Highly active anti-retroviral therapy including nevirapine once daily versus twice daily after at least 12 weeks of nevirapine twice daily. A randomized, open, multicentre trial.

Submission date 13/07/2006	Recruitment status No longer recruiting	 Prospectively registere Protocol
Registration date 28/07/2006	Overall study status Completed	 [] Statistical analysis plan [X] Results
Last Edited 08/01/2021	Condition category Infections and Infestations	Individual participant da

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s) Scientific

Contact name Dr Daniel Podzamczer

Contact details

HIV Unit Infectious Disease Service Hospital Universitari de Bellvitge Feixa Llarga s/n L'Hospitalet de Llobregat Barcelona Spain 08907 +34 (0)93 260 7668 dpodzamczer@csub.scs.es

Additional identifiers

EudraCT/CTIS number

IRAS number

- ed
- ٦
- lata

ClinicalTrials.gov number

Secondary identifying numbers NODy-03

Study information

Scientific Title

Highly active anti-retroviral therapy including nevirapine once daily versus twice daily after at least 12 weeks of nevirapine twice daily. A randomized, open, multicentre trial.

Acronym

NODy

Study objectives

Patients tolerating a standard nevirapine regimen for at least 12 weeks will not present greater hepatic toxicity if switched to a once daily regimen comparing with continuing the standard twice a day (bid) regimen.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Approved 18/12/2003 by the Medicine Spanish Agency and the ethics boards of all participating hospitals.

Study design Randomized, open, multicentre trial

Primary study design Interventional

Secondary study design Randomised controlled trial

Study setting(s) Not specified

Study type(s) Treatment

Participant information sheet

Health condition(s) or problem(s) studied

Human immunodeficiency virus (HIV) infection

Interventions

Patients will be stratified according to whether their CD4 level is more than, equal to or less than 200 cells/ul and whether they are hepatitis C virus (HCV) positive or negative, and centrally randomized to one of these arms:

- 1. Switch to nevirapine 400 mg once daily
- 2. Continue with nevirapine 200 mg bid

Intervention Type

Drug

Phase Not Specified

Drug/device/biological/vaccine name(s)

Nevirapine

Primary outcome measure

Proportion of patients with ALT or aspartate aminotransferase (AST) more than or equal to grade three (more than five times above normal values)

Secondary outcome measures

 Time to ALT and time to AST to reach more than five times above baseline values
 Virological (virological rebound), immunological (CD4 response) and clinical (progression to acquired immune deficiency syndrome [AIDS]) efficacy
 Clinical hepatitis

Overall study start date

30/04/2004

Completion date

30/12/2006

Eligibility

Key inclusion criteria

1. Human immunodeficiency virus (HIV)-positive confirmed by Western blot

2. Adult 18 years or over

3. Under treatment with a highly active anti-retroviral therapy (HAART) regimen including nevirapine 200 mg bid for at least 12 weeks. Females with cluster of differentiation subset four molecules (CD4) >250 cells/ul need to have been receiving the nevirapine bid regimen for at least 18 weeks.

- 4. Alanine aminotransferase (ALT) <2.5 times the upper limit normal
- 5. Undetectable viral load (with the test used in each center)

6. Written informed consent

Participant type(s)

Patient

Age group Adult

Lower age limit 18 Years **Sex** Both

Target number of participants 308 (154 per arm)

Total final enrolment 289

Key exclusion criteria

1. Concomitant participation in another clinical trial

- 2. Clinical suspicion of hepatic cirrhosis
- 3. Renal failure with creatinine clearance <50 ml/min
- 4. Any of the following laboratory parameter alterations: amylases more than three times above normal values, haemoglobin <8 mg/dl, neutrophils <500 cells/ul, platelets <30,000/ul

5. Pregnancy

- 6. Active infection within the last four weeks
- 7. Treatment for neoplasms
- 8. Treatment with methadone

Date of first enrolment

30/04/2004

Date of final enrolment 30/12/2006

Locations

Countries of recruitment Spain

Study participating centre HIV Unit Barcelona Spain 08907

Sponsor information

Organisation

Institute of Biomedical Investigations of Bellvitge (Institut d'Investigació Biomèdica de Bellvitge) (IDIBELL) (Spain)

Sponsor details

Av. Gran via s/n km 2,7 L'Hospitalet de Llobregat Barcelona Spain 08907

Sponsor type Hospital/treatment centre

Website http://www.idibell.es

ROR https://ror.org/0008xqs48

Funder(s)

Funder type Industry

Funder Name Boehringer Ingelheim, Spain

Results and Publications

Publication and dissemination plan Not provided at time of registration

Intention to publish date

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?
<u>Results article</u>	results	01/04/2009	08/01/2021	Yes	No