

NIDA Clinical Trials Network

Demographics (DEM)

Web Version: 1.0; 1.00; 06-03-11

1. Date of birth:(DEBRTHDT)

 (mm/dd/yyyy)

2. Sex:(DEGENDER)

 Male Female Participant chooses not to answer

3. Ethnicity:(DEETHNIC)

 Hispanic or Latino Not Hispanic or Latino Participant chooses not to answer

4. Race:

American Indian or Alaska Native(DEAMEIND)

 No Yes

Asian(DEASIAN)

 No Yes

Black or African American(DEBLACK)

 No Yes

Native Hawaiian or Pacific Islander(DEHAWAII)

 No Yes

White(DEWHITE)

 No Yes

Other(DEOTHER)

 No Yes

If "Yes", specify:(DEOTHRSP)

OR

Unknown(DEUNKNOW)

 Yes

Participant chooses not to provide their race(DENORACE)

 Yes

Comments:(DEMCOMM)

NIDA Clinical Trials Network

0049Z (ENR)

Web Version: 1.0; ; 09-16-13

Date of assessment: (R6ASMDT)

(mm/dd/yyyy) [Click here for calendar](#)

Inclusion Criteria

In order to meet eligibility ALL Inclusion answers must be "Yes".

1. Participant is at least 18 years old:(R6PTAGE) No Yes Unknown
2. Participant signed HIPAA and/or ROI to abstract hospital records to verify CD4 and VL eligibility criteria: (R6HIPAA) No Yes Unknown
3. Participant reports living in the vicinity and is able to return for follow-up visits:(R6LIVRET) No Yes Unknown
4. Participant is able to communicate in English:(R6ENGLSH) No Yes Unknown
5. Participant was admitted to a hospital and was HIV-infected at the time of recruitment:(R6ADMIT) No Yes Unknown
6. Participant has a Karnofsky performance scale index score of greater than or equal to 60:(R6KARNOF) No Yes Unknown
7. Participant reports any opioid and/or stimulant and/or heavy alcohol use within the past year:(R6DRGUSE) No Yes Unknown
8. Participant has an indication of any opioid and/or stimulant and/or heavy alcohol use within the past 12 months:(R6DRGUSE)
 - a. If female, the AUDIT-C score is greater than or equal to 3 or if male, the AUDIT-C score is greater than or equal to 4:(R6AUC) No Yes Unknown
 - b. In the past year, participant has used one of the following for non-medical reasons: Ecstasy, Heroin, Methamphetamine, Powdered Cocaine, Rock Cocaine, or Recreational use of prescription drugs or pain killers to get high:(R6SUB) No Yes Unknown
 - c. The hospital system medical record for this participant has evidence of opioid or stimulant use in the past 12 months:(R6SUBHR) No Yes Unknown
 - d. The hospital system medical record for this participant has evidence of heavy alcohol use in the past 12 months:(R6AUCHR) No Yes Unknown
9. Participant has a detectable (>200 copies/mL) viral load or unknown level in the past 6 months: (R6VRLOAD)
 - a. If "Yes" or "No", specify viral load: (R6LOADVL) (xxxxxxx) copies/mL -or- (R6VRUNK) Unknown
 - b. Date viral load obtained:(R6LOADDT) (mm/dd/yyyy)
10. Participant has a baseline CD4 count <350 cells/uL in the past 6 months:(R6BSECD4)
 - a. If "Yes" or "No", specify CD4 count (R6CD4VAL) (xxxxx) cells/uL
 - b. Date CD4 count obtained:(R6CD4DT) (mm/dd/yyyy)
11. Participant meets one or more of the following:(R6VLCD4)
 - a. Participant has an AIDS-defining illness during the current hospital admission:(R6AIDS) No Yes Unknown
 - b. Within the past 6 months, participant's most recent CD4 count performed is less than 350 cells/uL and viral load is greater than 200 copies/mL:(R6VLCD6) No Yes Unknown
 - c. Within the past 12 months, participant's most recent CD4 count performed is less than or equal to 500 cells/uL and viral load is greater than 200 copies/mL or unknown **AND** the Site PI's discretion indicates that the participant is likely to currently have a viral load greater than 200 copies/mL, is not currently successfully/correctly taking ART, and needs to be on ART:(R6VLCD12) No Yes Unknown

If the above criteria are "Yes," the patient is eligible to enroll in the study (proceed with main consent and locator information form).

12. Participant provided informed consent for baseline assessments: (R6INFORM)

a. Date informed consent signed for baseline assessment: (R6CNSTDT)

b. If "No", specify: (R6CNSNO)

No Yes

(mm/dd/yyyy) [Click here for calendar](#)

If "Judgment of study personnel", specify: (R6CNSJUD)

If "Other", specify: (R6CNSOSP)

c. Provided consent for audio recording: (R6INFAUD)

d. Provided consent to be contacted for optional future studies: (R6INFFUT)

No Yes

No Yes

No Yes Unknown

No Yes Unknown

No Yes

13. Participant provides sufficient locator information: (R6LCATOR)

14. A baseline blood draw has been completed for this participant: (R6BLOOD)

15. Participant has completed the baseline CAPI assessments: (R6CAPI)

Exclusion Criteria

In order to meet eligibility ALL Exclusion answers must be "No".

1. Participant has a significant cognitive or developmental impairment to the extent that they are unable to provide informed consent: (R6SIGCOG)

No Yes Unknown

2. Participant is terminated via site PI decision with agreement from study LI: (R6TERM)

No Yes Unknown

If "Yes", specify: (R6TERMSP)

Eligibility for Randomization

1. Is the participant eligible for the study? (R6PTELIG)

No Yes

2. Is the participant eligible for randomization? (R6PTRAND)

No Yes

a. If "No", specify: (R6NORASP)

b. If "Judgment of CTP/research staff", specify: (R6JUDGSP)

c. If "Other", specify: (R6OTHRSP)

Comments:(R6COMM)

