Subjective well-being, craving for cannabis and compliance or medication switch in a randomized doubleblind study with olanzapine and risperidone.

No registrations found.

Ethical review Positive opinion

Status Recruitment stopped

Health condition type -

Study type Interventional

Summary

ID

NL-OMON22184

Source

NTR

Brief title

SUB.CAN.OLA.RIS

Intervention

Outcome measures

Primary outcome

- 1. Subjective Well-being under Neuroleptics scale (SWN);
- 2. Obsessive Compulsive Drug Use Scale (OCDUS);
- 3. Positive And Negative Symptoms Scale (PANSS) based on information from the semistructured interview (SCI- PANSS);
- 4. Calgary Depression Rating Scale (CDRS);
- 5. ESRS: Extra pyramidal Symptom Rating Scale;
 - 1 Subjective well-being, craving for cannabis and compliance or medication switch ... 3-05-2025

- 6. Clinical Global Impression (CGI);
- 7. Y-BOCS (Yale Brown Obsessive Compulsive Scale);
- 8. Desires for Drugs Questionnaire (DDQ);
- 9. Drug Use Self Report (DUSR);
- 10. Recent Drug Use Urinalysis (RDUU).

Secondary outcome

- 1. Drop out from the study;
- 2. Medication compliance and medication switch, symptoms and rehospitalisations during one year follow up, measured with the Life Chart Schedule (LCS).

Study description

Background summary

N/A

Study objective

N/A

Study design

N/A

Intervention

Patients are treated double blind with olanzapine (5-20 mg) or risperidone (1,25-5 mg) for 6 weeks. At t=0, t=7 days and t=42 days, questionnaires are taken and after 6 weeks the medication is disclosed. The physician and patient decide if this neuroleptic will be continued. After one year the questionnaires are taken once more.

Contacts

Public

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Eligibility criteria

Inclusion criteria

- 1. Patients should be able to understand the study description and give informed consent;
- 2. Diagnosis of schizophrenia, schizoaffective disorder or schizophreniform disorder according to DSM IV;
- 3. Patients experience a first or second psychotic episode;
- 4. Age is between 18 and 30 years;
- 5. No current use of clozapine;
- 6. Patients must be reliable. They must agree to co operate with all tests and examinations required by the protocol.

Exclusion criteria

- 1. Pregnancy;
- 2. Lactating women;
- 3. Female subject without adequate contraconception;
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- 4. Known hypersensitivity to any ingredient of olanzapine or risperidone;
- 5. Concomitant use of any other antipsychotic drug than olanzapine or rsiperidone;
- 6. Patients are not allowed to have received depot- antipsychotics for a period of at least 3 months prior to the study;
- 7. Use of other psychotropic medication other than oxazepam or biperideen;
- 8. Narrow-angle glaucoma;
- 9. Known neurological or endocrine disease.

Study design

Design

Study type: Interventional

Intervention model: Parallel

Allocation: Randomized controlled trial

Masking: Double blinded (masking used)

Control: Active

Recruitment

NI

Recruitment status: Recruitment stopped

Start date (anticipated): 01-07-2003

Enrollment: 120

Type: Actual

Ethics review

Positive opinion

Date: 22-11-2004

Application type: First submission

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

RegisterIDNTR-newNL4NTR-oldNTR28Other: N/A

ISRCTN ISRCTN46365995

Study results

Summary results

N/A