## **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 & Presentat	ion		
	Presentation & 0	QA		
	ICD, Presentatio	on & QA		
1	Questionnaire Nu	mber:		
2	Highest level of E	ducation:		
	Primary school	High school	Tertiary education	Illiterate
3	Race:			
	Black	White	Coloured	Asian
4	Date of Birth:			
	Day (dd)	Month (mm)	Year (yy)	l
5	Gender:			
	Male	Female		

1.	. Participation in this clinical trial is:			
N	Required by the South-African ledicines Control Council (MCC)	Voluntary	Required by the sponsoring company	Required by South- African government
2.	<ol> <li>If you withdraw from the clinical trial you will lose the following benefits you would otherwise be entitled to:</li> </ol>			
	Partial compensation	All compensation	Follow-up care/visits	None
3.	3. The purpose of this trial is to establish:			
E	The difference between 3 favirenz 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.	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.	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 experi	ence:		
Bladder infection	High Blood Pressure	Ear Ache	Drowsiness, sleeplessness
10. As a result of yo	our participation, you wi	II:	
Receive the newest care available	Not have access to a Doctor	Have a complete medical evaluation	Become famous
11. As a result of th	is 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 wil	I be protected in the foll	owing 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 res	sults 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 invo	lved 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 d	octor'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