

Blended Alcohol Depression Ehealth

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We expect to find an 25% treatment response for TAU and 50% treatment response for TAU + online alcohol reduction intervention.

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

Samenvatting

ID

NL-OMON23390

Bron

NTR

Verkorte titel

BLADE

Aandoening

Depression, alcohol misuse

Ondersteuning

Primaire sponsor: Arkin Institute for Mental Health

Overige ondersteuning: ZonMw (The Netherlands Organisation for Health Research and Development)

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

Treatment respons: the treatment response outcome measure combines alcohol and depression measures into a composite score which indicates whether treatment has been successful or not. The treatment is deemed successful if all three conditions below are met:
- Drinking less than 21 (males) / 14 (females) glasses of alcohol in the week prior to

measurement. 'Glasses' is operationalized as standard drinks which contain 10 g of ethanol (the European standard);

- 0 days with 4 or more (women), or 5 or more (men) drinks reported in the last 7 days;
- Center for Epidemiological Studies-Depression (CES-D) score < 16 or a reduction of 40% relative to CES-D at baseline.

Toelichting onderzoek

Achtergrond van het onderzoek

Objective: To evaluate the (cost-) effectiveness of adding an internet-based alcohol reduction intervention to depression treatment (TAU), in reducing alcohol and depression outcomes compared to TAU alone among young adults aged 18-35 years.

Study design: A multicentre, 2-arm randomized controlled trial. Assessments take place at baseline, 3 months, 6 months (primary endpoint) and 12 months.

Study population: Patients diagnosed with a depressive disorder enrolling for or in depression treatment, age 18-35, and a total score of ≥ 8 for men and ≥ 5 for women on the Alcohol Use Disorder Identification Test (AUDIT).

Intervention: The intervention group receives TAU + guided online alcohol reduction intervention (self-help modules). The content of these modules aimed at reducing alcohol use and is based on existing internet-based self-help materials and on cognitive behaviour therapy and motivational interviewing (CBT/MI) and are tailored for young people in depression therapy. TAU consists regular CBT or other evidence-based psychotherapy combined with medication if necessary. TAU is directed at activation and identification of maladaptive cognitions. Control group receives TAU alone.

Main study parameters/endpoints: The primary outcome is treatment response, a composite score that combines alcohol and depression measures which indicates whether treatment has been successful or not.

Doel van het onderzoek

We expect to find an 25% treatment response for TAU and 50% treatment response for TAU + online alcohol reduction intervention.

Onderzoeksopzet

T0 = Baseline

T1 = 3 months follow-up

T2 = 6 months follow-up (primary endpoint RCT)

T3 = 12 months follow-up

Onderzoeksproduct en/of interventie

The intervention group receives blended TAU + a guided web-based alcohol reduction intervention. Guidance is focused on process support (no care related guidance). The content of the modules is based on existing internet-based (guided) self-help materials developed by Arkin/Jellinek and is based on cognitive behaviour therapy and motivational interviewing (CBT/MI) and tailored for young people in depression therapy. The online alcohol reduction intervention consist of 5 modules + 1 aftercare module, conceptually in line with the evidence-based brief CBT for adults and exists mainly of information (in text and video) and short assignments and registration of daily alcohol intake and mood. A prototype of the online intervention was evaluated for usability in focus groups by clients and therapists. Based on the input from these focus groups adjustments were made to the final online alcohol intervention. The online alcohol reduction intervention can be accessed on computer and other mobile devices (e.g. smartphone/tablet).

Both groups will receive TAU. The control group will receive TAU alone. The TAU has a duration of 4-6 months. TAU consists of 8 to 16 45-minutes sessions of regular Cognitive-Behavioural Therapy (CBT) or other evidence-based psychotherapy (e.g. interpersonal psychotherapy, problem solving therapy), combined with medication if necessary.

Contactpersonen

Publiek

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Wetenschappelijk

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Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

- Depressive disorder as a diagnosis
- Age 18-35
- AUDIT score of ≥ 8 for men and ≥ 5 for women
- Moderately proficient in Dutch
- Willing to provide contact details including (mobile)phone,
- Healthcare insurance coverage
- Computer/tablet at home and willingness to use this for treatment and research purposes
- Informed consent regarding the study provided by the patient

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

- Acute psychosis
- Alcohol dependence as primary diagnosis
- Dementia
- Waitlisted for in-patient mental health care
- Pregnancy

Onderzoeksopzet

Opzet

Type:	Interventie onderzoek
Onderzoeksmodel:	Parallel
Toewijzing:	Gerandomiseerd
Blinding:	Enkelblind
Controle:	Geneesmiddel

Deelname

Nederland	
Status:	Werving nog niet gestart
(Verwachte) startdatum:	18-11-2019
Aantal proefpersonen:	156
Type:	Verwachte startdatum

Voornemen beschikbaar stellen Individuele Patiënten Data (IPD)

Wordt de data na het onderzoek gedeeld: Nog niet bepaald

Toelichting

NA

Ethische beoordeling

Positief advies

Datum: 29-10-2019

Soort: Eerste indiening

Registraties

Opgevolgd door onderstaande (mogelijk meer actuele) registratie

ID: 49219

Bron: ToetsingOnline

Titel:

Andere (mogelijk minder actuele) registraties in dit register

Geen registraties gevonden.

In overige registers

Register	ID
NTR-new	NL8122
CCMO	NL66899.100.18
OMON	NL-OMON49219

Resultaten

Samenvatting resultaten

NA