

Registraties

Opgevolgd door onderstaande (mogelijk meer actuele) registratie

ID: 46908

Bron: ToetsingOnline

Titel:

Andere (mogelijk minder actuele) registraties in dit register

Geen registraties gevonden.

In andere registers

Register	ID
NTR-new	NL3831
NTR-old	NTR3999
CCMO	NL39579.058.12
ISRCTN	ISRCTN wordt niet meer aangevraagd.
OMON	NL-OMON46908

Resultaten