

Medication strategies in first onset schizophrenia (Mesifos)

Submission date 19/12/2005	Recruitment status No longer recruiting	☐ Prospectively registered
		☐ Protocol
Registration date 19/12/2005	Overall study status Completed	☐ Statistical analysis plan
		☒ Results
Last Edited 05/07/2013	Condition category Mental and Behavioural Disorders	☐ Individual participant data

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
NTR374; DO 0945-01-001

Study information

Scientific Title

Acronym

Mesifos

Study objectives

Overall research question: Is there a difference in quality of life between patients with a first psychotic episode, treated with targeted and maintenance treatment?

Detailed questions:

1. Do both treatment strategies differ with respect to quality of life, subjectively as well as objectively, regarding work, daily activities, housing, social network, satisfaction and wellbeing, including (para)suicide, aggressive behaviors towards others, contacts with police, days in jail, and to social role functioning?
2. Do both treatment strategies differ with respect to the course of the illness (relapse, quality of remission), side-effects of medication (dyskinesia, EPS, subjective well-being), and dependence on care facilities (including involuntary admission)?
3. Does the psychosocially oriented treatment lead to better compliance and earlier recognition of prodromal signs with the possibility of prevention of full blown psychosis by targeted pharmacological treatment?
4. Can we identify predictors of successful drug withdrawal/discontinuation?
5. To what extent are these treatment strategies acceptable to this patient population?
6. To what extent do early drop out and refusal make a difference with respect to mental health care consumption and social outcome?
7. Do direct medical costs differ between the two strategies?
8. Is there a difference regarding indirect costs and burden on the family?

Ethics approval required

Old ethics approval format

Ethics approval(s)

Received from local medical ethics committee

Primary study design

Interventional

Study design

Multicentre randomised open label active controlled parallel group trial

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Non affective psychosis, schizophrenia

Interventions

Maintenance treatment was carried out according to the guidelines of the APA. This entailed the preferred use of second-generation antipsychotics in low dose.

In targeted treatment the dose was gradually tapered in one or two months and discontinued, if possible. Tapering was allowed to be more gradual, subject to symptom levels and individual preferences of patients. If early warning signs of relapse emerged or positive symptoms recurred, clinicians were to reinstate or increase the dose of antipsychotic medication, not only in targeted, but also in maintenance treatment. If feasible and considered safe, in targeted treatment discontinuation was tried again.

Intervention Type

Other

Phase

Not Specified

Primary outcome(s)

Quality of life

Key secondary outcome(s)

1. Symptomatology
2. Relapse
3. Side effects
4. Social functioning
5. Burden on the family

Completion date

01/08/2005

Eligibility**Key inclusion criteria**

1. Suffering from a first episode of psychosis
2. 18-45 years of age
3. Treatment naïve
4. Responding to medication (remission of positive symptoms) within 6 months and remaining stable for another 6 months

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 Years

Upper age limit

45 Years

Sex

All

Key exclusion criteria

No remission or relapse within 6 months

Date of first enrolment

01/08/2001

Date of final enrolment

01/08/2005

Locations

Countries of recruitment

Netherlands

Study participating centre

University Medical Center Groningen

Groningen

Netherlands

9700 RB

Sponsor information

Organisation

University Medical Centre Groningen (UMCG) (Netherlands)

ROR

<https://ror.org/03cv38k47>

Funder(s)

Funder type

Industry

Funder Name

Eli Lilly B.V. (Netherlands)

Funder Name

Service Foundation (Stichting Diensbetoon) (Netherlands)

Funder Name

Support Foundation (Stichting Steun) (Netherlands)

Funder Name

Netherlands Organisation for Health Research and Development (ZonMw) (Netherlands)

Alternative Name(s)

Netherlands Organisation for Health Research and Development

Funding Body Type

Private sector organisation

Funding Body Subtype

Other non-profit organizations

Location

Netherlands

Results and Publications

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

Not provided at time of registration

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	01/04/2006		Yes	No
Results article	results	01/09/2013		Yes	No