

Do both healthcare professionals and patients benefit from suicide prevention training of professionals?

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Our primary outcome is change in suicide ideation in patients. We hypothesise that, as a result of the improved skills and confidence of healthcare professionals in the experimental condition due to the intervention, suicidal patients are better...

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

Samenvatting

ID

NL-OMON26081

Bron

NTR

Verkorte titel

PITSTOPsuicide

Aandoening

suicide suicide ideation depression anxiety knowledge confidence skills of professionals

Ondersteuning

Primaire sponsor: VU Amsterdam, clinical psychology

Overige ondersteuning: ZonMW

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

Our primary outcome is change in suicide ideation in patients. We hypothesize that, as a result of the improved skills and confidence of healthcare professionals in the experimental condition due to the intervention, suicidal patients are better assessed and treated and will therefore more quickly recover from suicidal ideation. Also we expect less episodes of suicidal ideation, deliberate self-harm, non fatal suicide attempts and less health care consumption of patients in the units of the experimental condition.

Toelichting onderzoek

Achtergrond van het onderzoek

Adherence to evidence based practice guidelines in mental health care is not self-evident. Merely developing and publishing guidelines will not bring the wanted change because of time-restraint, resistance to change and organizational complexity. In order to gain greater acceptance and adherence to guidelines, guideline implementation is imperative.

The objective of this study is to examine if patients, professionals and the organizations benefit from a train the trainer course, that was developed to implement the new Dutch multidisciplinary practice guideline for assessment and treatment of suicidality.

Doel van het onderzoek

Our primary outcome is change in suicide ideation in patients. We hypothesize that, as a result of the improved skills and confidence of healthcare professionals in the experimental condition due to the intervention, suicidal patients are better assessed and treated and will therefore more quickly recover from suicidal ideation. Also we expect less episodes of suicidal ideation, deliberate self-harm, non fatal suicide attempts and less health care consumption of patients in the units of the experimental condition.

Onderzoeksopzet

Patients:

Via routine outcome measurement:

1. Start;
2. 1month;
3. 3months;
4. 6months.

Professional:

1. 2 weeks before intervention;
2. Directly after intervention;
3. After 3 months.

Onderzoeksproduct en/of interventie

We implement the guideline with a train the trainer course called PITSTOP suicide that consists of:

1. A multidisciplinary interactive group training;
2. Given by role model clinicians;
3. Combined with an e-learning module;
4. And textbook material.

The training is one day with the e-learning module taking one hour.

In the experimental condition, hundred percent of the registered nurses, psychologists, physicians and psychiatrists will be trained in their own team, where personalized feedback is possible.

In the control group, the guideline is not implemented, but spread in the "normal" way, via internet, lectures, leaflets etc.

Contactpersonen

Publiek

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Wetenschappelijk

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Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

1. > 18 years;
2. Fluent in dutch;
3. New resident in mental health care.

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

1. < 18 years;
2. Non fluent in dutch;
3. Psychotic.

Onderzoeksopzet

Opzet

Type:	Interventie onderzoek
Onderzoeksmodel:	Parallel
Toewijzing:	Gerandomiseerd

Blindering:	Enkelblind
Controle:	Geneesmiddel

Deelname

Nederland	
Status:	Werving nog niet gestart
(Verwachte) startdatum:	01-01-2012
Aantal proefpersonen:	8000
Type:	Verwachte startdatum

Ethische beoordeling

Positief advies	
Datum:	04-10-2011
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	NL2945
NTR-old	NTR3092
Ander register	METC VUmc / Wetenschapscommissie : 2011/151 / 2010-018;
ISRCTN	ISRCTN wordt niet meer aangevraagd.

Resultaten

Samenvatting resultaten

N/A