NIDA Clinical Trials Network

Demographics (DEM)

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

. Date of birth: (DEBRTHDT)	(mm/dd/yyyy)
. Sex:(DEGENDER)	☐ Male ☐ Female ☐ Participant chooses not to answer
. Ethnicity: (DEETHNIC)	☐ Hispanic or Latino ☐ Not Hispanic or Latino ☐ Participant chooses not to answer
Race: American Indian or Alaska Native(DEAMEIND) Asian(DEASIAN) Black or African American(DEBLACK) Native Hawaiian or Pacific Islander(DEHAWAII) White(DEWHITE) Other(DEOTHER) If "Yes", specify:(DEOTHRSP)	No Yes No Yes No Yes No Yes No Yes No Yes
OR	
Unknown(DEUNKNOW) Participant chooses not to provide their race(DENORACE)	☐ Yes ☐ Yes
Comments: (DEMCOMM)	

NIDA Clinical Trials Network
00497 (FNR)

Web Version: 1.0; ; 09-16-13

	0049Z (ENR)
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
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
 Participant reports any opioid and/or stimulant and/or heavy alcohol use within the past year: (R6DRGUSE) 	□ No □ Yes □ Unknown
 Participant has an indication of any opioid and/or stimulant and/or heavy alcohol use within the past 12 months:(R6DRGUSE) 	□ No □ Yes □ Unknown
 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) 	
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:(R6A UCHR)	□ No □ Yes □ Unknown
 Participant has a detectable (>200 copies/mL) viral load or unknown level in the past 6 months: (R6VRLOAD) 	□ No □ Yes □ Unknown
a. If "Yes" or "No", specify viral load: (R6LOADVL)	(xxxxxxxx) copies/mL -or- (R6 VR UNK) ☐ 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)	□ No □ Yes □ Unknown
a. If "Yes" or "No", specify CD4 count: (R6CD4 VAL)	(xxxxx) cells/uL
b. Date CD4 count obtained:(R6CD4DT)	(mm/dd/yyyy)
11. Participant meets on e or more of the following: (R6VLCD4)	□ No □ Yes
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:(R6VL CD6)	
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 Pl'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)	

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)	□ No □ Yes
a. Date informed consent signed for baseline assessment:(R6CNSTDT)	(mm/dd/yyyy) Click here for calendar
b. If "No", specify: (R6CNSNO)	
If "Judgment of study personnel", specify: (R6CNSJUD)	
If "Other", specify:(R6CNSOSP)	
c. Provided consent for audio recording:(R6INFAUD)	□ No □ Yes
d. Provided consent to be contacted for optional future studies: (R6INFFUT)	□ No □ Yes
13. Participant provides sufficient locator information:(R6L CATOR)	□ No □ Yes □ Unknown
14. A baseline blood draw has been completed for this participant: (R6BLO OD)	□ No □ Yes □ Unknown
15. Participant has completed the baseline CAPI assessments: (R6CAPI)	□ No □ Yes
Exclusion Criteria	
In order to meet eligibility ALL Exclusion answers must be "No".	
4. Dadicional han a similar and a south and the same and similar and the state of the same and t	
 Participant has a significant cognitive or developmental impairment to the extent that they are unable to provide informed consent: (R6SIGCOG) 	□ No □ Yes □ Un kno wn
2. Participant is terminated via site PI decision with agreement from study LI: (R6TERM)	□ No □ Yes □ Unknown
If "Yes", specify:(R6TERMSP)	
Flinibility for Dondomination	
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)	NO E les
b. If "Judgment of CTP/research staff", specify:(R6JUDGSP)	
c. If "Other", specify:(R60THRSP)	
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Comments:(R6COMM)	