

Innovation in the treatment of Dissociative Identity Disorder: The application of schema therapy.

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Investigate the efficacy of schema therapy in Dissociative Identity Disorder

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

Samenvatting

ID

NL-OMON29102

Bron

NTR

Aandoening

Dissociative Identity Disorder

Ondersteuning

Primaire sponsor: University of Groningen

Overige ondersteuning: University of Groningen

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

Dissociative symptoms (DSS), presence of DID (SCID-D), dropout

Toelichting onderzoek

Achtergrond van het onderzoek

Patients suffering from Dissociative Identity Disorder (DID) were found to have the highest psychiatric treatment costs among patients receiving medical care. There are no consensus treatment guidelines for dissociative disorders and current clinical practice entails lengthy treatment with a high dropout rate. In the current proposal, the efficacy of an alternative treatment for DID (i.e., schema therapy) is tested. Due to the rarity of the disorder and the lengthy treatment of these patients, a case series experimental approach will be used (non-concurrent multiple baseline design). Ten DID outpatients will be included from several community mental health institutes in the Netherlands. A process-measure of dissociative symptoms will be included as index of change. Additionally, various pre-, post-, and follow-up measures will be included encompassing an assessment of the presence of DID, trait dissociative symptoms, comorbid symptomatology, and global symptomatic distress.

Doel van het onderzoek

Investigate the efficacy of schema therapy in Dissociative Identity Disorder

Onderzoeksopzet

Multicenter, non-concurrent multiple baseline design. Patients are randomly assigned to one of the predetermined baseline lengths. Baseline observations are carried out, and after an education phase, the treatment is implemented.

Timepoints: Process measure weekly during baseline and education phase, other measures before and after baseline period, after education phase, each 6 months during intervention, at the end of the intervention and at follow up (six months after the end of treatment).

Additional baseline measures SCID I and II, CTQ.

An additional follow-up assessment will be included one year after the end of treatment (i.e., 6 months after the end of the booster sessions).

Onderzoeksproduct en/of interventie

Treatment will consist of two individual sessions a week during the first two years, followed by one individual session a week during the third year, with each session lasting 50 minutes. The treatment will be theoretically consistent with the model described in Young, Klosko, and Weishaar (2003), which was expanded for the treatment of BPD patients (van Genderen & Arntz 2010; Farrell & Shaw, 2012), and adapted for the specific treatment needs of DID patients (Farrell & Shaw, 2013).

Six monthly booster sessions will be included after the end of treatment.

Contactpersonen

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Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

- 1) Main diagnosis of DID
- 2) Age between 18 and 60
- 3) Dutch literacy
- 4) Stable use of medication

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

- 1) Mental retardation (IQ < 80)
- 2) Drug/alcohol dependency
- 3) Acute suicide risk
- 4) Florid psychotic episodes

Onderzoeksopzet

Opzet

Type:	Interventie onderzoek
Onderzoeksmodel:	Parallel
Toewijzing:	N.v.t. / één studie arm
Blinding:	Open / niet geblindeerd
Controle:	N.v.t. / onbekend

Deelname

Nederland	
Status:	Werving nog niet gestart
(Verwachte) startdatum:	01-06-2014
Aantal proefpersonen:	10
Type:	Verwachte startdatum

Voornemen beschikbaar stellen Individuele Patiënten Data (IPD)

Wordt de data na het onderzoek gedeeld: Nog niet bepaald

Ethische beoordeling

Positief advies	
Datum:	07-04-2014
Soort:	Eerste indiening

Registraties

Opgevolgd door onderstaande (mogelijk meer actuele) registratie

Geen registraties gevonden.

Andere (mogelijk minder actuele) registraties in dit register

Geen registraties gevonden.

In overige registers

Register	ID
NTR-new	NL4356
NTR-old	NTR4496
Ander register	Ethische Toetsingscommissie RUG ppo-013-110 : METC UMCG Groningen METc 2013/420

Resultaten