

Informed Consent Questionnaire

Instructions:

- *This questionnaire is intended to establish your understanding of the clinical trial that you are enrolling in. It will NOT affect your eligibility to enroll into the study.*
- *The questionnaire is anonymous- please DO NOT write your name (or any part of it) on the questionnaire*
- *Please tick the correct answer with a "x"*
- *Note that there may be more than one correct answer*
- *Thank you for completing the questionnaire*

ARM

- ☐ ICD Only
- ☐ ICD & QA
- ☐ ICD & Presentation
- ☐ Presentation & QA
- ☐ ICD, Presentation & QA

1 Questionnaire Number:

2 Highest level of Education:

Primary school	High school	Tertiary education	Illiterate
<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

3 Race:

Black	White	Coloured	Asian
<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

4 Date of Birth:

Day (dd)	Month (mm)	Year (yy)
<input type="text"/>	<input type="text"/>	<input type="text"/>

5 Gender:

Male	Female
<input type="checkbox"/>	<input type="checkbox"/>

1. Participation in this clinical trial is:Required by the
South-African
Medicines Control
Council (MCC)☐

Voluntary

☐Required by the
sponsoring company☐Required by South-
African government☐**2. If you withdraw from the clinical trial you will lose the following benefits you would otherwise be entitled to:**Partial
compensation☐

All compensation

☐

Follow-up care/visits

☐

None

☐**3. The purpose of this trial is to establish:**The difference
between 3
Efavirenz capsule
products☐If the product to be
tested (Efavirenz) is not
safe☐To prove that Efavirenz
is the best☐To prove that Efavirenz
needs to be used in
conjunction with
Neurontin☐**4. The following staff will know which volunteers has received which Efavirenz product:**

Clinic staff only

☐

Laboratory staff only

☐Both clinic staff and
Laboratory staff☐

Nobody

☐**5. What is the duration of this study?**

19 days

☐

135 days

☐

26 days

☐

53 days

☐**6. You will be compensated for your participation in the following way:**

R6570.00

☐R6570.00 and petrol
money☐Food, drink and
accommodation☐None, participation is
voluntary☐

7. After the trial:

You will have to read the newspaper to get the results

☐

The results will be broadcasted on T.V.

☐

You will be informed of all findings related to your individual case

☐

You will be informed in writing about the results of the trial.

☐
8. You are entitled to:

Nothing- you have been paid

☐

The right of access to all the data collected during this trial

☐

The right to all laboratory samples taken for all other volunteers

☐

The right of access to your data

☐
9. You may experience:

Bladder infection

☐

High Blood Pressure

☐

Ear Ache

☐

Drowsiness, sleeplessness

☐
10. As a result of your participation, you will:

Receive the newest care available

☐

Not have access to a Doctor

☐

Have a complete medical evaluation

☐

Become famous

☐
11. As a result of this study:

You might contract HIV

☐

The treatment of HIV will be improved

☐

You will never contract HIV

☐

People in Africa will gain access to medicines which prevents HIV

☐
12. After the study you will

Receive medication free of charge for 3 years

☐

Be phoned to participate in a follow-up study

☐

Follow-up investigations will be conducted within 72 hours

☐

Never have HIV

☐

13. Currently, the following alternative treatment is available for HIV:

Hypnosis

☐

Acupuncture

☐Efavirenz 600mg once
daily☐No alternative
treatment is available☐**14. Your privacy will be protected in the following way:**Your telephone
number will not be
recorded
anywhere at all☐Only your participant
number will be used
during analysis☐Staff will have access
to your personal details☐By confirming your
details with your next of
kin☐**15. Any personal results from this study**Will not be
communicated to
anybody without
your permission☐Will be communicated
to your next of kin only☐Will be communicated
to your insurance
company☐Will be communicated
to anybody who needs
to know☐**16. This study is sponsored by:**

GSK

☐

Novartis CH

☐

Adcock Ingram Ltd.

☐South-African
Government☐**17. Any records/data collected from you during the course of this study will**Be made
available for all
follow-up studies
as well☐Not be used for any
other purpose other
than this study☐May be used to
develop training
material☐May be used to collect
national statistics☐**18. After the clinical trial, your biological samples (i.e. blood or urine) will be:**Stored for 10
years☐

Stored for 15 years

☐

Stored for 5 years

☐

Destroyed

☐

19. A commercial product will be developed from your biological sample:

True

☐

False

☐**20. The doctor involved in the study will be acting as:**Your physician
only☐

The study doctor only

☐The study doctor and
your physician☐

None

☐**21. It is the Study doctor's responsibility to**Provide only
seizure related
care☐

Protect life and health

☐Protect, life, health and
privacy☐Protect, life, health,
dignity and privacy☐**22. Should you experience any adverse effects due to your participation to this study, you will:**Receive medical
care free of
charge until you
are healthy again☐Have to pay for 10% of
your medical care☐Your medical aid will
have to pay for your
medical care☐

Receive nothing

☐**23. Should you die or be disabled as a result of your participation in this study, you or your family or dependants will receive:**R100 000 from
the Medicines
Control Council
(MCC)☐

No compensation

☐R100 000 from Adcock
Ingram SA☐Insurance coverage
from Santam to
indemnify you of any
costs☐**24. The right to compensation is legally guaranteed:**

True

☐

False

☐

25. The protocol for this study has been approved by the Ethics Committee for Medical Research of the University of the Freestate and the South-African Medicines Control Council:

True

☐

False

☐