

The validity of the Diagnostic Screening Instrument (DSI) according to the SCID-5-S

Gepubliceerd: 02-09-2020 Laatste bijgewerkt: 18-08-2022

To determine if the DSI is valid.

Ethische beoordeling	Positief advies
Status	Werving gestart
Type aandoening	-
Onderzoekstype	Observationeel onderzoek, zonder invasieve metingen

Samenvatting

ID

NL-OMON27151

Bron

Nationaal Trial Register

Verkorte titel

TBA

Aandoening

Psychological complaints for which is referred to outpatient treatment

Ondersteuning

Primaire sponsor: HSK Groep

Overige ondersteuning: HSK Groep

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

In this study the DSI will be found to be valid if the level of agreement between the DSI and the SCID-5-S at both the classification level as at the disorder level has a Cohen's Kappa of

0.7 or more.

Toelichting onderzoek

Achtergrond van het onderzoek

Various studies have shown that in routine clinical practice, the reported levels of interrater reliability for classifications according to the DSM (APA, 2013) is low. Therefore, the use of semi-structured interviews is advised to guarantee the validity of a classification according to the DSM. The disadvantage of most (semi-) structured interviews is that they are long, contain many questions and that extensive training is required to conduct them properly. The DSI is a shorter and digital (semi-) structured questionnaire. However, this has not yet been assessed for validity. In this study, the criterion validity of the DSI will be assessed by administering the DSI as well as the SCID-5-S by each subjects in the same week.

In this study, the criterion validity of the DSI is assessed, with the following research questions:

- 1) Is the DSI sufficiently valid for setting a DSM classification at category level?
- 2) Is the DSI sufficiently valid for setting a DSM classification at the disorder level?
- 3) Is there indeed a difference in the time it takes to administer the DSI and the SCID-5-S?

Doel van het onderzoek

To determine if the DSI is valid.

Onderzoeksopzet

Two time points within one week

Onderzoeksproduct en/of interventie

With each participants the DSI and the SCID-5-S will be administered within one week.

Contactpersonen

Publiek

HSK
Emelie Tan

0031619283420

Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

Anyone who qualifies for outpatient treatment.

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

HSK has certain exclusion criteria for entering treatment. These are: schizophrenia, psychosis or delusional disorder; serious disorders in and through the use of substances such as alcohol and drugs; bipolar disorder as main reporting complaint; serious personality problems; eating disorders and eating problems; non-congenital brain injury (NAH) and/or IQ <80; developmental disorders: attention deficiency/hyperactivity disorder (AD(H)D), autism spectrum disorder (ASD); severe suicidality and/or crisis sensitivity and language barrier. It is not always clear in advance that a client, referred for treatment, meets one or more of the above disorders or criteria. In other words, these are not exclusion criteria for the study as such, but for the subsequent treatment. However, these clients may be referred less.

Onderzoeksopzet

Opzet

Type:	Observationeel onderzoek, zonder invasieve metingen
Onderzoeksmodel:	Anders
Toewijzing:	Niet-gerandomiseerd
Blinding:	Open / niet geblindeerd
Controle:	N.v.t. / onbekend

Deelname

Nederland
Status: Werving gestart
(Verwachte) startdatum: 01-08-2020
Aantal proefpersonen: 250
Type: Verwachte startdatum

Voornemen beschikbaar stellen Individuele Patiënten Data (IPD)

Wordt de data na het onderzoek gedeeld: Ja

Toelichting

N/A

Ethische beoordeling

Positief advies
Datum: 02-09-2020
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
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NTR-new	NL9743
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Ander register Commissie Mensgebonden Onderzoek regio Arnhem-Nijmegen : 2020-6308

Resultaten

Samenvatting resultaten

N/A