Development en validation of the 'Yale-Brown Obsessive Compulsive Scale for Body Dysmorphic Disorder Self Rating version' (Y-BOCS BDD-SR)

Published: 17-07-2008 Last updated: 07-05-2024

This study will determine whether the *Yale-Brown Obsessive-Compulsive Scale modified for Body Dysmorphic Disorder (Y-BOCS-BDD) interview* can be replaced by a questionaire (selfrating (SR)).

Ethical review	Approved WMO
Status	Pending
Health condition type	Somatic symptom and related disorders
Study type	Observational non invasive

Summary

ID

NL-OMON32136

Source ToetsingOnline

Brief title Development en validation of the Y-BOCS BDD-SR

Condition

• Somatic symptom and related disorders

Synonym body dysmorphic disorder

Research involving Human

Sponsors and support

Primary sponsor: Leids Universitair Medisch Centrum

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Source(s) of monetary or material Support: Ministerie van OC&W

Intervention

Keyword: body dysmorphic disorder, questionnaire versus interview, Y-BOCS-BDD

Outcome measures

Primary outcome

The results of the interview and questionaire will be compared.

Secondary outcome

Not applicable.

Study description

Background summary

The standardised interview Y-BOCS-BDD is an internationally used instrument to determine the severity of Body Dismorphic Disorder (Phillips, 1997). The YBOCS modified for BDD (Phillips, 1997 52 /id) has become the gold standard for assessing outcome of treatment in randomised controlled trials (Veale, 2003). Validity and reliability of the American version is good (Philips 1997). In the Netherlands we use a translation of the original standardised interview (Van Rood, Bouman, 2001). Recent analysis of the dutch version (Van Rood en Bouman (2007)) show equally good qualities.

The De Y-BOCS-BDD questionaire is based on the original American interview. The version is reviewed by Veale and Philips and their comments have been used to improve the questionaire.

Study objective

This study will determine whether the *Yale-Brown Obsessive-Compulsive Scale modified for Body Dysmorphic Disorder (Y-BOCS-BDD) interview* can be replaced by a questionaire (self-rating (SR)).

Study design

Observational reseach using a questionnaire and interview

Study burden and risks

The burden associated with participation is restricted the time needed to fill in the questionnaire and being interviewed. The total burden for the patient will be 50 to 150 minutes during a 12 month period. There will be no financial compensation for the time invested. There are no risks associated with participation.

There are no direct benefits for participants, but in the future the new questionaire could mean a lesser burden for new participants of ROM because the questionaire takes far less time.

Contacts

Public Leids Universitair Medisch Centrum

Albinusdreef 2300 RC Leiden Nederland **Scientific** Leids Universitair Medisch Centrum

Albinusdreef 2300 RC Leiden Nederland

Trial sites

Listed location countries

Netherlands

Eligibility criteria

Age Adults (18-64 years) Elderly (65 years and older)

Inclusion criteria

BDD Diagnosis and/ or answering "yes" at first question of BDD-screener

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Exclusion criteria

Not enough understanding of Dutch to fill in the questionnaires.

Study design

Design

Study type: Observational non invasive	
Masking:	Open (masking not used)
Control:	Uncontrolled
Primary purpose:	Diagnostic

Recruitment

NL	
Recruitment status:	Pending
Start date (anticipated):	01-04-2008
Enrollment:	100
Туре:	Anticipated

Ethics review

Approved WMO	
Application type:	First submission
Review commission:	METC Leids Universitair Medisch Centrum (Leiden)

Study registrations

Followed up by the following (possibly more current) registration

No registrations found.

Other (possibly less up-to-date) registrations in this register

No registrations found.

In other registers

Register

ССМО

ID NL21935.058.08