

Cognitive Behavioural Suicide Prevention for psychosis

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| Submission date 21/05/2010 | Recruitment status No longer recruiting | <input type="checkbox"/> Prospectively registered |
| Registration date 21/05/2010 | Overall study status Completed | <input type="checkbox"/> Protocol |
| Last Edited 09/09/2016 | Condition category Mental and Behavioural Disorders | <input type="checkbox"/> Statistical analysis plan |
| | | <input type="checkbox"/> Results |
| | | <input type="checkbox"/> Individual participant data |
| | | <input type="checkbox"/> Record updated in last year |

Plain English summary of protocol
Not provided at time of registration

Contact information

Type(s)
Scientific

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Additional identifiers

Protocol serial number
6329

Study information

Scientific Title
Prevention of Suicide risk in patients with psychosis through the development of a focused and conceptually-driven intervention

Acronym

CBSPP

Study objectives

Study aims:

1. To confirm and describe the psychological architecture that drives suicide behaviour, this consists of: information processing biases, suicide schema and appraisal systems. We intend to investigate hypotheses relating to determine the role of these in suicide behaviour specifically we will use information processing tasks such as autobiographical memory tasks to investigate bias; standardised assessment to investigate appraisal and schema.
2. To derive a method of assessing this architecture through semi-structured interview that will have clinical utility
3. To formulate and develop a manualised cognitive behavioural treatment programme for suicide prevention in psychosis and to test the clinical acceptance and feasibility of this intervention. It is hypothesised that the treatment will be acceptable, feasible and reduce suicide behaviour.
4. At the end of the programme we will be in position to design a clinical trial to test the efficacy of the intervention

Ethics approval required

Old ethics approval format

Ethics approval(s)

Stockport REC , 12/12/2008, ref: 08/H1012/97

Study design

Multicentre randomised interventional treatment trial

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Topic: Mental Health Research Network; Subtopic: Psychosis; Disease: Psychosis

Interventions

Those allocated to the experimental treatment will receive CBSPP as described in the manual in addition to treatment as usual (TAU). Treatment will take place over 12 weeks. Participants will be seen twice weekly. The control group will receive TAU alone. All participants will be re-assessed at 4 months and at 6 months.

Study entry: single randomisation only

Intervention Type

Other

Phase

Not Applicable

Primary outcome(s)

Suicidal behaviour, measured at 4 months and 6 months

Key secondary outcome(s)

Measured at baseline, 4 months and 6 months:

1. Beck Anxiety Inventory (BAI) to assess level of anxiety
2. Beck Hopelessness Scale
3. Calgary Depression Scale for Schizophrenia
4. Global assessment of functioning (DSM-IV) to assess overall functioning
5. Positive And Negative Symptoms Scale (PANSS) to assess severity of symptoms
6. Psychotic Symptom Rating Scales (PSYRATS) to assess control over symptoms
7. Questionnaire about the Process of Recovery (QPR)
8. Self Esteem Rating Scale
9. The Subjective Experiences of Psychotic Symptoms Scale (SEPSS)

Completion date

31/12/2010

Eligibility**Key inclusion criteria**

1. Diagnostic and Statistical Manual of Mental Disorders, 4th Edition (DSM-IV) diagnosis of schizophrenia, schizophreniform disorder, schizoaffective disorder, delusional disorder or psychotic disorder not otherwise specified (NOS)
2. Have had previous suicide attempts or current suicidal ideation
3. Are not currently acutely suicidal or considered a danger to themselves or others
4. Are receiving appropriate anti-psychotic medication
5. Are under the care of an appropriate clinical team and have contact with mental health service
6. Males and females aged between 18 and 65 years

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Sex

All

Key exclusion criteria

Not providing informed consent

Date of first enrolment

01/02/2009

Date of final enrolment

31/12/2010

Locations

Countries of recruitment

United Kingdom

England

Study participating centre**Zochonis Building**

Manchester

United Kingdom

M13 9PL

Sponsor information

Organisation

University of Manchester (UK)

ROR

<https://ror.org/027m9bs27>

Funder(s)

Funder type

Government

Funder Name

National Institute for Health Research (NIHR) (UK)

Results and Publications

Individual participant data (IPD) sharing plan**IPD sharing plan summary**

Not provided at time of registration