HPTN 065
TLC-Plus: A Study to Evaluate the Feasibility of an Enhanced Test, Link to Care, Plus Treat Approach for HIV Prevention in the United States

A Study of the HIV Prevention Trials Network

Sponsored by:
Division of AIDS (DAIDS), U.S. National Institute of Allergy and Infectious Diseases (NIAID)
U.S. National Institutes of Health (NIH)

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LIST OF ABBREVIATIONS AND ACRONYMS

AA   African American
ACASI  Audio Computer-Assisted Self Interview
ACTG  AIDS Clinical Trials Group
AIDS  Acquired Immunodeficiency Syndrome
ADAP  AIDS Drug Assistance Program
ALIVE  AIDS Link to Intravenous Experience
AMPATH  Academic Model for the Prevention and Treatment of HIV/AIDS
ART  Antiretroviral Therapy
ARTAS  Antiretroviral Treatment Access Study
ASD  Adult/Adolescent Spectrum of HIV Disease
ASO  AIDS Service Organization
BRFSS  Behavioral Risk Factor Surveillance System
CARE+  Computer Assessment and Risk Reduction Education for HIV-positives
CATI  Computer-Assisted Telephone Interviewing
CBO  Community-Based Organization
CD4  surface glycoprotein that denotes helper T cells
CDC  Centers for Disease Control and Prevention
CHS  Community Health Survey
CLIA  Clinical Laboratory Improvement Amendments
CORE  Coordinating and Operations Center
CRF  Case Report Form
CSTE  Counsel of State and Territorial Epidemiologists
CTRS  Counseling, Testing and Referral Services
DAIDS  Division of AIDS
DC  District of Columbia
DHHS  Department of Health and Human Services
DOB  Date of Birth
DOC  Department of Corrections
DOH  Department of Health
DOHMH  Department of Health and Mental Hygiene
ED  Emergency Department
eHARS  enhanced HIV/AIDS Reporting System
EIA  Enzyme Immunoassay
EMR  Electronic Medical Record
FI  Financial Incentive
FSU  Field Services Unit
FY  Fiscal Year
GE  General Electric
GEE  Generalized Estimating Equations
HbA1c  Hemoglobin A1C
LIST OF ABBREVIATIONS AND ACRONYMS (continued)

HAART  Highly Active Antiretroviral Therapy
HAHSTA  HIV/AIDS, Hepatitis, STD and TB Administration
HARS  HIV/AIDS Reporting System
HCPI  Health Communications/Public Information
HCSUS  HIV Cost and Services Utilization Study
HHS  Health and Human Services
HIPAA  Health Insurance Portability and Accountability Act
HIV  Human Immunodeficiency Virus
HOPS  HIV Outpatient Study
HOPWA  Housing Opportunities for People with AIDS
HPTN  HIV Prevention Trials Network
HRSA  Health Resources and Services Administration
ICC  Intra-class Correlation
ICF  Informed Consent Form
ICT  Information and Communication Technologies
ID  Identification
IDSA  Infectious Disease Society of America
IDU  Injecting Drug User
INR  International Normalized Ratio
IPV  Intimate Partner Violence
IRB  Institutional Review Board
JAIDS  Journal of Acquired Immune Deficiency Syndromes
LoA  Letter of Amendment
MEMS  Medication Event Monitoring System
MCO  Managed Care Organization
MMP  Medical Monitoring Project
MMP  Morbidity Monitoring Project
MMWR  Morbidity and Mortality Weekly Report
MSM  Men who have Sex with Men
MTRH  Moi Teaching and Referral Hospital
NAAT  Nucleic Acid Amplification Testing
NHANES  National Health and Nutrition Examination Survey
NHBS  National HIV Behavioral Surveillance
NHBS-HET  National HIV Behavioral Surveillance – At Risk Heterosexuals
NHBS-IDU  National HIV Behavioral Surveillance – Injection Drug Users
NHBS-MSM  National HIV Behavioral Surveillance – Men who have Sex with Men
NHIS  National Health Interview Survey
NIAID  National Institute of Allergy and Infectious Diseases
NIH  National Institutes of Health
NIMH  National Institutes of Mental Health
NL  Network Laboratory
NY  New York
LIST OF ABBREVIATIONS AND ACRONYMS (continued)

NYC New York City
OARAC Office of AIDS Research Advisory Committee
OHRP Office for Human Research Protections
PEMS Program Evaluation and Monitoring System
PHMC Public Health Management Corporation
PLWHA People Living With HIV/AIDS
PRO Protocol Registration Office
RSC Regulatory Support Center
RCT Randomized Controlled Trial
RFA Request for Applications
RNA Ribonucleic Acid
SCHARP Statistical Center for HIV/AIDS Research and Prevention
SDMC Statistics and Data Management Center
SES Socio-economic Status
SHAS Supplement to HIV/AIDS Surveillance
SMART Strategies for Management of Antiretroviral Therapy
SMC Study Monitoring Committee
SOC Standard of Care
SQL Structured Query Language
SSP Study-Specific Procedures Manual
STD Sexually Transmitted Disease
TA Technical Assistance
TB Tuberculosis
TLC Test and Link to Care
UODT Universal Offer Development Team
U.S. United States
USPHS United States Public Health Service
USA United States of America
VL Viral Load
WB Western Blot
WHO World Health Organization
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INVESTIGATOR SIGNATURE PAGE

A Study of the HIV Prevention Trials Network (HPTN)

Sponsored by:
Division of AIDS (DAIDS), U.S. National Institute of Allergy and Infectious Diseases (NIAID)
U.S. National Institutes of Health (NIH)

I, the Investigator of Record, agree to conduct this study in full accordance with the provisions of this protocol. I agree to maintain all study documentation for a minimum of three years after submission of the site’s final Financial Status Report to the Division of AIDS (DAIDS), unless otherwise specified by DAIDS or the HIV Prevention Trials Network (HPTN) Coordinating and Operations Center. Publication of the results of this study will be governed by HPTN policies. Any presentation, abstract or manuscript will be made available by the investigators to the HPTN Manuscript Review Committee and DAIDS for review prior to submission.

I have read and understand the information in this protocol and will ensure that all associates, colleagues and employees assisting in the conduct of the study are informed about the obligations incurred by their contribution to the study.

__________________________________
Name of Investigator of Record

__________________________________   _________________________________
Signature of Investigator of Record   Date
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TLC-Plus: A Study to Evaluate the Feasibility of an Enhanced Test, Link to Care, Plus Treat Approach for HIV Prevention in the United States

SCHEMA

Purpose: The main purpose of this study is to evaluate the feasibility of an enhanced community-level test, link to care, plus treat strategy in the United States. The study includes the following components:

- Expanded Human Immunodeficiency Virus (HIV) Testing
- Linkage-to-Care
- Viral Suppression
- Prevention for Positives
- Patient and Provider Surveys

Design: Each component of the study involves an independent design but is interrelated to the other components.

- The Expanded HIV Testing component involves social mobilization, with targeted messaging to promote testing, and implementation of the universal offer of HIV testing in emergency departments (EDs) and hospital inpatient admissions.

- The Linkage-to-Care and Viral Suppression components involve site randomization to test the effectiveness of a financial incentive (FI) intervention compared with the standard of care (SOC).

- The Prevention for Positives component uses individual randomization to compare the SOC plus a computer-delivered intervention with the SOC.

- The Patient and Provider Surveys will be administered at specific time points during the study to assess knowledge, attitudes and practices regarding early initiation of antiretroviral therapy (ART) and the FI interventions.

Study Sites: The study will be conducted in two intervention communities (the Bronx, New York and Washington, D.C.) and surveillance data from these communities will be compared with that from four non-intervention communities (Chicago, Illinois; Houston, Texas; Miami, Florida; and Philadelphia, Pennsylvania).

Study Population: This study will primarily target individuals age 18 years and older, and will also include younger individuals who are legally able to consent for HIV testing and care according to the state or local law in the two study communities. All of the study components will include HIV-positive individuals, with the exception of the provider survey sub-component.

Study Objectives: The main objective of the study is to determine the feasibility of a community focused enhanced test and link-to-care strategy in the United States. The study includes feasibility objectives for the Expanded HIV Testing, Linkage-to-Care and Viral Suppression components, and effectiveness objectives for the Linkage-to-Care, Viral Suppression and Prevention for Positive components.

Study Duration: The study will take place over 36 months.
Study Components:

The study includes the following five components:

**Expanded HIV Testing:** The expanded HIV testing intervention will supplement ongoing social mobilization and HIV testing efforts already in place in the two intervention communities. Refined messages will be added to ongoing social marketing and social networking efforts, targeted to increase HIV testing and testing frequency among men who have sex with men (MSM) and other subpopulations disproportionately affected by HIV. The intervention for EDs and hospital admissions includes encouraging hospital leadership and staff to institute standing orders for universal HIV testing. It also includes providing financial support for an increased number of HIV tests, and use of novel mechanisms to deliver information about HIV testing to patients.

**Linkage-to-Care:** HIV test sites will be randomized to either the FI intervention or to the SOC for linkage of HIV-positive patients identified at the testing sites to HIV care sites. At HIV test sites assigned to FIs, HIV-positive patients will be provided with a coupon to redeem at participating HIV care sites in the community. Upon completion of HIV laboratory testing, patients with coupons will be given an FI ($25 gift card) at an HIV care site. A $100 gift card will be provided to patients upon completion of a care visit that includes interaction with a healthcare provider and discussion of HIV laboratory test results (e.g. CD4 cell count and viral load (VL) measurements).

**Viral Suppression:** HIV care sites will be randomized to either the FI intervention or the SOC for the achievement and maintenance of viral suppression. HIV care sites assigned to the FI intervention will provide an FI ($70 gift card) to HIV-positive patients on antiretroviral therapy (ART) demonstrating a suppressed VL (as defined by <400 copies/mL) at quarterly care visits.

**Prevention for Positives:** In a subset of patients enrolled from select HIV care sites in the two intervention communities, patients will be randomized either to an intervention arm (receiving SOC prevention activities plus a computer-delivered intervention for sexual and behavioral risk reduction) or to the control arm (receiving only the SOC prevention activities at the care site). In the intervention arm a modified version of the computerized counseling platform called **Computer Assessment and Risk Reduction Education for HIV-positives (CARE+)** will be used with an Audio Computer Assisted Self-Interviewing technique (ACASI). The CARE+/ACASI intervention session will ascertain behavioral risk, assess self-efficacy/motivation, and provide tailored feedback on specific risk behaviors. The computer-delivered intervention session will be administered every three months for one year. Participants in the control arm will also have CARE+/ACASI sessions every three months. However, participants in the control arm will only be administered behavior assessments via CARE+/ACASI and will not receive prevention messaging.
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SCHEMA (Continued)

Study Components (continued):

Patient and Provider Surveys: The same computer system used for the Prevention for Positives intervention will be used to administer a survey to patients enrolled in that study component. The survey will assess their knowledge and attitudes towards ART use for treatment and prevention, ART adherence and FIs. Providers from these sites will be invited to complete a Web-based survey regarding their knowledge, attitudes and practices concerning ART for treatment and prevention as well as use of FIs. Both surveys will also collect some key sociodemographic data on surveyed populations to allow more in depth characterization of these populations.

Study Size: Each of the five study components has a target sample size.

- The universal offer of HIV testing will be made in EDs and during hospital admission at ~seven facilities in the Bronx, NY and ~seven in Washington, D.C. Additional focused messages promoting testing will be targeted to the entire population of each intervention community: the Bronx (population 1.4 million) and Washington, D.C. (population 600,000).

- The Linkage-to-Care component includes 40 HIV test sites (20 in each intervention community) and 40 HIV care sites (20 in each intervention community). We project that, by the end of the study, approximately 3000 new individuals in the two intervention communities will have tested positive for HIV.

- The Viral Suppression component includes the 40 HIV care sites (20 in each intervention community). Throughout the study duration, approximately 30,000 HIV-positive individuals will be in care, with an estimated 75% (22,500) eligible for ART in the two intervention communities.

- The Prevention for Positives component will be conducted at a total of twelve sites (six in each intervention community) with up to a total of 1320 patients participating.

- The patients at the twelve sites participating in the Prevention for Positives component will be surveyed. Providers at all participating HIV care sites will be invited to complete the provider survey.
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OVERVIEW OF STUDY DESIGN AND RANDOMIZATION SCHEME

### Summary of Study Components

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STUDY TIMELINE

Year 1

Year 2

Year 3

Expanded HIV Testing Component (36 m)

Linkage-to-Care Component (24 m)

Provider Training and Survey

Viral Load Suppression Component (24 m)

Provider Survey

Patient Survey

Prevention for Positives Component (18 m)

Patient Survey
1.0 INTRODUCTION

1.1 Background and Prior Research

1.1.1 The Test-and-Treat Approach

The test-and-treat (TNT) strategy to prevent human immunodeficiency virus (HIV) transmission is based on expanded HIV testing to identify undiagnosed HIV infection, combined with prompt and effective initiation of ART to lower HIV viral load (VL) levels. While the potential effect of ART on HIV transmission has been previously studied by several investigators (Cohen, Gay et al. 2007), it received renewed attention in 2009. Granich and colleagues at the World Health Organization (WHO) published results of a modeling exercise that assessed the potential effect of such an approach on the HIV epidemic in South Africa (Granich, Gilks et al. 2009). The assumptions in the model included annual HIV testing of all adults older than 15 years of age, prompt initiation of ART in all those infected irrespective of disease stage, and a 99% decrease in infectiousness with highly effective first-line ART. The model indicated a dramatic drop in HIV incidence within 10 years and reduction of HIV prevalence to 1 percent in 50 years. The results of this modeling exercise have generated interest in the evaluation of such an approach (Assefa and Lera 2009; Cohen, Mastro et al. 2009; Dieffenbach and Fauci 2009; Epstein 2009; Granich, Gilks et al. 2009; Hsieh and de Arazoza 2009; Jurgens, Cohen et al. 2009; Wagner and Blower 2009).

The TNT strategy is hypothesized to achieve its effect on HIV transmission through the following two pathways:

- HIV testing identifies HIV-positive persons who, after learning their status, adopt safer behaviors, which decreases HIV transmission.

- HIV-positive individuals who initiate ART, and then maintain high levels of adherence and achieve viral suppression, are less infectious, which decreases HIV transmission.

Figure 1: Test and Treat Concept
HIV screening has been found to be as cost-effective as other routine health interventions (Paltiel, Weinstein et al. 2005; Sanders, Bayoumi et al. 2005) and initiatives to expand HIV screening are underway in the United States. However, to optimize health outcomes, expanded testing efforts must be coupled with initiatives that ensure that both the newly diagnosed and those already known to have HIV infection are effectively linked to HIV care (Walensky, Weinstein et al. 2005) and receive ART as indicated with optimal adherence and suppression of viral replication.

The use of ART for prevention is supported by observational data from discordant couples that suggest that use of ART by the HIV sero-positive partner is associated with lower HIV incidence in the sero-negative partner (Quinn, Wawer et al. 2000; Sullivan, Kayitenkore et al. 2009). Other data also indicate a decrease in rates of HIV infection with use of ART (Bunnell, Ekwaru et al. 2006). However, definitive data for the effect of ART on transmission awaits the outcome of HIV Prevention Trials Network (HPTN) 052/AIDS Clinical Trials Group (ACTG) 5245. In addition, if ART is to be used widely for prevention purposes, its risk/benefit needs to be defined for a population not eligible for ART based on current therapeutic guidelines. Definitive data are lacking for the optimal timing of ART initiation for clinical benefit (Wilkin and Gulick 2008). Two ongoing randomized clinical trials (RCTs), the INSIGHT START study and HPTN 052/ACTG 5245, should yield such data on the efficacy and safety of initiation of ART at higher CD4 cell counts than the currently recommended threshold (Panel on Antiretroviral Guidelines for Adults and Adolescents 2009). In HPTN 052, HIV-positive individuals with CD4 count between 350 and 550 cells/mm$^3$ and who have HIV-discordant partners are randomized to initiate ART at CD4 cell count between 350 and 550 cells/mm$^3$ or at CD4 cell count between 200 and 250 cells/mm$^3$. The purpose of the latter study is to determine the effectiveness of ART on the sexual transmission of HIV to the uninfected partner as well as to determine the long-term effectiveness and safety of use of ART. In the INSIGHT START Study, HIV-positive patients with CD4 cell count of $>$500 cells/mm$^3$ are randomized to immediate ART initiation versus deferral of ART to when the CD4 cell count falls below 350 cells/mm$^3$.

1.1.2 HIV Testing in the United States, Washington, D.C., and the Bronx

1.1.2.1 Current HIV Testing in the United States

Based on data from the 2006 National Health Interview Survey (NHIS), the Centers for Disease Control and Prevention (CDC) estimates that 40% of all non-elderly adults (18 - 64 years) in the United States have been tested for HIV. A much smaller percentage of this population (10%) reported testing for HIV within the year prior to the survey. These levels of HIV testing, for both having ever been tested and for those tested in the past year, have remained constant from 2001 to 2006, suggesting that some individuals are tested repeatedly, while the majority of the U.S. population (60%) remains untested.

Approximately 21% of HIV-positive patients in the United States are not aware of their HIV infection (Campsmith, Rhodes et al. 2008) and not only may miss the benefits of HIV care and timely ART but also may continue to transmit HIV to their partners.

To increase the level of HIV testing in the United States and reduce the frequency of late HIV diagnoses, the CDC revised its recommendations for HIV testing in healthcare
settings and several federal agencies have new initiatives to increase testing in specific subpopulations (Duran, Beltrami et al. 2008). In 2006, the CDC recommended opt-out HIV screening as part of routine clinical care for adults and adolescents, including pregnant women, in all healthcare settings (Branson, Handsfield et al. 2006). In order to lower some of the barriers to testing, CDC recommendations require neither separate written consent nor HIV-prevention counseling as mandatory elements of HIV testing.

In September 2007, CDC funded an Expanded Testing Initiative in the 26 U.S. jurisdictions with the largest number of Acquired Immunodeficiency Syndrome (AIDS) cases. Through June 2009, 1.2 million persons have been tested under the Expanded Testing Initiative, with identification of nearly 14,400 new HIV diagnoses, 63% of which were among persons previously unaware they were infected. These tests were performed in Emergency Departments EDs (34%), sexually transmitted disease (STD) clinics (26%), community health centers (18%), corrections facilities (13%), and hospital inpatient settings (1.4%).

1.1.2.2 Current HIV Testing Activities in New York City and the Bronx

In the past three years, the New York City (NYC) Department of Health and Mental Hygiene (DOHMH) has scaled up its efforts to support routine HIV testing throughout its jurisdiction. As part of that effort, “The Bronx Knows” project was conceptualized. This program was initiated in early 2007 with the intention of scaling up HIV testing throughout the borough of the Bronx. After one year of capacity building, “The Bronx Knows” initiative was launched publicly on June 27, 2008 (National HIV Testing Day). The goals of the initiative are three-fold: to ensure that all Bronx residents who have never had an HIV test are screened for HIV over three years, to identify Bronx residents with undiagnosed HIV, and to link these HIV-positive individuals to high quality care and supportive services. While these are the stated goals of the initiative, residents with ongoing risk for HIV acquisition are also encouraged to be tested at least annually, per CDC recommendations.

As part of “The Bronx Knows,” the NYC DOHMH provides free test kits to organizations that offer HIV testing to uninsured individuals and to Community-Based Organizations (CBOs) that do not have dedicated funds for HIV testing. The DOHMH also provides technical assistance (TA) in the following areas: logistical/infrastructure change to support routine HIV screening (including on-site TA), obtaining clinical laboratory improvement amendment (CLIA) waivers, billing/reimbursement, rapid HIV testing technologies and data collection. In addition, the NYC DOHMH conducts sector-based workshops and Webinars in which participants review best practices and barriers experienced in their own sector (hospital, clinic or CBO). Finally, DOHMH provides ongoing social marketing and media campaigns targeting both Bronx residents (promoting routine HIV testing) and providers (promoting routine screening of patients for HIV).

By January 2009, more than 70 organizations had joined “The Bronx Knows.” These include seven of eight Bronx hospitals, 39 community health clinics and 20 of the borough's largest CBOs. Other participants include colleges/universities, faith-based institutions, local commercial establishments and community boards. Preliminary
aggregate data show that HIV testing among reporting organizations has increased by 28% since the launch of the initiative, despite New York State’s requirement of separate signed informed consent for HIV testing.

1.1.2.3 Current HIV Testing Activities in Washington, D.C.

The Washington, D.C. Department of Health (DOH) has made HIV its number one health priority. Efforts have focused on aggressive diagnosis and treatment of HIV throughout Washington, D.C. Washington, D.C. does not require separate signed consent for HIV testing and was the first jurisdiction in the country to commit to and implement a policy of routine, opt-out HIV testing for residents. In June 2006, Washington, D.C. launched its, “Come Together D.C. - Get Screened for HIV” campaign. This campaign promoted testing among residents and recruited new testing organizations and testing sites. Starting in fiscal year 2007 (FY07) and continuing through FY08, testing expansion focused on scaling up routine, opt-out testing in the Department of Corrections (DOC) jail settings as well as in medical settings with special emphasis on hospital EDs, primary medical settings, managed care organizations (MCOs) and CBOs. In 2008, Washington, D.C. began implementation of a six-pronged strategy to scale up routine HIV screening citywide. The goals of this strategy are to: 1) increase the scope and scale of routine screening for HIV in clinical settings; 2) provide HIV testing and referral services, with an emphasis on identifying newly infected persons and ensuring that test results are received; 3) establish models to more efficiently link HIV-positive individuals to care; 4) collaborate with DOCs, medical care entities and CBOs to encourage and support Counseling, Testing and Referral Services (CTRS); 5) collect and analyze data to determine the scope and reach of CTRS programs; and 6) develop and disseminate tools to address myths and barriers related to routine HIV screening.

Despite efforts to increase provider awareness and practice of routinely offering HIV testing, behavioral survey and testing data suggest missed opportunities for routine testing in medical settings are still quite frequent, with nearly 75% of newly diagnosed HIV-positive persons reporting having seen a healthcare provider in the past twelve months without having been diagnosed (NHBS-Het Survey 2009). As a complement to efforts to increase provider participation in routine HIV testing, Washington, D.C.’s expansion efforts also include a direct-to-consumer marketing campaign called “Ask for the Test,” which seeks to drive consumer demand by encouraging clients to ask for HIV testing if it is not routinely offered. It is anticipated that the approaches described above could result in an additional 85,000 tests per year.

1.1.3 HIV Care and ART Utilization in the United States, Washington, D.C. and the Bronx

The most recent population-based estimates on the proportion of HIV-positive Americans who are in HIV care and receiving ART are from 2003. These estimates depend on: 1) the fraction of HIV-positive persons who are undiagnosed and unaware of their HIV infection; 2) the fraction who have been diagnosed with HIV but are not in care; and 3) the changing recommendations with respect to timing of initiation of ART. Using data from the CDC’s national HIV surveillance system and its 10-city Adult/Adolescent Spectrum of HIV Disease (ASD) project, the CDC estimated that, as of 2003, there were
480,000 Americans between the ages of 15 and 49 living with HIV or AIDS who were eligible for ART at a threshold of CD4 cell count < 350 cells/mm$^3$ (Teshale, Kamimoto et al. 2005). Of all eligible persons, approximately 340,000 (71%) were diagnosed with HIV and in care. Only 268,000 (55% of all eligible) were receiving ART. Based on these findings, the CDC recommended that three critical components be incorporated into the national HIV-prevention strategy: 1) increasing the number of HIV-positive individuals who are aware of their status; 2) linking them to HIV-prevention and care services; and 3) increasing the number receiving ART per federal guidelines (Branson, Handsfield et al. 2006). Given the growing number of HIV-infected persons living in the United States, estimated at 1.1 million as of 2006 (Campsmith, Rhodes et al. 2008), and the trend for initiation of therapy at CD4 cell count > 350 cells/mm$^3$ for some subsets of patients (Panel on Antiretroviral Guidelines for Adults and Adolescents 2009), it is likely that a greater number of persons (than the 0.5 million estimated in 2003 by Teshale et al.) are eligible for and in need of ART nationwide today.

In 2007, the CDC launched the Medical Monitoring Project (MMP) based on a national probability sample of HIV-positive persons receiving care in the United States to obtain data on patterns of use and quality of HIV care and prevention services among such persons, and their clinical and virologic status (McNaghten, Wolfe et al. 2007). Preliminary data from the MMP project are expected in 2010. Available data on current patterns of HIV care, ART treatment and ART adherence come primarily from a few large U.S. HIV cohorts (Palella, Delaney et al. 1998; Lazo, Gange et al. 2007; Mugavero, Lin et al. 2009). For instance, the CDC-funded HIV Outpatient Study (HOPS) has shown persistent reductions in mortality and hospitalizations due to increasing use of highly active antiretroviral therapy (HAART) over time (Palella, Delaney et al. 1998; Buchacz, Baker et al. 2008). Data from the HIV Cost and Services Utilization Study (HCSUS) (Cunningham, Markson et al. 2000) and the HIV Research Network (Gebo, JAIDS 2005) point to disparities in use of ART by sociodemographic characteristics (e.g., race, gender and insurance status), some of which appear to persist to the present (Palella, Armon et al. 2008).

Linkage-to-care is required for HIV-positive persons to realize the benefits of both HIV care and prompt initiation of ART. Recently updated Infectious Disease Society of America (IDSA) guidelines for the management of persons with HIV emphasized the importance of linkage and retention in primary care (Aberg, Kaplan et al. 2009). Poor engagement in care has been found to be a predictor of higher mortality. Specifically, HIV-positive persons with poor retention in care have been found to have 50% higher mortality rates (Giordano, Gifford et al. 2007). Appropriate care includes determining stage of HIV disease through clinical evaluation and measurement of CD4 cell count and HIV ribonucleic acid (RNA) levels, other health maintenance interventions such as Pap smears for women, tuberculin skin testing, provision of drugs for prevention of opportunistic infections, health education, risk reduction, supportive counseling and ART for patients eligible for such treatment (Panel on Antiretroviral Guidelines for Adults and Adolescents 2009). However, data from multiple studies indicate both failure and delay in linkage-to-care (Shapiro, Morton et al. 1999; Giordano, Visnegarwala et al. 2005; Tobias, Cunningham et al. 2007). Nationally, various studies estimate that only 60-75% of persons are linked to HIV care within three to six months of receipt of HIV diagnosis (Torian, Wiewel et al. 2008; Reed, Hanson et al. 2009; Zetola, Bernstein et al. 2009).
In the supplement to HIV/AIDS surveillance (SHAS) project, analyses of interview data collected during 2000-2004 from over 20 U.S. cities, counties, and states, indicated that 72% of persons reported entering HIV care within three months of testing HIV-positive (Reed, Hanson et al. 2009). Barriers associated with failure or delayed entry into HIV care include structural, financial and personal/cultural factors. Such factors include first-time HIV testing and anonymous HIV testing (Reed, Hanson et al. 2009), longer waiting time for initial care appointment (Mugavero, Lin et al. 2007) or being diagnosed at earlier stage of HIV disease when patients report “feeling well” (Tobias, Cunningham et al. 2007). In a NYC study using 2003 HIV surveillance data, Torian et al. found that 64% of patients initiated HIV care within three months of HIV diagnosis, 19% initiated care more than three months after diagnosis, and 17% never initiated care. Delay in initiation of care was associated with HIV testing in a community site, in the correctional system, at a sexually transmitted infection or tuberculosis (TB) clinic, non-white race, injection drug use and foreign birth (Torian, Wiewel et al. 2008).

The national five-year multisite Outreach Initiative funded by Health Resources and Services Administration (HRSA) in 2001 identified a number of effective programmatic interventions for engaging and retaining HIV-positive persons in care, including education and outreach, strengthening of referrals, provision of linkage coordination and navigation services, and case management (Rajabiun, Mallinson et al. 2007; Tobias, Cunningham et al. 2007). For instance, an “HIV system navigation” approach was successful in reducing barriers to establishing care and improved health outcomes in a multisite study over 12-month period (Bradford, Coleman et al. 2007). The CDC-funded Antiretroviral Treatment Access Study (ARTAS) strengths-based case management intervention, delivered at CBOs and health departments in 10 sites across the United States during 2005-2006, resulted in 79% of recently diagnosed HIV-positive persons receiving HIV medical care within six months of enrolling in the study (Gardner, Metsch et al. 2005; Craw, Gardner et al. 2008).

### 1.1.3.1 Expected Rates of Viral Suppression

The epidemiology of HIV infection in the United States reveals a disproportionate percentage of African American (AA) people (41%) living with HIV compared to the general population (12%). Outcomes of virologic efficacy in all populations impacted by HIV in the United States have been assessed in studies to determine rates of success of such treatment and its effect on mortality and morbidity. An analysis from 1995 –2001 revealed that mortality rates for HIV-positive white men in the United States declined by 85% compared to 50% and 65% for HIV-positive AA women and men respectively (Prevention 2002). The latter finding may relate to factors such as late diagnosis of HIV, access to ART and challenges in achieving high rates of adherence with treatment.

A number of studies to evaluate these differences between race and virologic efficacy have been published. Weintrob et al. (Weintrob, Grandits et al. 2009) demonstrated significantly lower odds of obtaining virologic suppression in AA compared to Caucasians at six and 12 months in a cohort of military personnel. This group was studied because such individuals are likely to have similar access to medications and care. However, several factors may have still influenced this finding including adherence with ART, provider bias, or differences in education status impacting selection bias.
regarding provider selection of initial therapy, or how long patients stay on a given regimen. Anastos et al. (Anastos, Schneider et al. 2005) reported similar findings in terms of rates of viral suppression in 961 HIV-infected women enrolled in the Women’s Interagency HIV Study (WIHS). AA women were 30% less likely to achieve viral suppression and were also 30% more likely to rebound after achieving viral suppression. The reasons for these differences remains unclear, as subsequent adjustment for multiple variables including biologic and behavioral variables did not eliminate the differences noted by race. There have been other studies showing genetically determined factors that influence drug levels, but studies that link these same factors to virologic success or toxicity have been inconclusive.

Thus, evidence suggests that, in addition to the disproportionate impact of HIV on racial/ethnic minorities in the United States, there are racial disparities in terms of outcomes of HIV disease and these persist in HIV treatment. It is unclear if this disparity in outcomes points toward race as a biologic determinant or as an underlying complex sociologic marker. Nonetheless, the selection of measure of virologic success at an HIV RNA level of < 400 copies/mL is a reasonable measurement that should reflect success of ART in individuals irrespective of racial group.

1.1.3.2 Access to Support Services for Mental Health, Substance Use and Homelessness

Linkage to HIV care is critical for those infected with HIV; however, underlying conditions, such as substance use (Robison, Westfall et al. 2008; Wood, Kerr et al. 2008; Applebaum, Reilly et al. 2009; Norman, Basso et al. 2009), mental illness (Berg, Cooperman et al. 2009; Kapetanovic, Christensen et al. 2009; Roux, Carrieri et al. 2009), and homelessness (Royal, Kidder et al. 2009) can have significant impact on a person’s ability to become successfully linked to HIV care, as well as to adhere to ART. Depression and anxiety are the most common psychiatric diagnoses in HIV-positive persons and are 5-10 times more common in this population than in the general population (Pence 2009). A limited number of randomized clinical trials have demonstrated the beneficial effect of various psychotherapy-based interventions on ART adherence (Weber, Christen et al. 2004; Wyatt, Longshore et al. 2004; Safren, Knauz et al. 2006). In addition, substance use may impact side effects of ART. For example, Cheng et al. noted that subjects with high alcohol use had higher odds of experiencing lipodystrophy (odds = 2.07, adjusted odds ratio = 0.90, 95% confidence interval = 4.73) compared to those with lower alcohol use (Cheng, Libman et al. 2009). It is estimated that at least 13% of HIV patients have mental health and substance use disorders (Weaver, Conover et al. 2009).

Access to supportive services such as substance use management, mental health, homelessness prevention and adherence support are some of the key components in comprehensive HIV programs. Such supportive services are available in both of the intervention cities. Study test sites will maintain the ability to refer clients with urgent needs (e.g., mental health) for necessary services. However, test sites will not routinely link patients to support services. The DOHs in both intervention communities actively encourage a single referral of HIV-positive patients from test sites to care sites. Multiple referrals can be overwhelming to newly HIV-diagnosed patients and may give the wrong impression that an individual must be drug-free in order to engage in HIV care, and may
paradoxically delay entry into care. Additionally, testing sites usually do not have ongoing relationships with persons in need of support services, and are thus not well suited for evaluating sustained linkages to these resources or adequacy of services to meet patient needs.

Participating HIV care sites will serve as the comprehensive ‘medical home’ for coordination of all HIV-positive patient care needs, including substance use, mental health, and other support services. As newly diagnosed HIV-positive patients present for care, care sites will evaluate CD4 cell count and VL. Care sites will then create an appropriate medical plan for each patient, consistent with current HIV care and treatment guidelines. Prior to study implementation, all HIV care site staff will be trained on the importance of linking HIV-positive patients to appropriate support services.

1.1.3.3 HIV Care and ART Utilization in Washington, D.C.

Historically, Washington, D.C., like several other jurisdictions, has had limited success in ensuring that clients newly diagnosed with HIV are routinely and rapidly linked to a primary HIV “medical home.” In 2006, before the HIV testing scale up, only approximately 50% of persons newly diagnosed with HIV were linked to care, as evidenced by a baseline CD4 cell count within six months of diagnosis reported to HIV surveillance. In contrast, in 2007, approximately 67% of the newly diagnosed HIV cases had laboratory evidence (CD4 cell count or VL) of a first HIV-related medical care visit within six months of the initial diagnosis.

In 2008, Washington, D.C. began to look beyond initial linkage-to-care to continuum-of-care after diagnosis. The number of HIV/AIDS cases reported between 2004 and 2007 increased 20.3% from 1,239 cases to 1,490 cases; data from 2008 are still preliminary but include 1,198 reports to date. Sixty-seven percent (67%) of cases had a first CD4 cell count, percentage or VL reported within three months of HIV diagnosis in 2008. The median CD4 cell count among newly diagnosed cases increased 57% from 216 cells/µl in 2004 to 340 cells/µl in 2008. Among newly diagnosed AIDS cases, the proportion of late testers decreased from 66% in 2004 to 57% in 2008.

Washington, D.C. supports full access to HIV care and treatment for all HIV-positive residents through a combination of regular Medicaid/Medicare, the Medicaid 1115 waiver, and Ryan White-funded services, including the AIDS Drug Assistance Program (ADAP). However, despite availability of services, use and outcomes of HIV care in Washington, D.C. remain sub-optimal. Preliminary review of ADAP data suggests irregular utilization at client level, with approximately 60% of enrollees picking up medications in a given month. Moreover, a preliminary review of Ryan White-funded, primary HIV-care providers suggests that only between 25-60% of clients in care are on ART, with approximately 50% of those being virally suppressed (unpublished data from ADAP and grantee reports).

Washington, D.C. is starting to address sub-optimal utilization and outcomes through a number of initiatives. First, a mass media information campaign promoting the availability of treatment was instrumental in increasing ADAP enrollment over 50% within 18 months. Since 2007, the HIV/AIDS, Hepatitis, STD and TB administration (HAHSTA) has also conducted treatment promotion through the “It’s Free to Treat Your
HIV” campaign, which is currently being retooled with patient components (emphasizing treatment enrollment and retention) and a provider toolkit (improving patient linkages and outcomes). Second, Washington, D.C. is collaborating with CDC to identify ways to better utilize routine ADAP pharmacy data to identify irregular utilization and build rapid feedback loops to providers. Third, the Ryan White Planning Council and HAHSTA have adopted a “health first” approach, making navigation (for linkage-to-care from positive test), re-capture (bringing persons lost to follow-up back into HIV care), and retention core priorities for all new funding opportunities for 2009-2012.

HIV care in Washington, D.C. is supported by a range of funding sources, to ensure that every individual in need of HIV treatment is provided both primary care and the supportive services necessary to remain in care. HAHSTA directly supports primary medical care programs at nine organizations in Washington, D.C. These primary medical care programs serve as a hub for services, with the responsibility of coordinating the services needed by their clients. In 2008 HAHSTA began implementing a best practice, four Rs (recruitment, recapture, retaining and results) approach to improving the continuum and continuity of HIV care in Washington, D.C. Recruitment or rapid entry to care links individuals who receive a positive HIV test immediately into primary care facilities, and includes special programs, such as: 1) Red Carpet Entry to Care services; 2) adult and adolescent healthcare navigation services; and 3) Rapid Entry to Care initiatives. To recapture patients into care, HAHSTA partnered with several funded care providers over three months to bring back into care nearly 900 people known to be living with HIV but found to not have had a CD4 cell count or VL test in the previous six months. For example, Family Medical Counseling Services re-established care with approximately 70% of 450 clients identified as lost to care, often after more than 10 contacts. Finally, retaining people living with HIV into care and ensuring quality results is a critical component of ensuring the continuity and continuum of care. HAHSTA has supported best-practice, outcome-based management of HIV-positive clients in care with affiliated primary care providers. HAHSTA has partnered with funded providers to use E-clinical Works and other electronic medical record (EMR) systems to institute evidence-based best practices such as deploying coordinated care teams, using focused, comprehensive health messaging, and identifying outcomes at every clinical encounter, with the goals of reducing missed medical appointments, increasing efficiency, increasing prophylaxis and increasing viral suppression.

Another specific strategy to increase linkage-to-care in Washington, D.C. is the requirement that all individuals with preliminary positive rapid HIV tests be linked immediately into care. When confirmatory testing is needed, it is performed at the initial medical visit, along with CD4 cell count, VL and other routine tests for HIV-positive individuals.

Most recently, HAHSTA funded Navigator Services, consisting of an Adult Navigator and an Adolescent Navigator, to facilitate the linkage of HIV-positive individuals into care and treatment. This new citywide resource supports the efforts of independent clinicians who have historically diagnosed up to 40% of our new HIV-positive persons. Navigator services provide individualized support to link newly diagnosed and previously identified positives not in care to a medical home for ongoing specialized HIV care and treatment, as well as evaluate these clients for any other social service needs.
1.1.3.4 HIV Care and ART Utilization in the Bronx, New York

Recent data evaluating linkage-to-care in NYC generally, and the Bronx specifically, have shown slightly more robust linkage rates than in Washington, D.C. However, retention in care is comparable. Most recent surveillance data for NYC show that approximately 70% of newly diagnosed persons have laboratory evidence of a first HIV-related medical care visit (demonstrating linkage-to-care) within three months of their initial diagnosis of HIV. The linkage rate in the Bronx is equivalent to that citywide. Continuity of care, defined by evidence of a medical visit (CD4 cell count or VL reported in the HIV/AIDS Registry) at least every six months, was 54.4% for NYC overall and 49.3% in the Bronx. Among persons initiating care within three months of HIV (non-AIDS) diagnosis, median CD4 cell count was 439 cells/mm$^3$ citywide and 451 cells/mm$^3$ in the Bronx. Finally, 69% of persons eligible under Department of Health and Human Services (DHHS) guidelines to receive ART citywide (CD4<350), and 70% of eligible persons in the Bronx, had achieved an undetectable VL within a median of six months after initiation of care.

HIV-related treatment and economic services are widely available for all HIV-positive New Yorkers, regardless of means. A combination of Medicaid/Medicare services, Ryan White-funded services, including the ADAP, Housing Opportunities for People with AIDS (HOPWA), and benefits available through the HIV/AIDS Services Administration of NYC's Human Resources Administration ensure broad coverage to meet treatment and social service needs. Despite this network of support, outcomes of HIV care for New Yorkers including Bronx residents, as described above, indicate some potential for gain.

NYC has undertaken at least two major initiatives to attempt to make headway in these areas. Beginning in December 2009, the NYC DOH will begin funding 27 agencies throughout NYC, including six agencies in the Bronx, for comprehensive coordination of care using Ryan White funds. Coordination of care will include health system navigation, medical case management with treatment adherence, and educational coaching with goals of viral suppression and self-sufficiency for individual patients. Additionally, an expanding Field Services Unit (FSU), created in 2006, stations public health advisors on site at ten tertiary hospitals in the highest prevalence neighborhoods of NYC, as well as at Rikers Island jail, a large correctional facility. FSU staff assists index patients and physicians with partner elicitation, partner notification and testing of partners. FSU staff also work to recapture patients who have fallen out of medical care.

1.1.4 Adherence to ART and Viral Suppression in the United States, Washington, D.C. and the Bronx

Non-adherence to ART is common among patients in the United States, with the percentage of prescribed doses taken estimated between 60% to 70% (Simoni, Pearson et al. 2006). Somewhat higher adherence levels are reported over shorter recall periods (Mugavero, Ostermann et al. 2006; Lazo, Gange et al. 2007). However, levels of adherence to ART as high as 80% to 95%, depending on the type of regimen, may be necessary to achieve and maintain maximal viral suppression and optimize clinical outcomes (Bangsberg 2008). Incomplete adherence has been among the most important factors related to virologic failure, the emergence of drug resistance, and ultimately
progression of HIV disease to AIDS and death (Lazo, Gange et al. 2007; Lima, Geller et al. 2007; Horberg, Silverberg et al. 2008). Non-adherence also contributes to the transmission of drug-resistant HIV strains (Sethi, Celentano et al. 2003).

Recently introduced ART regimens are less toxic, more tolerable and simpler, particularly if available in fixed-dose combinations (ARV treatment guidelines). These include ART regimens with new agents from established classes and new classes of ART that have been recently shown to be effective in suppression of resistant viral strains (Steigbigel, Cooper et al. 2008; Markowitz, Nguyen et al. 2009). Multiple clinical trials have shown the ability to achieve suppressed viral replication with the use of such regimens in a large proportion of patients. However, suboptimal adherence remains common, and young age, active drug and alcohol use, and depression are associated with poor adherence to ART (Levine, Hinkin et al. 2005; Lazo, Gange et al. 2007; Horberg, Silverberg et al. 2008). In clinical practice, adherence is typically assessed by self-report (Simoni, Kurth et al. 2006) and can be corroborated by the use of MEMS caps, pill counts, biologic markers (e.g., plasma drug levels), pharmacy refill data, (Bangsberg 2008) and ultimately by the measurement of HIV VL.

In addition to optimizing ART regimens, effective approaches to improving ART adherence include ART-readiness training, adherence-related case management, various forms of counseling, pharmacist-based support, telephone support, reminder devices, and directly observed therapy (Bartlett 2002; Bangsberg 2008). A recent meta-analysis of randomized behavioral interventions conducted mostly in the United States found that participants receiving adherence interventions were 1.5 times as likely to report 95% adherence and 1.25 times as likely to achieve undetectable VL than controls (Simoni, Pearson et al. 2006). Improving ART adherence requires a combination of methods appropriate to the patient and clinical setting. Among the most important predictors of ART adherence are engagement in medical care and continuous adherence with medical visits (Aberg, Kaplan et al. 2009). Alterable factors known to impact adherence, such as mental health problems, active substance use, homelessness, inadequate level of social/economic support (Magnus, Kuo et al. 2009), as well as the therapeutic relationship between patient and provider should be addressed proactively (Bakken, Holzemer et al. 2000; Bangsberg 2008).

Both in the Bronx, NY and Washington, D.C., the health departments consider adherence support services to be an essential component of a comprehensive HIV-care package. These health departments monitor trends in adherence by reviewing HIV VLs and CD4 cell count data reported to HIV surveillance systems. For example, HAHSTA approaches the promotion of treatment adherence as a responsibility shared by all providers or services including physicians, physician assistants, nurses, pharmacists and other clinical care providers. Upon enrollment, and periodically thereafter, each client’s records are reviewed for evidence of viral suppression as well as assessed for the need for specialized services to improve the ability of the client to take HIV medications effectively and remain consistently in care. A wide range of services is available for those in need, including linkages to mental health and substance use programs, education on HIV, programs designed to assist with the necessities of daily living, and ongoing individual or group psychosocial and peer support.
1.1.5 Prevention for Positives

Meta-analyses show that prevention efforts in HIV-positive individuals (Prevention for Positives) (Crepaz, Lyles et al. 2006; Johnson, Carey et al. 2006), can be effective in decreasing risky behaviors. However, most Prevention for Positives interventions have been assessed only in research contexts and require substantial investments of staff and resources to deliver with fidelity, suggesting that population-level uptake may not likely follow. Sexual risk assessments and risk-reduction counseling are not routinely performed in many HIV clinical settings, and even when they are, quality and consistency are variable (Metsch, Pereyra et al. 2004). Information and communication technologies (ICT) may facilitate scale-up of prevention interventions, as they utilize the client’s time for self-monitoring, deliver content with fidelity, can include multiple languages, and, once programmed, can be used for multiple computers with marginal cost beyond cost of the hardware. A recent meta-analysis found that these tools are efficacious for reducing HIV transmission risk (Noar, Black et al. 2009). A recent study by Gerbert et al. found that computer-delivered “video doctor” counseling to 960 HIV-positive patients was associated with reduced transmission risk behavior (Gilbert, Ciccarone et al. 2008). CARE+ Prevention is one example of an intervention that may be effective in decreasing sexual behaviors most likely to transmit HIV. Other studies have shown that Prevention for Positives interventions are less effective with patients with ongoing substance use and mental health conditions.

In Washington, D.C. HAHSTA is in the process of developing a Prevention for Positives initiative that includes consistent and repeated delivery of prevention interventions by clinicians to people living with HIV/AIDS (PLWHA) in care and treatment settings. The strengths of this clinic-based approach include linking the prevention of HIV transmission to the treatment of HIV, offering repeated opportunities to intervene in high-risk behavior, and serving as a source of information, support and linkage to needed services for the HIV-positive individual. The five areas of focus for this initiative will be: 1) social support for disclosure; 2) treatment adherence for viral suppression; 3) mental health; 4) substance use; and 5) general prevention interventions, such as distribution of condoms.

The NYC DOHMH has begun a concerted effort to prioritize prevention activities with HIV-positive individuals. Beginning in 2007, the department began funding 19 agencies to conduct 16 different behavioral interventions that have clear evidence of achieving behavior change. While many of these interventions work with HIV-positive and high-risk HIV-negative individuals, six agencies were funded to conduct interventions that focus primarily on HIV-positive individuals, such as "Healthy Relationships." Partner services activities as described above, operating primarily through the FSU and the Contact Notification Assistance Program, also focus directly on HIV-positive individuals as well as soliciting, notifying and testing partners of index patients. As part of their routine work, FSU staff members also assist in linkage and health navigation activities, discuss risk reduction, and provide free condoms, along with condom education as needed.

In 2009, the NYC DOHMH introduced a CDC-developed training course for NYC healthcare providers focusing exclusively on HIV prevention for positive patients called
“Ask, Screen, Intervene.” The course was offered quarterly and will continue in 2010. The Department further developed a comprehensive prevention-for-positives protocol for clinicians and non-clinician CBO staff, which will be made available in 2010. Finally, both the Ryan White comprehensive coordination of care program and a planned new funding stream will increase dedicated prevention activities for HIV-positive persons beginning in 2010.

1.1.6 Provider and Patient Attitudes Towards ART Use

Mathematical models that have addressed the issue of use of ART for prevention of HIV transmission have largely concluded that ART would be most effective for HIV prevention if used in the largest proportion of HIV-positive populations (Granich, Gilks et al. 2009). This implies that such treatment would be provided to HIV-positive individuals irrespective of their eligibility for ART initiation, as detailed in the available national or international guidelines. To date, such guidelines (Health and Human Services (HHS) guidelines, WHO ART guidelines) have been largely based on evidence supporting benefits versus risks of the use of ART for the individual with HIV infection, rather than based on prevention considerations.

Providers have largely utilized such recommendations to guide their decisions for when to initiate or use ART in HIV-positive individuals, aiming to provide patients in their care with the opportunity to achieve optimal outcomes for their individual health and well-being. Use of ART in HIV-positive individuals to benefit their partners and the community is a concept that goes beyond this framework and places providers in a dilemma, should the well-being of their patients be affected adversely by therapy whose aim is to reduce potential transmission to others in the community.

Similarly, from the patient’s perspective, ART use has been largely perceived to provide individual benefits. Thus, in making decisions regarding initiation of ART, patients make their assessment based on information provided to them regarding benefits and risks to their own health rather than based on potential effect on transmission of HIV to others. Adding reduction in infectiousness as a consideration will need to be weighed and patient knowledge and attitudes regarding such use carefully evaluated.

1.1.7 Financial Incentives

The use of financial incentives (FIs) to modify behaviors has become increasingly common in settings both within and outside the healthcare sector. Outside the healthcare setting, experiments have ranged from conditional cash transfers to alleviate poverty among low socio-economic status (SES) individuals in NYC to efforts to improve school performance within NYC, Chicago and Washington, D.C. Within the healthcare setting, in recent years, there has been increasingly widespread use of FIs among insurers and employers who see this as an approach with great promise to help individuals make better tradeoffs between unhealthy behaviors (that have immediate gratification) in favor of healthier behaviors (that have delayed benefits). For example, NYC is evaluating the effectiveness of FIs in achieving normal HbA1c levels in individuals with diabetes. In fact, such efforts have now been shown to be effective in changing health behavior and improving health outcomes in a variety of clinical contexts.
The interest in interventions aimed at improving patient behaviors is in part due to recognition that unhealthy behaviors may be a bigger contributor to poor health and premature mortality than inadequate healthcare delivery. Experts have estimated that unhealthy behaviors, such as medication non-adherence, smoking, poor diet and sedentary lifestyles account for as much as 40% of premature mortality in the United States, whereas deficiencies in healthcare delivery account for only 10% of premature mortality (Schroeder 2007). Many factors such as the social and structural environment, public policies, genetics and provider access and quality affect the rate of such behaviors. However, individuals’ behavioral choices are clearly a central driver and are potentially more amenable to incentives.

Early evidence on incentives suggests that incentive-based approaches can be highly effective in two areas in particular: (1) changing short-term health behaviors related to preventive services that involve a limited number of visits, and (2) reducing the use of addictive substances. Examples of effectiveness of incentives in increasing use of preventive services include studies that have shown increases in rates of follow-up for abnormal pap smears (Marcus, Kaplan et al. 1998), postpartum visits by adolescents (Stevens-Simon, O'Connor et al. 1994), TB test reading (Malotte, Rhodes et al. 1998), and the rate at which IV drug users received all three doses of hepatitis B vaccine (Seal, Kral et al. 2003). The evidence that such approaches are effective in reducing the rate of use of addictive substances, such as cocaine (Higgins and Silverman 1999; Lussier, Heil et al. 2006) and nicotine in the short term, (Donatelle, Hudson et al. 2004; Volpp, Gurmankin Levy et al. 2006), as well as for short-term weight loss (Jeffery, Thompson et al. 1978; Jeffery, Gerber et al. 1983; Finkelstein, Linnan et al. 2007), suggests that financial rewards designed to incent long-term changes in behavior could be applicable to a wide range of other health behaviors in which frequent reinforcement and longitudinal follow-up are necessary.

However, evidence of effectiveness of incentives is not limited to the areas cited above. A review of 11 randomized trials of FIs found that in 10 studies, FIs promoted adherence better than any tested alternative, leading to better blood pressure control, better appointment attendance and higher immunization rates (Giuffrida and Torgerson 1997). More recent reviews of economic incentives found that a wide range of incentive mechanisms are effective in changing behavior (Kane, Johnson et al. 2004; Sutherland, Christianson et al. 2008).

Another reason to consider further testing of incentive-based approaches is that many highly efficacious medical tests, treatments and medications have limited effectiveness due to patient behaviors. For example, by one year after having a myocardial infarction, nearly half of patients prescribed cholesterol medications have stopped taking them (Jackevicius, Mamdani et al. 2002). Similarly, the effectiveness of HIV medications would be much higher if rates of adherence increased to the point where benefits demonstrated in clinical trials could also be seen in high rates of effectiveness in communities across the United States.

More recent research has found that FIs double the rate of attendance at and completion of a smoking-cessation program, triple long-term smoking cessation rates and substantially increase medication adherence among patients on warfarin. For example, in
a two-arm RCT of 878 employees at General Electric (GE), (Volpp, Troxel et al. 2009) incentives of up to $750 resulted in quit rates at nine-12 months triple those in the control group (p<.001). Other studies have examined the use of incentives for weight loss (Volpp, John et al. 2008) and medication adherence (Volpp, Loewenstein et al. 2008).

Two studies have been published in which FIs were used for ART adherence. In both cases, the payments made a significant improvement in ART adherence (Rigsby, Rosen et al. 2000; Rosen, Dieckhaus et al. 2007). The findings that FIs can help modify difficult-to-change behaviors whose cessation requires ongoing reinforcement, such as quitting alcohol or cocaine use, are particularly striking and suggest that a well-designed incentive program can succeed in changing behaviors in clinical contexts in which many other approaches have been unsuccessful (Bigelow and Silverman 1999; Donatelle, Prows et al. 2000; Higgins, Wong et al. 2000).

The amounts we selected for use as FI in this study are based on the following principles: 1) review of relevant literature and experiences, 2) input from investigators with knowledge of target communities and populations, 3) input from community groups/advisors from the study communities and elsewhere, and 4) input from key staff in departments of health and public health entities in the relevant communities. Based on these factors, the amount of $125 for successful linkage-to-care, and a maximum of $630 for viral suppression throughout the study period were thought to be appropriate amounts for evaluation in this study.

With regard to the sustainability of any effect caused by the FI after it is removed, a recent study on incentives for smoking cessation (Volpp, Troxel et al. 2009) demonstrated that the ratio of tobacco cessation among incentive to control group participants at nine-12 months (2.9) remained significant six months after cessation of incentive payments (ratio of quit rates was 2.6 at 15-18 months). In other work, shorter durations of incentives have been associated with higher relapse rates. It is of interest that based on the findings of the latter study on smoking cessation, GE announced plans to implement a program for FIs for smoking cessation nationally for all its 152,000 employees.

Studies involving FIs have focused on both process measures (the behavior in question, such as medication adherence) and outcomes (smoking cessation). There are several precedents for the use of outcomes (e.g., VL) to assess the effect of FIs on medication adherence. Several studies that used FIs to improve adherence with medication (e.g., warfarin or insulin) have measured biologic outcomes to evaluate adherence (e.g., INR, HbA1c) (Volpp, Loewenstein et al. 2008). It is essential that FIs be provided in such a way that they yield a verifiable outcome. Because lack of VL suppression most often results from lack of adherence, the use of FIs to encourage ART adherence is plausible from both a medical and a behavioral-economics perspective. The relative effectiveness of focusing incentives on process versus outcomes is generally unknown and is ultimately an empirical question. In Washington, D.C., a demonstration project indicated that a focus on outcomes (rather than process) has been associated with improvements in the areas of HIV prevention and care (Hadar, personal communications). Specifically, rewarding HIV-care organizations for every three patients they successfully re-linked to care, regardless of the effort expended, yielded positive results. In this study, FIs will
focus on outcomes – the completion of care visits for the Linkage-to-Care component and in the Viral Suppression component.

In contingency management studies, non-monetary rewards have often been used in place of monetary rewards because of concerns that cash might increase the likelihood of substance use. However, studies have not supported such concern. In the AIDS Link to Intravenous Experience (ALIVE) study, ethnographers trailed the drug users after they received the money to assess where they went. They found that participants mostly went to food establishments after they received the money. There was no difference in the rates of death caused by overdose in participants when analyzed by days after study visit or amount of reimbursement (Vlahov, Tang et al. 2000).

1.1.8 HIV Surveillance Data

Since 1982, all 50 U.S. states and Washington, D.C. have reported AIDS cases to the CDC using a standardized case report form (CRF) (Schneider, Whitmore et al. 2008). In 1994, the CDC integrated national reporting of HIV with AIDS case reporting, at which time 25 states with confidential, name-based HIV reporting started submitting case reports to the CDC. Over time, additional states implemented name-based HIV reporting and started reporting these cases to CDC. By April 2008, all states and Washington, D.C. had implemented name-based HIV reporting and were reporting cases to the CDC.

HIV case surveillance data are collected as part of routine HIV surveillance, as mandated by state or local laws or regulations. Named reporting is required for all diagnoses of HIV and AIDS and all HIV-related illness according to the case definition. Information is collected on demographic characteristics of persons diagnosed with HIV, transmission risk factors, facility of diagnosis, diagnostic tests (e.g., positive Western Blot (WB) tests for HIV antibody) and death. All areas require laboratory reporting of VL and CD4 cell count values. While not all areas require reporting of all values of VL or CD4 cell count (some limit reporting to detectable VL and/or CD4 <200 or <14%), all areas collect the first results of such tests after HIV diagnosis through chart review or laboratory reporting, where all values are reported through voluntary laboratory reporting. Both intervention communities require reporting of all laboratory results. Some areas such as NY also require reporting of all HIV genotypes.

These population-based registries are continuously updated with new, de-duplicated diagnoses and laboratory results. Incoming diagnostic WB and VL reports from providers and laboratories that cannot be matched to an existing registry record initiate a field investigation to confirm the case, date and disposition of diagnosis and collect other data required for surveillance. Data are also obtained through matches with other disease registries, the state and local death registries, the National Death Index and the Social Security Death Master File. Assessments of duplicate cases occur both on the state and national level (potential duplicates are identified based on soundex code [a phonetic algorithm for indexing names by sound, as pronounced in English] and selected demographic characteristics), while elimination of such cases occurs at the state level.

HIV reporting and laboratory reporting requirements allow virtually complete surveillance of diagnoses of HIV, stage of disease at HIV diagnosis, and number of...
people diagnosed and presumed to be living with HIV. Using CD4 cell count and VL test ordering as proxy measures for initiating HIV primary care, after the first positive WB test, allows for calculation of the time between diagnosis and initiation of care. Frequency of visits, regularity of U.S. DHHS-recommended laboratory monitoring, and estimates of the proportion of cases eligible for ART are now possible with CD4 cell count and VL result reports. These laboratory indicators also allow estimates of the number and characteristics of cases not in care. Clusters of highly resistant HIV will be detectable when state genotype reporting systems become operational.

The CDC has developed a data entry and reporting system, the Program Evaluation and Monitoring System (PEMS), to strengthen monitoring and evaluation of HIV-prevention programs. PEMS is used by health departments and CBOs funded through CDC HIV-prevention cooperative agreements. PEMS allows grantees to collect agency data, community planning data, program plan data, and client-level data. This assures a comprehensive set of standardized variables are available.

Some areas have implemented additional surveillance activities, such as behavioral risk factor surveys and supplemental surveillance activities supported by the CDC (e.g., incidence surveillance). The National HIV Behavioral Surveillance (NHBS), which takes place in 25 cities, can provide information on testing among various population groups including among MSM, a special emphasis group in this project. Where available, the Morbidity Monitoring Project (MMP, a 26-site study) may provide information on treatment among those in care. In addition, data are collected on testing conducted in all sites receiving funding from the CDC and in additional sites as required by state or local laws or regulations.

1.2 Purpose

The main purpose of this study is to assess the feasibility of a community-level test, link to care, plus treat strategy in the United States. The study will involve evaluation of the feasibility of some components and the effectiveness of others.

The study will assess the following:

- The feasibility of expanding HIV testing via social mobilization and universal offer of HIV testing in EDs and inpatient hospital admissions
- The feasibility and effectiveness of facilitating the linkage of HIV-positive patients to HIV care sites
- The feasibility and effectiveness of different strategies for assuring maximum initiation of ART for clinically eligible patients according to current guidelines
- The feasibility and effectiveness of different strategies for promoting high adherence to ART and maintenance of VL suppression
- The effectiveness of a computer-assisted program for Prevention for Positives
- Patient and provider attitudes towards the initiation of ART in early HIV disease
The primary outcomes of this study’s package of interventions will be determined through measurement of change from baseline (at the initiation of the study) over the duration of the study (see study timeline) in key parameters in two intervention communities in the United States. Observations in four non-intervention communities will help assess the influence of current trends in HIV testing and care expansion in the United States.

This study will serve as a proof-of-concept, formative study. It will provide key information to guide the design and anticipate the costs of a future large, randomized, community-level clinical trial of full implementation of a test-and-treat strategy in the United States. This study uses innovative approaches including: 1) a community focus; 2) multi-component strategies that include behavioral and biomedical interventions; 3) the use of routinely reported HIV surveillance data to determine key outcomes; and 4) partnership with both local DOHs and the CDC. Findings from this study could also inform test-and-treat efforts in other developed countries with epidemics similar to that in the United States.

HPTN 065 will help generate estimates for some of the parameters that, taken together, will describe an “index of participation.” Such an index of participation (upon which the test-and-treat strategy would ultimately depend) is based on a cascade: the percentage of HIV-infected persons tested and identified within a community, the percentage of such individuals linked-to-care, and the percentage who initiate and remain adherent to ART and maintain ongoing viral suppression. For example, even with optimistic assumptions (80% of infected persons tested; 80% linked to care, and 80% adherence to ART) the index of participation would be 51%. These parameters have, to date, been treated as assumptions in test-and-treat models. The mathematical model described by Granich et al. assumed that all adults would be tested annually with a 100% sensitive, 100% specific test; that all HIV-infected persons would enter care and start ART as soon as they were diagnosed irrespective of HIV disease stage; that there would be a decrease of 99% in their infectiousness and that only 1.5% of subjects would discontinue ART each year, for an unrealistic index of participation of 98.5% (Granich, Gilks et al. 2009).

HPTN 065 will examine the feasibility and relative effectiveness of various interventions that individually aim at optimizing each component related to the index of participation. These insights together with data on the magnitude of the effect of ART on infectiousness (to be obtained from other studies), could be incorporated into future models that aim to determine the potential effect of a test and treat strategy on incidence in various communities in the United States.

1.3 Rationale

1.3.1 Rationale for a Test-and-Treat Approach

More than one million persons in the United States are currently living with HIV and about 21% are unaware of their HIV-positive status. The HIV epidemic in the United States affects specific subpopulations and localized “hot spots.” Individuals may be at risk for HIV because of their own risky behaviors or the high prevalence of HIV among
persons whom they encounter, or a combination of both. Risk may be compounded by high rates of undiagnosed HIV infection within these same communities.

In 2006, there were an estimated 56,000 new HIV infections in the United States; a number that has been approximately stable for the past decade (Hall, Song et al. 2008). Certain municipalities and communities in the United States bear the brunt of the HIV epidemic. This includes certain geographic areas in the United States and specific subsets of individuals defined by behavior and/or by race/ethnicity. The data from 2006 indicate that MSM and AAs are most severely affected. For example, in Washington, D.C., three percent of the population is HIV-positive. National Health and Nutrition Examination Survey (NHANES) data indicate that HIV prevalence among AAs age 40-49 is nearly 3% among women and 4.5% among men (McQuillan, Kruszon-Moran et al. 2006).

The central factors that drive the HIV epidemic in the United States are: (1) the number of individuals unaware of their HIV infection who continue behaviors likely to transmit HIV; (2) the frequency of late diagnosis of HIV infection; and (3) delay in access to care, delay in initiation of ART and suboptimal adherence, leading to failure of sustained viral suppression. Approximately 40% of persons are diagnosed with AIDS within a year of their first HIV-positive test (Valdiserri, Holtgrave et al. 1999; Castilla, Sobrino et al. 2002; (CDC) 2009), too late to realize the full benefit of advances in HIV management. Persons with undiagnosed HIV infection may also unwittingly transmit HIV to partners. While they represent less than one quarter of the entire HIV-infected population, persons with undiagnosed HIV infection account for more than 50-70% of new sexually transmitted infections in the United States and are 2.5 times more likely to transmit HIV than persons who are aware they are infected (Marks, Crepaz et al. 2006). Evidence also indicates that many diagnosed individuals delay or fail to engage in HIV care (Torian, Wiewel et al. 2008), and for those who have initiated ART, retention in care and adherence to ART is suboptimal (Lazo, Gange et al. 2007; Mugavero, Lin et al. 2009).

Since the beginning of this decade, evidence has accumulated that higher plasma HIV RNA is associated with increased risk of HIV transmission (Quinn, Wawer et al. 2000; Fideli, Allen et al. 2001). More recent data suggest that use of ART is associated with a substantial decrease in rate of HIV transmission from the HIV-infected to uninfected sexual partner (Bunnell, Ekwaru et al. 2006; Sullivan, Kayitenkore et al. 2009). The availability of various ART regimens capable of suppressing viral replication in various subsets of HIV-positive individuals offers the opportunity to determine the potential effectiveness of use of ART on HIV incidence in a community. While various models suggest that such a strategy may be associated with a successful impact on the trajectory of the HIV epidemic, empiric data are needed to confirm this hypothesis.

It is widely anticipated that if a TNT approach is to achieve its desired impact, it would require the following ambitious effort: universal HIV testing efforts; prompt and effective linkage of all individuals with HIV infection to HIV care; timely initiation of ART; and sustained suppression of HIV replication in all HIV-positive individuals. It is evident that assessment of such an intervention requires a deeper understanding of each of its components.
Thus, we will examine five study components highly relevant to the TNT approach. Four study components are interventions and one is a survey. The four study interventions to be assessed include: expanded HIV testing, linkage from HIV testing sites to HIV care sites, viral suppression with use of ART, and a computer-delivered intervention to achieve safer behaviors among HIV-positive individuals. In addition, surveys for both HIV-positive persons and HIV-care providers will be done to determine knowledge and attitudes regarding early use of ART and financial incentives.

1.3.2 Rationale for Choice of Intervention Communities

This study, Test, Link-to-Care, Plus Treat, will assess five components in two communities in the United States, the Bronx, NY and Washington, D.C. These communities were selected due to: (1) the severe impact of the epidemic on these communities (NYC has the largest number of cases of HIV of all cities in the United States and Washington, D.C. has the highest HIV seroprevalence), and (2) the substantial efforts these cities have already made to improve HIV diagnosis, facilitate linkage-to-care, and support adherence.

NYC has the oldest, largest and most heterogeneous epidemic in the Western World. By June 30, 2008, a cumulative total of 207,687 persons in NYC had been diagnosed and reported with HIV. 43,344 (20.8%) of these persons were residents of the Bronx at the time of diagnosis. A cumulative total of 100,378 persons with HIV have died over the course of the epidemic (48.3% of cumulative diagnoses). There have been 20,052 deaths (19.9% of cumulative deaths citywide) among Bronx residents. In 2007 HIV was the third leading cause of death in persons under age 65.

As of June 30, 2008, there were 104,234 persons diagnosed, reported and living with HIV in NYC. 22,479 (21.6%) were residents of the Bronx. Citywide, the rate of new HIV diagnoses was 46.8/100,000 population. The diagnosis rate in the Bronx was 62.4/100,000 population. The Bronx rate was second to that of Manhattan, which reported a diagnosis rate of 70.5/100,000 population. Citywide, 26.1% of newly diagnosed persons have already progressed to AIDS at the time they first learn they are infected with HIV (concurrent diagnosis of HIV and AIDS). The borough with the highest proportion of concurrent diagnoses (27.3%) is the Bronx.

Washington, D.C. is in the midst of a generalized HIV epidemic. At the end of December, 2007, there were 15,120 persons diagnosed and reported living with HIV in Washington, D.C. Between 2003 and 2007, there were 7,432 new HIV/AIDS cases reported, bringing to more than 3% the proportion of Washington, D.C.’s adult population diagnosed and living with HIV. It is estimated that one-third to one-half of infected persons may be unaware of their HIV infection (DC NHBS data). All wards in Washington, D.C. (with the exception of Ward 3) have prevalence rates above the Joint United Nations Programme on HIV/AIDS (UNAIDS)/WHO threshold for a generalized HIV epidemic. More than two-thirds of new HIV diagnoses are in people over age 30 years. More than 7% of Washington, D.C. residents age 40-49 and 5.2% of D.C. residents age 50-59 years are diagnosed and living with HIV. The rate of HIV is highest among black males (6.5%), but 3.0% of Hispanic males and 2.5% of white males and black females are estimated to be living with HIV. In examining the most recent trend
data on HIV (not AIDS) and newly reported AIDS cases, mode of transmission trends are changing. Heterosexuals account for over 37% of newly reported HIV cases and 31.5% of newly reported AIDS cases, followed by MSM (25.8% and 29.0%, respectively) and injection drug users (13.2% and 21.7% respectively). With the advent of ART, deaths due to AIDS have significantly decreased over the last 15 years. HIV/AIDS, however, is still the leading cause of death in Washington, D.C. residents age 25-44 (Washington 2009). In 2007, there were 138 deaths from AIDS or AIDS-related complications.

1.3.3 Rationale for Choice of Interventions

The study will focus on expansion of HIV testing in hospital emergency departments and inpatient units. The rationale for expanding the offering of HIV screening in EDs emanates from numerous studies in urban EDs that indicate a 0.7% - 1% yield of new HIV diagnoses among those tested. Many patients with HIV infection visit EDs, but remain undiagnosed. For example, of all patients with new HIV diagnoses in South Carolina from 2001-2005, 73% had previously visited healthcare facilities, but were not tested for HIV; 79% of these visits were to an ED. Although many EDs in both the intervention and non-intervention cities have initiated HIV screening, most ED programs are unable to offer screening 24/7, and many are unable to test all patients who agree to be tested.

A similar yield of new HIV diagnoses might be expected from testing hospital admissions. Data from the NY Health and Hospitals Corporation (the organization that operates 11 public hospitals in NYC where expanded HIV screening began in 2006), and from Boston Medical Center’s HIV Inpatient Testing Service, indicated an HIV positivity rate of 2.5% - 5% among inpatients who agreed to be tested (selectively thus which may or may not reflect the prevalence of HIV infection among all inpatients, including those who refused HIV testing or were not offered testing). No data exist on the yield of routine HIV screening of all hospital admissions. This approach was traditionally a significant component of syphilis screening efforts. Similarly, this approach may be a cost-effective way to expand HIV screening and to ensure that no HIV-positive patient who encounters the healthcare system remains undiagnosed. This component of the HPTN 065 study (i.e. supporting expanded ED- and hospital-based offering of testing) will be a supplement to ongoing testing efforts in these communities.

The study will also aim to use social mobilization to increase HIV testing volume and frequency. Because of the bimodal nature of the HIV epidemic in the Bronx and in Washington, D.C. (generalized, especially among AAs, and concentrated among MSM) the study will also include another strategy in the testing component. Through social mobilization, focused messages will be created for the intervention communities on the importance of getting tested for HIV and knowing one’s status (tested in past year). In addition, specific messages will be created to promote more frequent retesting for sexually active MSM—preferably twice a year, but at a minimum, annually.

In this study, we will not aim to identify acute infections. Unpublished data from the CDC’s Acute HIV Infection Study demonstrate that HIV retesting as described above is considerably more cost-effective than Nucleic Acid Amplification Testing (NAAT) screening for acute infection.
The study will utilize FIs to promote linkage of HIV-positive individuals from testing to care sites and to achieve and maintain viral suppression. As described in Section 1.1.7 FIs hold great promise for achieving the target behaviors. The study will use a site-randomized approach to compare the effectiveness of FIs to the standard of care (SOC) in achieving these two goals.

Some TNT models assume that all HIV-positive individuals will initiate ART irrespective of their CD4 cell count. Evidence from observational studies supports earlier initiation of ART, with the goal of delaying HIV disease progression and reducing mortality rates (Kitahata, Gange et al. 2009; Sterne, May et al. 2009; When to Start Consortium 2009). The availability of safer, simpler, more tolerable and potent ART regimens has reduced, but not eliminated, some concerns about the development of long-term toxicities and complications among HIV-positive patients (Justice 2006; Friis-Moller, Reiss et al. 2007) and reduced concerns about development of antiretroviral resistance (Phillips, Leen et al. 2007; von Wyl, Yerly et al. 2009). However, in this study, the provision of ART will be guided by the prevailing DHHS Guidelines for the Use of ART in HIV-Positive Adults and Adolescents (Panel on Antiretroviral Guidelines for Adults and Adolescents 2009). Providers at participating sites will receive educational trainings on best practices in HIV management, per current guidelines, and will be encouraged to promptly and appropriately evaluate their patients for eligibility for ART, maximize ART initiation when indicated per current guidelines, and switch regimens when indicated.

Lastly, while TNT models have not directly addressed the contribution of efforts to promote safer behaviors to reduce transmission from HIV-positive persons, unsafe sex behaviors by HIV-positive persons are a major factor contributing to ongoing HIV transmission. Therefore, in this study, we will evaluate an individually randomized computer-delivered prevention-for-positives component to reduce unsafe sexual and injection-drug using behaviors. In this manner, we will gather feasibility and effectiveness data to inform about the value of such an intervention for future studies.

1.3.4 Rationale for Use of Surveillance and Routinely-collected Data

The study will utilize routinely reported HIV surveillance data collected in the United States to determine the effects of the various interventions. The United States is fortunate to have a robust HIV surveillance system that was established at the advent of the epidemic in the early 1980’s. Over the ensuing years, this system has been expanded and refined. At present, in many communities, it includes data on all positive HIV tests, demographic and risk behavior data on all those found to be HIV-positive, collection of all CD4 cell count and HIV RNA results, and information on AIDS-related deaths, among other data elements. Thus, this system offers a remarkable opportunity to assess various site-level or community-level interventions.

The evaluation of each of the present study’s components, and the feasibility of use of existing HIV surveillance data, will be critical to the design, implementation and evaluation of a large, future, community-focused test-and-treat study, with HIV incidence as the endpoint of interest. The collaborative process utilized in the development of this study that includes partnership among the National Institutes of Health (NIH)-funded
HPTN, the CDC, and departments of health is noteworthy. In addition, the engagement of diverse settings, and organizations, including HIV testing sites, HIV care sites, health facilities, providers, CBOs, advocates and leaders is critical to its success and necessary for future large studies of TNT strategies.

1.4 Protocol Structure

This protocol describes the five study components: 1) expansion of HIV testing, 2) enhancing the linkage of HIV-positive individuals to HIV care sites, 3) maintenance of VL suppression, 4) use of a computer-delivered intervention to decrease high-risk sex behaviors (Prevention for Positives), and 5) provider and patient surveys to gather data on knowledge, attitudes and practices regarding ART and ART adherence (potentially starting at higher CD4 cells counts) and FIs. Sections 2-6 are devoted to the objectives; study design, interventions, and procedures; data and statistical analysis; and ethical issues of each study component. Sections 7-8 contain information applicable to all study components (HIV surveillance data, data sources and administrative/operational issues).

2.0 EXPANDED HIV TESTING

2.1 Study Objectives for Expanded HIV Testing

The intervention communities have committed substantial programmatic funds to expanded testing. The study will aim for expansion of HIV testing efforts in the intervention communities building on ongoing efforts in these communities. The study will focus on two elements: 1) social mobilization and 2) universal offer of HIV testing and counseling in hospital EDs and inpatient units.

True evaluation of the efficacy of increased testing and its impact on detecting undiagnosed infections could only be achieved through a high-quality random sample, representative of the general population, to obtain information about HIV status and knowledge. The expense and scale of such an effort is beyond the scope of a feasibility study. Thus, we have adopted the planned ecological approach of monitoring process and outcome measures to capture increase in HIV testing and change in number and characteristics of newly detected infections throughout the course of the study.

The primary feasibility objectives for expanded HIV testing, through a combination of focused and enhanced social mobilization efforts, will be measured by assessing the following trends in the intervention cities:

- Volume of testing in publicly funded testing sites
- Number of newly identified HIV-positive individuals
- Site of diagnosis for newly identified individuals
- Initial CD4 cell count results after first positive HIV test (to assess proportion eligible for ART at different initiation thresholds)
- Proportion of newly identified individuals with HIV concurrently diagnosed with AIDS
- Proportion of population tested for HIV in the last year (local behavioral surveys)
Because expanded testing initiatives are already underway in numerous jurisdictions, the magnitude and trends of changes in the intervention cities will be compared to similar measures in the non-intervention cities, in an attempt to account for secular trends independent of the study components.

- The following variables in the intervention cities will be assessed:
  - Costs of enhanced testing activities in EDs and for hospital admissions
  - Costs of social mobilization
  - Intensity of social mobilization activities (content, number, timing of activities, and type of activity, such as TV, radio, brochures, posters, community leaders, events, etc.)

The protocol team will track all of the expenses listed above. Funds for these items will be held centrally and only disbursed when line item bills are presented. Funds provided by HPTN 065 will be a small supplement to DOH program efforts in Washington, D.C. and the Bronx, NY. The protocol team will request costs of existing activities from the DOHs.

The **primary feasibility objectives** for expanded HIV testing in EDs and for hospital admissions will be measured by determining the following:

- Proportion of visits to EDs and admissions to hospitals who receive an HIV test (including all individuals permitted by local or state law to consent for HIV testing in the study communities)
- Number of patients tested in EDs newly identified as HIV-positive and their demographic characteristics
- Number of patients tested during hospital admissions newly identified as HIV-positive and their demographic characteristics
- Number of tested patients identified with previously diagnosed HIV who are not in care
- Cost of support for additional staff and HIV tests

### 2.2 Design for Expanded HIV Testing

This study component will aim to substantially expand HIV testing activities in the two intervention communities. The Expanded HIV Testing component is a descriptive ecologic study. For this component of the study, outcomes will be evaluated through routinely reported surveillance and process data. Because extensive interventions to expand testing have already been undertaken, the study team will catalogue the existing activities in the intervention communities as they relate to social mobilization, enhanced testing and linkage activities, in collaboration with departments of health in the two communities.

### 2.3 Study Population for Expanded HIV Testing

The target age population for the social mobilization element of expanded HIV testing will be all persons age 13 and older in the Bronx, NY and Washington, D.C., with special emphasis on MSM.
The target population for the implementation of universal offer of testing in EDs and during hospital admissions will be individuals permitted to consent for HIV testing according to New York State or Washington, D.C. law.

2.3.1 Inclusion Criteria

The inclusion criteria for the universal offer of HIV testing are the following:

- Individuals who are permitted to consent for HIV testing according to New York State or Washington, D.C. law
- Capacity to understand and provide consent for HIV testing
- Admission to a Bronx, NY or Washington, D.C. ED and/or a Bronx, NY or Washington, D.C. hospital

2.3.2 Exclusion Criteria

The exclusion criteria for the universal offer of HIV testing are:

- Lacks the capacity to provide consent for HIV testing
- Acute or urgent medical condition that might be adversely affected by process of obtaining consent or performing HIV test

2.4 Study Sites for Expanded HIV Testing

2.4.1 Hospital/ED Study Sites in the Bronx, NY

Efforts to expand HIV testing will be undertaken at approximately seven Bronx hospitals with at least a Level I trauma center/ED and with daily hospital admissions. Overall, total ED visits at these hospitals in 2006 ranged from 49,635 to 134,969 per hospital.

2.4.2 Hospital/ED Study Sites in Washington, D.C.

Efforts to expand HIV testing will be undertaken at approximately seven Washington, D.C. hospitals. Routine HIV testing through the ED at George Washington University Hospital and throughout Howard University Hospital, begun in 2006, accounted for nearly 25% of all publicly supported tests in Washington, D.C. and 20% of reported positives in the HIV testing data in 2008.

2.5 Interventions for Enhanced HIV Testing Activities

The goal of the enhanced HIV testing activities to be implemented in the intervention communities is to increase the proportion of individuals from the community who have been tested within the prior year, with the intent of achieving earlier detection of HIV infection. Two populations are being targeted: those with ongoing risk through their own or partner risk behaviors and those with prevalent HIV-infection who are unaware of
their HIV infection. The former are targeted for regular HIV testing, at least annually, the latter are targeted for one-time “capture” testing.

At least three factors hamper the existing HIV testing promotion programs: limitations on the ability to influence hospital infrastructure; the inability to collect enhanced data from unfunded agencies; and limited resources to sustain social marketing campaigns. Within each institution, multiple stakeholders (including physicians, administrators and Department of Health staff) have identified necessary logistical changes that would be required to bring HIV screening to scale. Because staff have many other duties that often preclude their ability to focus on these key logistical or infrastructure change, the necessary coordination and follow-through between departments has not taken place. The ability to hire dedicated staff solely on overcoming key logistical barriers to routine HIV screening within each institution should significantly address this first limitation.

Programs such as “The Bronx Knows” have been able to collect aggregate data on HIV screening from all of its testing partners; however, because there has been no direct funding to each organization to support this data collection, the complexity and quantity of process data available to evaluate the initiatives are limited. The study will fund enhanced data collection directly at each site and allow creation of a more robust data set to evaluate various aspects of the testing promotion.

Although funds provided by the CDC in 2008 and 2009 have allowed for the development and dissemination of initial phases of consumer campaigns encouraging HIV testing, resource limitations hampered the sustainability of social messaging campaigns. Supplemental funding from the study will improve both sustainability of ongoing messaging and the creative development of additional components, including a module geared to medical providers. In both jurisdictions, the messaging to the general adolescent/adult population has been to get tested – either annually (Washington, D.C. message) or at least once (Bronx message). However, in both jurisdictions, some sub-populations, such as men who have sex with men, would be best served by more frequent repeated testing. Delivery of targeted enhanced testing messages alongside general testing messages will be supported by the TLC-Plus study.

In addition, the earliest models of routine HIV testing in medical settings were often stand-alone projects working in parallel with the existing medical services, necessary in order to prove feasibility in an environment where routine testing was felt to be burdensome or risky. These models have often relied extensively on the availability of rapid point-of-care HIV tests. However, as routine testing expands new models of integrated testing and lab-based large platform testing may be more appropriate to sustainably meet volume, workload, and cost-efficiency needs. The “early adopter paradigm” applies here – once a program has initial success in routine testing, support is needed for continued evolution or shift of that model. The TLC-Plus study will supplement CDC funding for expanded testing to support this evolution.

The study will build on current social mobilization efforts and communications plans in the intervention communities by adding two refined messages:
• All sexually active individuals age 13 and older should have had an HIV test within the prior year

• Sexually active MSM should seek HIV testing, at least annually, but ideally every six months

This study component will include social marketing to the general population in the two intervention communities, with outreach to the MSM community, healthcare providers, CBOs and social networking systems.

The study aims to achieve the goal of universal offer of HIV testing in EDs and during hospital admissions through the combination of the following approaches, tailored to the needs of each facility:

• Outreach to facility directors and key leaders at these institutions

• Establishment of HIV-testing goals for each facility

• Establishment of HIV testing as part of routine admission orders

• Provision of financial support for HIV-testing kits needed for increased testing

• Development and implementation of computer-delivered information on HIV testing

• Provision of computer terminals

2.6 Study Procedures for Expanded HIV Testing

2.6.1 Study Procedures for Social Mobilization

Both the Bronx, NY and Washington, D.C. have ongoing social mobilization activities as part of their HIV campaigns (“The Bronx Knows” and “DC Takes on HIV”). The HPTN 065 study will provide additional resources to craft and fine tune key messages aimed at increased testing and testing frequency as well as increase testing frequency in specific populations. In addition, the study will also support the development of messages that highlight importance of linkage to HIV care and HIV treatment. Focus groups will assist in developing these messages and in evaluating tools to determine their acceptability and effectiveness. The details of the social mobilization interventions will be determined at the beginning of the study; however, they may include, but are not limited to: flyers/brochures in clubs frequented by MSM; radio commercials; Internet ads/links to study information on sites commonly visited by MSM; and utilizing CBO staff to disseminate “word-of-mouth” messaging in the community.
2.6.1.1 Study Procedures for Social Mobilization in the Bronx

The study will supplement current efforts by the NYC DOHMH for the creative development and placement of social marketing materials throughout the Bronx, NY. Materials produced will both reinforce and supplement current social marketing messaging regarding the importance of routine HIV screening in healthcare settings and will further refine the messaging to the two target populations as indicated above.

The NYC DOHMH will develop social marketing materials using its standard procedures, including (but not limited to) contracting with local advertising agencies to create and produce materials, per NYC contracting practices.

All messaging will be subject to approval by the NYC materials review board, as is the practice for other CDC-funded social marketing materials developed by the NYC DOHMH. Sample materials produced also will be provided to the NYC DOHMH Institutional Review Board (IRB) for review.

Decisions regarding dissemination of social marketing materials will be made by the Bureau of HIV/AIDS Prevention and Control in the NYC DOHMH.

In general, placement of social marketing materials developed for MSM will focus on locations frequently visited by MSM including the Bronx Community Pride Center and local MSM-friendly Bronx clubs, such as Mi Gente. Where possible and appropriate, social marketing materials will be placed on the Internet (e.g. on social networking sites frequented by MSM in the Bronx).

2.6.1.2 Study Procedures for Social Mobilization in Washington, D.C.

Financial support will be provided to supplement the comprehensive social marketing program the Washington, D.C. DOH has already developed. This multi-phased program seeks to scale up routine HIV testing, promote treatment and promote behaviors to reduce risk of infection. The program is branded with the umbrella message “DC Takes On HIV” and promotes routine HIV testing in medical settings. Other aspects of the social marketing campaign include a consumer component entitled “Ask for the Test” and a provider component, “We Offer the Test.” Using principles successfully implemented by the pharmaceutical industry, the campaign aims to drive consumers to ask for HIV testing when they visit their doctor. The message for providers is that patients will expect HIV testing as part of their standard healthcare. The consumer program features both traditional media (public transit, newspaper, radio and television advertising) and new media (Internet and text messaging). The DOH has developed an umbrella Web site www.DCTakesOnHIV.com with links to resources about HIV testing and services. The DOH has also established a text messaging service where residents can text “DCTEST” to 365247, receive a health fact, and then search by zip code for the nearest free HIV-testing location.

The provider component, “We Offer The Test,” includes a toolkit (handbook, pocket card, poster, information cards for patients and appointment cards for HIV specialists) to make it easier for practitioners to implement routine HIV testing. The provider materials
include practical steps to make HIV testing routine, including sample scripts for use with patients, information on billing codes, information cards outlining what patients should know about both negative and positive results, and a “refusal” card that informs patients of the health risk from not getting an HIV test. DOH has formed a partnership with the Global Business Coalition and Pfizer to implement a pilot program under which Pfizer sales representatives will promote routine testing and provide the toolkit during their regular visits to medical practitioners. The pilot will start with 200 physicians and practices in Washington, D.C.

Washington, D.C. will also use social networking media (social Web sites such as Facebook and MySpace) to increase access to Washington, D.C. residents, and will use Twitter as a vehicle to communicate relevant information on HIV. Currently, the DOH has established these sites and a Twitter account for its free condom distribution program. DOH has entered into a contract with a public relations firm to advance the social marketing materials.

### 2.6.2 Study Procedures for Universal Offer of HIV Testing in EDs and Hospital Admissions in the Bronx

#### 2.6.2.1 Emergency Departments

New York State has requirements for specific information that must be provided to patients before they sign separate, informed consent for HIV testing. HIV testing is already being offered in some EDs in the Bronx. Under this protocol, testing will be expanded at the EDs where it is currently offered and introduced in those that are not testing. Given the complexity and unique exigencies of each hospital's own logistics, this study will work within each facility to establish a procedure for universal offer of HIV testing. A universal offer development team (UODT), hired at the beginning of the study period for each intervention community, will conduct a series of meetings with staff at multiple levels of seniority within each hospital who have some stake in ED care. The UODT will work with these staff to identify and overcome current barriers to universal offer of HIV testing in their ED. Key staff may include the ED Director, the chief of nursing, laboratory director, chief ED residents and chief administrators. The UODT will also work with the ED leadership to set goals for testing and will provide quarterly results to each hospital to create and implement strategies for improvement.

Where necessary to improve logistics, laptop computers attached to rotating stands that can be easily moved from bay to bay will be provided by the study team to the ED site. Videos that meet New York State requirements for pre-test counseling will be loaded onto the laptops. Patients can view the videos and indicate whether they agree to test. Where possible, the video will be set up so that when a patient indicates interactively that he or she agrees to an HIV test, a tester (a nurse, physician or dedicated tester) will be notified and the test will be performed. While the patient waits for results, post-test, HIV-prevention messaging will also be viewable, if the patient chooses to watch it.

All EDs will be provided with free HIV test kits for individuals who are uninsured and cannot bill an insurance carrier (including Medicaid) for HIV screening. Free test kits
will be provided in proportion to the percentage of uninsured individuals seen in that ED who receive an offer of HIV testing at baseline.

The study will also provide EDs with a downloadable computer widget application on routine HIV testing that can be placed on the desktops of all ED computer terminals (either an updated version of the tool created by St. Vincent's Medical Center, with funding from the New York State AIDS Institute, or a similar tool). This widget will not only provide the full CDC Guidelines for routine HIV screening, but will include brief checklists that walk providers through the process of obtaining written consent (currently still required in NY), conducting an HIV test, providing results and handling a positive HIV diagnosis.

In working with the complex logistics in EDs, the study team will also operate within existing hospital admission processes as much as possible.

2.6.2.2 Hospital Admissions

Although some NYC hospitals have initiated programs to offer HIV testing for inpatients, none has yet implemented universal offer of testing at the time of hospital admission. At the beginning of the study period, the UODT will conduct meetings at hospitals with staff at multiple levels of seniority who have some stake in the inpatient admissions process and inpatient medical care. The UODT will work with these staff to identify and overcome current barriers to a universal offer of HIV testing in the hospital admissions process. Key staff will include the medical director, the chief of nursing, the laboratory director, internal medicine and surgery chief residents and chief administrators. The UODT will also work with the medical director and/or other designated hospital leadership to set goals for testing and will provide quarterly results to each hospital to create and implement strategies for improvement.

For any hospital that has set standing (basic, preset) admitting orders, the UODT will work with the Medical Director (or her/his designate) and the hospital's information technology department to have HIV testing added to these standing admitting orders, in accordance with hospital policies. The UODT will also work with the hospital's laboratory to overcome any barriers to placing HIV testing on hospital standing orders. All HIV testing completed as part of this protocol, whether in the ED or during hospital admission, will be conducted according to the laws of New York State. The UODT will work to ensure that all necessary paperwork is streamlined and available on all floors for the admitting teams and for admitting nurses.

All hospital admitting teams will ascertain if the patient has already been tested in the ED during his/her current visit, so that duplicate HIV testing is not performed. If possible, prior tests offered in the ED will be entered electronically, so that the admitting team will know whether the patient had been offered the test.

Admitting teams will be instructed to review the key pre-test counseling points that are required by New York State, obtain consent and, if possible, order the test with routine admitting phlebotomy. If ordering the test with routine phlebotomy is not possible, an HIV rapid test will be performed.
Alternatively (if desired) hospital admitting teams can be provided with laptop computers attached to rotating stands that can be easily moved from patient to patient. Videos that meet New York State requirements for pre-test counseling will be loaded onto the laptops. Patients can view the videos and indicate whether they agree to testing. Where possible, the video will be set up so that when patients indicate interactively that they agree to an HIV test, a tester (a nurse, physician or dedicated tester) will be notified and the test will be performed.

As with EDs, free HIV tests will be provided to the facilities for uninsured persons offered testing on admission. The percentage of tests offered will be derived in similar fashion to the percentage offered for ED testing. Where possible, the same downloadable widget provided to EDs will be provided to inpatient admitting teams for use on their computer terminals.

2.6.3 Study Procedures for Universal Offer of HIV Testing in EDs and Hospital Admissions in Washington, D.C.

2.6.3.1 Emergency Departments

Washington, D.C. EDs have been focal points for the implementation of routine HIV screening because many Washington, D.C. residents only access healthcare through the city’s EDs. HIV testing in EDs can be integrated as a routine, opt-out procedure because Washington, D.C. does not require a separate written consent form for HIV testing. Patients are provided with written or verbal information about HIV testing, advised that a test is recommended, and (depending on available staff) a point-of-care rapid test is performed, unless the patient declines. In addition to the two institutions currently conducting routine testing in EDs, the Washington, D.C. DOH has begun to engage six additional EDs to undertake routine, opt-out HIV testing. HAHSTA recently completed review of a Request for Applications (RFA) to support expansion of routine HIV screening in EDs and hospital centers by providing support for a part-time coordinator at each hospital to promote implementation of routine testing. A coordinator at each facility will work with hospital leadership and department heads (CEOs and medical, lab and data staff) to develop plans for routine HIV screening in the ED and in other departments in the hospital system. In Washington, D.C., health insurers are required to reimburse for a voluntary HIV test performed during an insured’s visit, regardless of the reason for that ED visit. Study funds will be used to support the cost of tests for patients who are uninsured. In addition, the study will support a part-time coordinator at each institution, working with management and ED and laboratory staff toward implementation of routine screening. The Washington, D.C. DOH will provide TA, including staff training, to all partnering hospitals.

Washington, D.C. has developed a provider toolkit for routine, opt-out HIV testing that includes an introductory brochure, pocket card, result cards and opt-out card. These tools define routine, opt-out HIV testing and provide checklists for the screening process, from test introduction to the provision of follow-up appointments for HIV-positive individuals. These materials will be available for wide distribution to the staff of partnering EDs participating in routine, opt-out HIV testing and to clinical providers throughout Washington, D.C.
The Washington, D.C. DOH will also provide test kits to partner EDs for use with uninsured patients. Although Washington, D.C. legislation mandates reimbursement for HIV testing performed in EDs, none of our partner hospitals has yet developed a protocol for billing for HIV tests. Free tests kits will be made available to hospitals while they develop systems to achieve sustainability through billing and reimbursement procedures. As those systems mature, the hospitals will assume responsibility for procuring test kits independently.

The study will provide financial support to participating EDs in order to increase staffing for HIV testing activities.

2.6.3.2 Hospital Admissions

Washington, D.C. hospitals are already at varying stages in the process of offering HIV testing to inpatients, but none conduct routine screening at admission. Seven of Washington, D.C.’s eight hospitals are performing rapid HIV testing in labor and delivery units for women without a documented HIV test result. Howard University Hospital currently makes rapid HIV testing available during regular business hours, upon physician or patient request. United Medical Center’s long-range goal is to implement routine, opt-out HIV testing throughout the hospital.

Intensive TA will be provided to hospitals for admission testing, as with ED HIV testing. In an effort to promote the most cost-effective method of HIV testing, the DOH and study-supported coordinator at each institution will work with hospitals to add routine, opt-out HIV testing to standing orders for admission blood work for all hospital admissions. Washington, D.C.’s lack of a requirement for separate signed consent for HIV testing makes this a feasible option. Hospitals will also be encouraged to offer point-of-care tests for persons who are not tested at the time of admission.

Hospital admissions staff will have access to the routine, opt-out HIV testing reference materials originally developed for EDs. The Washington, D.C. DOH will also provide HIV test kits to hospitals for inpatient testing that must be performed at point-of-care (when not done as part of admitting blood work) until they achieve sustainability through billing and reimbursements.

The study will provide financial support to participating hospitals in order to increase staffing for HIV testing activities.

2.7 Study Duration for Expanded HIV Testing

The expanded HIV testing component of this protocol will continue for the duration of the feasibility study, currently projected to be 36 months. It is expected—by virtue of emphasizing stakeholder buy-in, providing training and logistical support, and improving staff and systems capacity—that several key activities undertaken as part of this protocol, such as establishing HIV testing as part of routine admission orders, will be sustained after the study is completed.
2.8 Statistics and Data Analysis for Expanded HIV Testing

2.8.1 Endpoints

2.8.1.1 Endpoints for feasibility of enhancing HIV testing through a combination of focused and enhanced social mobilization efforts

- Number and results of HIV tests per month in publicly funded testing sites (local health department data)
- Number, transmission category and testing source of newly identified cases in HIV surveillance data
- Initial CD4 cell count of newly identified HIV cases in surveillance data
- Number of newly identified HIV cases concomitantly diagnosed with AIDS in surveillance data
- Proportion of persons in the community tested for HIV in the last year (local population-based behavioral surveys)

2.8.1.2 Endpoints for feasibility of routine offer of HIV testing at emergency departments and inpatient units

- Proportion and number of total ED visits and admissions to hospital where patients receive HIV testing
- Number of HIV tests in EDs where HIV infection is newly identified
- Number of HIV tests in hospital admissions where patients receive HIV testing
- Proportion of hospital admissions who have newly identified HIV infection
- Number of tested patients identified with previously diagnosed HIV who are not in care
- Cost of support for additional staff and HIV tests

2.8.2 HIV Testing Intervention Goals

More than 72,000 publicly supported HIV tests were conducted in Washington, D.C. in 2008, of which approximately 1000 were new positive diagnoses. Likewise, between April 2008 and March 2009, 161,619 HIV tests were performed as part of “The Bronx Knows” initiative, of which approximately 700 were new, positive diagnoses. This study will evaluate the increase in overall testing in Washington, D.C. and the Bronx. In addition to the volume of testing and number of new HIV diagnoses, the study will evaluate trends in CD4 cell count at first diagnosis.

Enhanced access to HIV testing in both ED and inpatient settings will occur throughout the entire 36-month study period. With regard to specific targets for the testing effort, the study will aim to offer HIV testing to 80% of eligible individuals during 80% of visits
with a 60% acceptance rate in the EDs. Likewise, the study will aim to offer HIV testing to 80% of eligible individuals with a 70% acceptance for inpatient admissions. The absolute numbers that these percentages represent will vary from hospital to hospital. The study will evaluate the ED and inpatient HIV data for increases in volume of testers, increases in number of HIV-positive tests and increased volume and percentage of new HIV infections detected.

2.8.3 HIV Testing Baseline Data

Table 1 contains the current levels of testing volume and newly identified cases from Health Departments and Surveillance data in each of the six cities (intervention and non-intervention) for 2007 and 2008.

Table 1. Testing Volume and Results in the Six Intervention and Non-Intervention Communities

<table>
<thead>
<tr>
<th>Cities</th>
<th>2007</th>
<th>2008</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. of HIT Tests N</td>
<td>No. of Sites N</td>
</tr>
<tr>
<td>Bronx(^a)</td>
<td>127,947</td>
<td>21</td>
</tr>
<tr>
<td>Washington, D.C.</td>
<td>43,271</td>
<td>47</td>
</tr>
<tr>
<td>Chicago</td>
<td>Not Available</td>
<td>77,616</td>
</tr>
<tr>
<td>Houston</td>
<td>38,612</td>
<td>16</td>
</tr>
<tr>
<td>Philadelphia</td>
<td>44,504</td>
<td>315</td>
</tr>
<tr>
<td>Miami</td>
<td>48,696</td>
<td>100</td>
</tr>
</tbody>
</table>

a) Data from The Bronx Knows Initiative
b) Data on linkage-to-care and newly diagnosed are only available for 19 agencies participating in The Bronx Knows for 2007.
c) The Bronx Knows agencies began reporting data in April of 2008; data reported here therefore covers the period of April 2008-March 2009 to provide one year’s worth of data.
d) Data on newly identified HIV+ in Houston included only a subset of testing facilities. Numerators and denominators for these data reflect only that subset of tests/facilities.

2.8.4 Data Analysis

Data to be analyzed for this study component will be obtained from routine HIV surveillance sources and from utilization data collected by participating hospitals.

Endpoint measures of the potential feasibility of large-scale implementation of an expanded HIV testing strategy will be obtained by two analytic strategies: 1) comparison within the intervention communities before vs. after the intervention, and 2) comparison of change between communities with vs. without the intervention. For the within-community comparison, we will monitor the change from pre- to post-
intervention, with the primary interest in the change achieved at the end of the intervention period. For the between-community comparison, we will compare the change in communities with vs. without the intervention. Thus, we will test two indicators of change as a result of the intervention:

- **Within**: Comparison of outcomes of interest before vs. after intervention in an intervention community

- **Difference of differences**: Comparison of the change in a measure during the study period in communities with vs. without the intervention cities

Measurement of change or temporal trends in any of the outcome measures of the intervention implies that both of the above indicators of change will be computed.

### 2.9 Human Subjects/Ethical Considerations

The expanded HIV testing study component is a public health practice. Two geographic areas will participate in this study component: Washington, D.C. and the Bronx, NYC. Social mobilization and emergency department testing is already taking place in the two intervention communities (Bronx and Washington, D.C.) and is intended to be specific to the needs of those populations. While some observations from the overall TLC Plus project with regard to social mobilization and expanded testing may be applicable elsewhere in the United States, these activities were initiated originally for the specific benefit of the respective communities, and do not constitute research (CDC 1999).

The protocol will be submitted to appropriate IRBs (a central and/or local site IRBs) for ethical review prior to study initiation. Any subsequent modifications to the protocol will be submitted to appropriate IRBs, and, at a minimum, the protocol will be submitted annually for continuing review and approval by these same ethics boards. Because the expanded HIV testing component is public health practice applied in the two study communities, the study team will request that IRBs reviewing the protocol as a whole consider only the expanded testing component to be a non-research component, and therefore not subject to the requirements of 45CFR46. The other three components of this protocol constitute research and will be addressed in separate sections.

In Washington, D.C., written informed consent is not required for routine HIV testing; only oral consent, as part of standard voluntary opt-out HIV testing, is required. In contrast, New York State requires written informed consent for individuals undergoing HIV testing. For the expanded HIV testing component of this study, participants will undergo HIV testing according to the SOC and legal requirements of their jurisdiction.

To assess the impact of the study on HIV testing in the communities of Washington, D.C. and the Bronx, surveillance data routinely collected by the DOHs will be analyzed. No individual data, other than what exists in the surveillance databases, will be collected from study participants in the HIV-testing component of this study.

No study-specific laboratory testing will be conducted under this protocol. Therefore, no additional study-related test results will be reported to authorities. HIV-testing data for
tests performed in the EDs, hospitals and community test sites in Washington, D.C. and the Bronx, NY will be reported per local HIV reporting requirements.

The study team will neither collect nor report Adverse Events because there is no biomedical intervention. Again, due to the nature of this study component, the team will not collect or report Social Harms.

3.0 LINKAGE-TO-CARE

3.1 Study Objectives for Linkage-to-Care

The primary feasibility objective for using FIs to facilitate linkage of HIV-positive patients to HIV care sites will be measured by determining the following:

- Overall cost of the program, including staffing, program materials and incentives
- Extent of other available linkage-to-care activities (case managers, peer navigation, etc.)

The primary effectiveness objective for using FIs to facilitate linkage of patients from HIV testing to HIV care sites is the following:

- To increase the proportion of newly HIV-diagnosed and out-of-care patients linking to care within three months of identification

The protocol team will track all of the expenses listed above. Funds for these items will be held centrally and only disbursed when line-item bills are presented.

In order to monitor the implementation of the FI program, certain parameters will be gathered from a subset of participating sites at various time points to ensure program quality. These parameters will include: the proportion and number of patients eligible for incentives (the number of HIV–positives at a participating site), the proportion and number of patients receiving incentives, and the amount received compared with the total number who are eligible.

3.2 Design for Linkage-to-Care

The Linkage-to-Care component of the study is a two-arm, site-randomized, prospective, effectiveness clinical trial conducted within each intervention community. This study component compares the effectiveness of an FI intervention to link HIV-positive individuals from HIV test sites to HIV care sites with the usual SOC procedures. Each HIV test site will be randomized to either the intervention or SOC arm of the study. For this component of the study, study outcomes will be evaluated through routinely reported HIV surveillance data. Using surveillance data, study outcomes will continue to be evaluated for a period of one year after the FI intervention ends.

In order to identify the SOC for linkage, against which the incentives intervention will be assessed, an appropriate facility administrator will complete a brief survey that will collect key attributes of HIV testing sites, including types of navigation and support services already available to patients to facilitate linkage-to-care. Data will be collected from sites annually from 2009 - 2013.
For the purposes of this protocol, we consider an HIV-positive individual linked to care when that person has a VL or CD4 cell count assessment at an HIV care site within 3 months of confirmatory WB testing.

### 3.3 Study Population for Linkage-to-Care

The Linkage-to-Care component of the study will include all individuals ages 12 and older who are permitted to consent, or can be consented for HIV care by a parent/legal guardian according to New York State or Washington, D.C. law, and who are newly found to be HIV-positive at HIV test sites participating in the study. This study component will also include individuals who have been previously diagnosed with HIV but have been out of care for at least a year and are reconfirmed for HIV infection by standard laboratory tests.

### 3.4 Study Sites for Linkage-to-Care

Twenty HIV test sites will be selected from Washington, D.C. and 20 HIV test sites will be selected from the Bronx to participate in the Linkage-to-Care component of this study. HIV test sites will be selected based on two primary criteria: 1) site agreement to participate in this component of the study, and 2) sites with the highest volume of HIV-positive individuals identified in the previous year. Additional site selection criteria may be considered if a total of 40 HIV test sites cannot be chosen based on the primary criteria.

In each community, these 20 sites will be randomized such that 10 will use the FI intervention, described in Section 3.5, to link HIV-patients to care and 10 will use the SOC only.

The site randomization will be balanced by the following two baseline characteristics:

- The number of HIV-positive individuals identified in the previous year
- The rate of linkage-to-care within three months of HIV diagnosis over the course of the calendar year prior to study initiation

In addition, 20 care sites will be selected in Washington, D.C. and 20 care sites will be selected in the Bronx, NY. These care sites will redeem the coupons provided to patients by the test sites selected for the FI arm.

### 3.5 Intervention for Linkage-to-Care

At HIV test sites assigned to FIs, individuals permitted to consent for HIV care according to New York State or Washington, D.C. law who test HIV-positive will be provided with a coupon that is redeemable at participating HIV care sites. These coupons will be designed so they will not breach patient confidentiality (for example, there will be no patient names, no indication of HIV status, and no clinic names on these coupons). Patients presenting coupons to participating HIV care sites will be given an FI upon completion of a blood draw/lab visit ($25), which is usually the first visit, and another FI
($100) upon an interaction with their healthcare provider, which is usually at a second visit. Alternatively, for those patients who complete a comprehensive visit at participating care sites, a visit that includes both the lab and provider components for Linkage-to-Care, a $125 FI gift card will be provided.

The coupons distributed at the HIV test sites must be redeemed at the HIV care sites within three months of the date that a participant receives them.

The proportion of persons successfully linked to care within three months of their HIV-positive test will be compared, through the use of routinely collected HIV surveillance data, between sites implementing the incentives intervention and those with SOC procedures for linkage-to-care.

3.6 **Study Procedures for Linkage-to-Care**

The specific procedures for HIV test sites to obtain and distribute the FI coupons for linkage-to-care, as well as the procedures for the redemption of these coupons at HIV care sites, are outlined in the HPTN 065 Study-Specific Procedures Manual (SSP).

3.6.1 **Procedures at Test Sites**

Participating test sites will link all HIV-positive patients to an HIV care site. Each test site will be provided with a listing of participating HIV care sites (with contact information) to give to HIV-positive patients. In addition, test sites will give HIV-positive patients coupons redeemable for FIs at HIV care sites.

Prior to study implementation, all HIV test site staff will be trained on the procedures for the FI intervention. Providers will also be trained on HIV prevention counseling, and the importance of linking HIV-positive patients to care.

Study test sites will maintain the ability to refer clients with urgent needs (e.g., mental health) for necessary services. However, test sites will not routinely link patients to support services. The DOHs in both intervention communities actively encourage a single referral of HIV-positive patients from test sites to care sites. Multiple referrals can be overwhelming to newly HIV-diagnosed patients and may give the wrong impression that an individual must be drug-free in order to engage in HIV care, and may paradoxically delay entry into care. Additionally, testing sites usually do not have ongoing relationships with persons in need of support services, and are thus not well suited for evaluating sustained linkages to these resources or adequacy of services to meet patient needs.

3.6.2 **Procedures at Care Sites**

Participating HIV care sites will serve as the comprehensive ‘medical home’ for coordination of all HIV-positive patient care needs, including substance use, mental health, and other support services.

As newly diagnosed HIV-positive patients present for care, care sites will evaluate CD4 cell count and VL. Care sites will then create an appropriate medical plan for each patient, consistent with current HIV care and treatment guidelines. Simultaneously, care
sites will also assess patients and link them as appropriate to support services. Finally, valid coupons distributed by HIV test sites will be redeemed with FI gift cards.

Prior to study implementation, all HIV care site staff will be trained on the procedures for the FI intervention, ART initiation according to current guidelines, and the importance of linking HIV-positive patients to appropriate support services. In addition, as part of study orientation and training for HIV care providers, existing support service linkages will be reviewed.

To collect clinic-level information on services available to patients in the two jurisdictions, HIV care sites will be surveyed at baseline and annually thereafter to ascertain support services available on- and off-site (e.g., social services, substance use treatment, support groups, mental health resources, etc). The study team will develop and maintain a comprehensive list of support services in the two intervention communities. In addition, the study will regularly update lists for care sites of contact persons and referral processes at support service agencies.

Because the study will use surveillance data to measure outcomes and will not collect data on individual patients, the study will not be able to document or track linkages to ancillary services in the Linkage-to-care and Viral Suppression study components. Information on referrals made and completed by patients is not part of the surveillance data that will be analyzed for study outcomes. Likewise, offering FIs for completion of linkages to drug treatment or mental health services is not possible, as it would also require following individual patients with explicit data collection. However, for the subset of patients completing Patient Survey, we will collect data on the frequency of use, ease of access, and patient experiences with support services (including mental health, drug abuse, case management, etc).

3.6.3 Monitoring HIV Test Sites

It is possible that members of the intervention communities will learn of the FI to strengthen linkage-to-care and seek out HIV test sites offering these incentives. In all likelihood, the majority of individuals who seek testing will not know their HIV status yet and, thus, will be unaware that incentives are being offered for linkage-to-care. In addition, many of the HIV test sites in the study (~seven out of the 20 participating HIV test sites in each city) will be hospitals in which HIV testing will take place in emergency departments and inpatient units. Individuals seen in these settings are usually not there for the purpose of obtaining an HIV test. Thus, it is unlikely that these individuals will seek medical services based on the availability of FIs for linkage to HIV care.

However, it is possible that some individuals, who are aware of their HIV infection, may “recycle” through the HIV test sites due to the availability of FIs. There are some natural barriers to this phenomenon, as it is likely that the staff at these HIV test sites will recognize individuals who are testing with undue frequency. In these cases, staff may withhold the FI coupons from those who have already received them for a prior HIV-positive test. Another natural barrier is that individuals are known at HIV care sites, so individuals will be unable to re-link repeatedly to the same HIV care site. In addition, the FI intervention is designed such that multiple activities (lab-work and provider
interaction, both of which take time and effort) are required for the full redemption, which may discourage some from repeated linkage.

It should also be noted that based on the HIV prevalence in the intervention cities, relatively few people test positive out of all of those who test. Thus, the number of people who could “recycle” through the HIV test sites is relatively small compared to the total testing done in these venues.

Despite all of the barriers to HIV-test-site migration and linkage recycling, FI may encourage these phenomena; thus, the study team will use surveillance data to monitor these potential situations. Specifically, the team will monitor the number of duplicate linkage-to-care events in the name/ID-based surveillance data system over time. So, for example, in the case where a person re-tests at five unique HIV test sites (all offering linkage-to-care incentives) and links to five unique participating HIV care sites over the course of six months, HIV surveillance data will capture this event.

In addition, the team will monitor whether the number of positive HIV tests becomes severely disproportionately distributed among the participating HIV test sites (a sign that HIV testing is much reduced or increased at some sites). This information will be examined in an on-going fashion, and if site migration or linkage recycling becomes highly extensive so that it becomes a burden on HIV test or care sites, or threatens the validity of the evaluation, this component of the study will be re-evaluated. Such re-evaluation may include discussions with the community advisory group to design other mechanisms to minimize such migration, or, if needed, a redesign of this component of the study may be embarked on by the study team.

3.7 Study Duration for Linkage-to-Care

The duration of the FI intervention for the Linkage-to-Care component of the study is depicted in Figure 2. Coupons will be available at the HIV test sites randomized to the FI arm for 21 months. These coupons can be redeemed after the initial visit with an HIV provider at HIV care sites for a total of 24 months (for the entire time coupons are distributed at certain HIV test sites plus an additional three months).

Figure 2: Duration of FI Intervention for Linkage-to-Care
3.8 Statistics and Data Analysis for Linkage-to-Care

3.8.1 Endpoints

3.8.1.1 Endpoints for the feasibility of using financial incentives in the HIV-positive population for Linkage-to-Care

The feasibility endpoints for the Linkage-to-Care component of the study are the following:

- Number of individuals eligible for incentives (see section 3.3) and number of individuals receiving incentives (upon linkage to HIV care) at participating sites
- Cost of the program including staffing, infrastructure and incentives

3.8.1.2 Endpoints for the Effectiveness of Linkage-to-Care

In this site-randomized design, Linkage-to-Care effectiveness endpoints will be evaluated for each site and intervention effectiveness assessed by comparing site outcomes for intervention vs. SOC sites.

The effectiveness endpoints for the Linkage-to-Care component of the study will be ascertained based on HIV surveillance data and will include the following:

- Proportion of HIV-positive individuals at each testing site, who either have a newly detected HIV infection (based on new surveillance case with confirmed WB) or who were previously diagnosed but were out of care (based on no VL or CD4 ascertainment in the year prior to a repeat WB test in an existing surveillance case), and who presently are linked to care as evidenced by having a CD4 cell count and/or VL measurement at a separate visit within three months of WB confirmation
- Mean time interval at each testing site from HIV diagnosis (WB confirmation) to first CD4 cell count or VL for those with newly detected HIV infection and those who were previously diagnosed but were out of care
- Proportion of HIV-positive individuals at a testing site (overall, and separately for those with newly detected HIV infection and those previously diagnosed who were out of care) with at least two CD4 cell count and VL measurements in the prior year

3.8.2 Sample Size and Power

The primary endpoint of this study component is proportion of cases (newly diagnosed and previously diagnosed but not in care) with a CD4 cell count and/or VL measurement within three months of the diagnostic WB.

Randomization is by testing site, with balance achieved for baseline volume of testing and baseline linkage-to-care rates. Accrual of participants into the Linkage-to-Care
component of the study will occur over the first 24 months of the study, when eligible persons receive the FI coupon at an HIV test site randomized to the FI arm of the study. There will be no follow-up for this component of the study, other than the two visits in care (or a combined lab/provider intake visit).

In 2007, in the 20 testing sites with the highest volume in the Bronx, NY, 611 newly diagnosed HIV-infected people were identified. Of those, 77% were liked to care within three months, with linkage at different testing sites ranging from 30% to 100%. The mean number of newly diagnosed cases per provider in 2008 was 31, and ranged from five to 93. The intra-class correlation (ICC) of proportion linked to care in the Bronx was 0.27.

In 2008, in the 20 testing sites in Washington, D.C, the jurisdiction with the highest volume of people testing HIV-positive, 783 newly infected cases were identified. Of these, 67% were linked to care, ranging across different testing sites from 5% to 100%. The mean number of newly diagnosed cases per provider in 2008 was 40, and ranged from eight to 143. The ICC of proportion linked was 0.31.

These estimates do not include the population of those re-linking to care. It is anticipated that linkage success in SOC for this population will be lower because of the history of loss to care. We could reasonably expect that: 1) the number of people eligible for linkage-to-care will be higher because of longer study duration and addition of re-linked to care cases; 2) the proportion linked to care will be lower because of the re-linked to care cases; and, therefore, 3) the ICC will be smaller. It is assumed that the number of cases to be linked to care during the 24-month study duration will be similar to these past case loads on an annual basis. We will randomize 20 testing sites in each intervention city, and we expect to identify between 1200 and 2400 individuals for linkage-to-care, a mean of 60-120 people per testing site.

Table 2 below gives the mean number of linkage cases needed per testing site to detect an increase in linkage-to-care given 20 sites per arm (i.e. 40 in all). With a mean ranging from 80 to 100 linkages per site during the 24 months of the study, we would have 80% power, even with a high ICC, to detect a 13% change in proportion linked to care. If higher numbers are eligible for intervention due to linkage-to-care (e.g. 140), we would attain 90% power to detect this magnitude of difference. Note that the ICC coefficient estimated from the linkage-to-care of newly detected cases is high in large part because of sites with a small number of newly detected cases and 100% success in linking to care. It is probable that the ICC observed in the trial will be lower.

This sample size calculation for clustered designs uses the approach described by Thomson, Hayes and Cousens (Hayes and Bennett 1999; Thomson, Hayes et al. 2009).
Table 2. Mean number at each site required to achieve sufficient power to detect differences in proportion of newly detected or reconnected to care within three months of testing. Number of clusters fixed at 20 per arm

<table>
<thead>
<tr>
<th>Mean percentage linked to care at a site</th>
<th>Standard deviation of percent linked to care</th>
<th>Intra-class correlation</th>
<th>Mean number of potential cases for linkage-to-care per testing site</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard of care</td>
<td>Intervention</td>
<td>Standard of care</td>
<td>Intervention</td>
</tr>
<tr>
<td>67%</td>
<td>80%</td>
<td>17%</td>
<td>12%</td>
</tr>
<tr>
<td>67%</td>
<td>80%</td>
<td>20%</td>
<td>15%</td>
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<tr>
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<td>20%</td>
<td>17%</td>
</tr>
<tr>
<td>67%</td>
<td>75%</td>
<td>26%</td>
<td>22%</td>
</tr>
</tbody>
</table>
3.8.3 Randomization Scheme

In the Linkage-to-Care study component, HIV testing sites will be randomized to use FIs or the SOC.

HIV test sites will be assigned to one of the following two arms:

- An arm offering FIs: Test sites assigned to this arm will provide coupons to all individuals testing HIV-positive, who are not already linked to HIV care. The coupons can be redeemed for the FI at a participating HIV care site.

- An arm continuing with SOC: Each person who tests HIV-positive, and is not currently in care, will be directed to HIV care sites using the site’s SOC procedures.

The site randomization will be balanced by the following two baseline characteristics:

- The number of HIV-positive individuals identified in the previous year

- The rate of linkage-to-care within three months of HIV diagnosis over the course of the calendar year prior to study initiation

3.8.4 Data Analysis

Data to be analyzed for this study component will be obtained from routine HIV surveillance data as well as from the participating sites. Participating sites will maintain tracking logs, both for coupons provided to HIV-positive individuals at the test sites and for FI disbursement at the care sites. Information from the tracking logs, with all participant-identifying information removed, will be provided to the study team.

The analysis approach for testing the effectiveness of an FI intervention to link HIV-positive individuals from HIV test sites to HIV care sites compared to the SOC is described below. Analysis of secondary endpoints will be detailed in the statistical analysis plan.

3.8.4.1 Primary Effectiveness Analysis

The population for assessment of Linkage-to-Care is the following:

1. All individuals with newly detected HIV infections reported to the HIV surveillance system for the DOH jurisdiction of Washington, D.C. and the Bronx, NY who were tested at any of the 40 testing sites in the study

2. All participants eligible for re-linkage to care, defined as a WB recorded in the enhanced HIV/AIDS Reporting System (eHARs), at any of the 40 testing sites in the study after the implementation of FIs, who had no VL or CD4 assessment by a provider in the previous 12 months
The endpoint is the percentage of participants with a VL or CD4 cell count assessment within three months of the WB confirmation (separate from clinical assessment done at the time of the confirmation visit). Participants in the FI arm are attributed to the linkage-to-care strategy in place at the site where the first HIV-positive result was detected.

The estimation of intervention effect will test for a change in odds ratio of linkage-to-care between the SOC and FI test sites, assuming a correlation within participants tested at the same site, using Generalized Estimating Equations (GEE) methods.

### 3.8.4.2 Feasibility Endpoints

Analysis of feasibility endpoints will use descriptive statistics to characterize the variability of uptake of FIs across types of testing venues.

### 3.9 Human Subjects/Ethical Considerations

The Linkage-to-Care component of the study involves a randomized intervention using FIs to link HIV-positive individuals to HIV care after diagnosis. This component is public health research. In this component, HIV test sites will be randomized, not individuals, and no de novo individual-level data will be collected. Instead, (de-identified) HIV surveillance data routinely collected by the Departments of Health (DOH) in the two communities will be analyzed. Because this research involves minimal risk and would be impracticable with informed consent, a waiver of patient informed consent will be requested under 45 CFR 46.116 (c) or (d).

The protocol will be submitted to appropriate IRBs (a central and/or local site IRBs) for ethical review prior to study initiation. Any subsequent modifications to the protocol will be submitted to appropriate IRBs, and, at a minimum, the protocol will be submitted annually for continuing review and approval by these same ethics boards.

This component of the study involves an intervention using FIs to link HIV-positive individuals to HIV care after diagnosis. These incentives are described in detail in the intervention section for this study component. No additional incentives (for example for transportation) will be given to individuals who participate in this component of the study.

No individual data, other than what exists in the surveillance databases, will be collected from study participants in the Linkage-to-Care component of this study. To assess the impact of the study on Linkage-to-Care in the communities of Washington, D.C. and the Bronx, HIV surveillance data routinely collected by the DOH will be analyzed. Data from the DOHs will be provided to the study team in a de-identified fashion, as per usual HIV surveillance procedures.

No study-specific laboratory testing will be conducted under this protocol. Therefore, no additional study-related test results will need to be reported to authorities. HIV testing and care data collected during routine clinical care will be reported per local HIV and AIDS reporting requirements.
3.10 Safety Monitoring and Adverse Event Reporting

The study team will not collect or report Adverse Events because there is no biomedical intervention. However, the team will collect and report all social harms that are brought to the attention of study staff members, using a study-specific incident report form. This form will be anonymous and will query common social harms such as altered personal relationships, forced change in housing, and physical violence. The form will also include space for a written narrative to document additional details of any social harm experienced. All research staff will be trained to properly complete the form. As a part of study training, research staff will also be trained on the provision of referrals to counseling and social service support. Reports of social harms will be reviewed quarterly or more often, if indicated, and reported to the medical officer together with any actions that are taken. Social harms will be summarized and reported to appropriate IRB(s) on an annual basis.

4.0 VIRAL SUPPRESSION

4.1 Study Objectives for Viral Suppression

The primary feasibility objectives for using FIs to incentivize HIV-positive patients to achieve and maintain viral suppression (HIV RNA < 400 copies/mL) will be measured by determining the following:

- Proportion and number of patients in care, on ART (eligible for incentives), and receiving incentives upon achieving VL suppression
- Number of incentives disbursed compared with the total number of incentives available to eligible patients over the study duration [measurement of duration of viral suppression]
- Overall cost of program including staffing, infrastructure and incentives
- Number of patients previously adherent to ART who change from non-intervention to intervention care sites

In order to monitor the implementation of the FI program, certain parameters will be gathered from a subset of participating sites at various time points to ensure program quality. These parameters will include: the proportion and number of patients in care, on ART (eligible for incentives) and receiving incentives, as well as the incentive amount received compared with the total number on ART who would be eligible.

The primary effectiveness objective is to compare an FI intervention with the SOC for achieving and maintaining suppressed VL in HIV-positive patients in care, managed under the prevailing guidelines for the initiation of ART. For sites in each intervention arm, this objective will be measured by the following:

- Comparing the mean proportion of patients in care at a site who have suppressed VL (HIV RNA <400 copies/mL) during the fixed 12-month calendar evaluation period beginning 12 months after initiation of the FIs at that site
- Comparing the trend in mean proportion of patients in care at a site who have suppressed VL over time since initiation of FIs
Comparing the mean proportion of patients who have suppressed VL (HIV RNA < 400 copies/mL) after cessation of incentives

4.2 Design for Viral Suppression

The viral suppression component of the study is a two-arm, site-randomized, prospective, effectiveness clinical trial. It will be conducted within each intervention community to assess the effectiveness of a FI intervention for achieving and maintaining suppressed VL (HIV RNA < 400 copies/mL) in HIV-positive individuals (who have initiated ART under prevailing guidelines) compared to SOC for supporting patients in achieving adherence and virologic suppression. Each HIV care site will be randomized to either the intervention or SOC arm of the study. For this component of the trial, study outcomes will be evaluated through routinely reported HIV surveillance data. Using surveillance data, study outcomes will continue to be evaluated for a period of one year after the FI intervention ends.

In order to identify the SOC for ART adherence/viral suppression support, against which the incentives intervention will be assessed, we will develop a brief survey to be completed by an appropriate facility administrator, which will collect key attributes of HIV care sites, including types of ART adherence support, case management services, and other support services already available to patients. Data will be collected from sites annually from 2009 - 2013.

4.3 Study Population for Viral Suppression

The study population for the viral suppression component of the study will include all individuals ages 12 and older who are permitted to consent, or can be consented for HIV care by a parent/legal guardian according to New York State or Washington, D.C. law, who have initiated care at participating HIV care sites.

4.4 Study Sites for Viral Suppression

Twenty HIV care sites will be selected from Washington, D.C. and 20 HIV care sites will be selected from the Bronx to participate in this component of the study. Sites will be selected based on two primary criteria: 1) site agreement to participate in this component of the study, and 2) sites with the highest number of HIV-positive patients in care in the previous year. Additional site selection criteria may be considered if a total of 40 HIV care sites cannot be chosen based on the primary criteria.

HIV care sites will be randomized 1:1 to the FI vs. SOC, stratified by city and balanced by size of patient population in a clinic and baseline levels of proportion of VL showing suppression. In each city, these 20 sites will be randomized such that 10 will use the FI intervention, described in Section 4.5, to promote viral suppression in HIV-patients on ART and 10 will continue the existing SOC.

The site randomization will be balanced by the following two baseline characteristics:

- The size of the site’s HIV-positive patient case load
• The proportion of HIV-positive patients with VL suppression at each site

4.5 Intervention for Viral Suppression

HIV-positive individuals at HIV care sites assigned to the FI intervention will receive FIs (gift cards worth $70) to reinforce adherence to ART as measured by confirmation of each suppressed VL measurement (<400 copies/mL). The gift cards will be distributed at the HIV Care sites during each routine, quarterly clinic visit that a study participant demonstrates a suppressed VL.

In the event participants report full adherence to their providers but have not succeeded in achieving VL suppression, providers will be expected to obtain resistance testing, as recommended by current treatment guidelines, and to alter their patients’ regimens accordingly. Once patients with resistant HIV are on appropriate regimens, they will again be eligible for FIs for viral suppression. The specific procedures for HIV care sites to distribute the FIs for viral suppression will be outlined in the HPTN 065 SSP.

This intervention will be compared to SOC support for ART adherence through the use of routinely collected HIV surveillance data.

4.6 Study Procedures for Viral Suppression

At the initiation of the study, the study team, along with experts in treatment and management of HIV disease, will provide training to providers in the intervention communities on current HIV-treatment guidelines. Training will encourage providers to maximize initiation of ART per current guidelines and will emphasize the following: eligibility criteria for ART as defined in the guidelines; assessment of patient readiness to start ART (if eligible); prompt initiation of ART (if eligible); use of recommended regimens; the provision of supportive services to ensure adherence; regular clinical and laboratory monitoring; the importance of assessment for evidence of treatment failure; reinforcement of adherence; obtaining HIV resistance testing (if appropriate); and modification of failing regimen with new effective regimen. All HIV care site staff will also be trained on the procedures for the FI intervention prior to implementation. Along with this training, the importance of linkage to supportive services will be emphasized, including income assistance, housing support, substance use management and mental health services. Each HIV care site, regardless of the study arm it is randomized to, will be provided with a listing of available resources in the community with contact names and numbers. Providers will also be trained on HIV prevention counseling.

A baseline survey will be conducted to determine the availability of supportive services at all participating HIV care sites (e.g. social services, adherence support, substance use treatment, support groups, mental health resources, etc.) and to determine whether these services are available on site or by referral. In addition, an effort will be made to monitor the number of new referrals and waiting times for appointments to assess program capacity.

The specific procedures for HIV care sites to distribute the FIs for viral suppression will be outlined in the HPTN 065 SSP.
4.6.1 Monitoring HIV Care Sites

It is possible that members of the intervention communities, particularly those who are HIV-positive, will learn of the FI intervention to promote viral suppression and seek out HIV care sites offering these incentives. In an effort to minimize significant migration of patients from HIV care sites without FIs to ones that offer them, the following rules have been put into place:

- A patient must have at least one VL measurement at the HIV care site between three and nine months prior to the first suppressed VL for which he or she may receive an incentive. Both of these VL measurements must be done at the same HIV care site.

- All subsequent VL measurements for which a patient may receive FIs must be performed at the same HIV care site.

If a patient changes HIV care sites during the course of the study, he or she will be required to “establish care” at the new HIV care site prior to receiving any FI for a suppressed VL as indicated above. Specifically, to receive an FI again, the individual must have a VL measurement at the new HIV care site and return in three to nine months for a repeat VL measurement. The results of the second test will determine whether the person receives the FI or not.

Despite this barrier to HIV-care-site migration, the FI intervention may encourage this phenomenon; thus, the study team will use surveillance data to monitor the case load at each site at frequent intervals. By monitoring HIV care site case load over time, the team will be able to determine if significant migration is taking place. This will motivate discussions with the community advisory group to design other mechanisms to minimize such migration or if needed a redesign of this component of the study may be embarked on by the study team.

4.7 Study Duration for Viral Suppression

The duration for the FI intervention for the viral suppression component of the study is depicted in Figure 3. Participants will be eligible to receive FIs for suppressed (< 400 copies/mL) VL measurements once every three months throughout the entire 24-month study period. Participants who miss a quarterly visit(s) remain eligible for incentives as long as they present back to care with a suppressed VL measurement.

Patients with HIV diagnoses returning to care after a hiatus (of at least a year) and individuals who switch HIV care sites during the study are all eligible for the FIs. However they must have at least one VL measurement at three months (or more) at the same HIV care site as the qualifying VL measurement for which they can receive an FI.

Accrual of participants into the Viral Suppression component of the study will occur over the entire 24 months of the study. Some participants may receive up to nine FI payments if they are seen at the same HIV care site throughout the entire study, had been receiving care at that the same site before the study began, and maintain viral suppression throughout the 24 months.
4.8 Statistics and Data Analysis for Viral Suppression

4.8.1 Endpoints

4.8.1.1 Endpoints for the feasibility of using financial incentives in the HIV-positive population for viral suppression

The feasibility endpoints for the viral suppression component of the study will include the following:

- Number of individuals eligible for incentives and receiving incentives at a select subset of sites for select time points
- Cost of program including staffing, infrastructure and incentives

4.8.1.2 Endpoints for the Effectiveness of Viral Suppression

The effectiveness endpoints for the viral suppression component of the study will include comparing between sites in the intervention and SOC arms:

- Probability of an HIV-positive patient in care at a site having a suppressed VL (<400 copies/mL) in the 12-month calendar assessment period beginning 12 months after initiation of the assessment period
- Number of identified HIV-positive patients in care who have sustained viral suppression (see below)
The primary endpoint of the intervention is evaluated using all VL ordered at each site in the 12-month period beginning 12 months after initiation of FIs. An individual will be defined as in care at a site if two VL or CD4 assessments were ordered at that site in the year prior to initiation of this study, or if care (defined by two VL/CD4 assessments) is initiated at the site during the study. PLWHA will be assigned to sites (and thus arms) based on the site where they are in care.

To avoid ascertainment bias that would arise from missing VL assessments in the in-care cohort, we will impute whether VL is suppressed in each quarter of the 12-month assessment period a patient is in care (this is based on an SOC where VL assessments are ordered approximately every three months). For example, a patient with a VL assessment in each quarter will have an endpoint assessment for each quarter. A patient who has a suppressed VL assessment in the first, third and fifth quarters, but no assessment in the second and fourth, will be assumed to be suppressed in the second and fourth. However a patient with a suppressed VL in the first quarter and no subsequent assessment throughout the study will be assumed not suppressed in all subsequent quarters. The algorithm for imputing VL suppression (incorporating the sequence of VL assessments and including management of the reporting lag in surveillance data), inclusion of assessments in the quarters preceding and following the assessment period, handling of cases in treatment at more that one site, and censoring because of moving outside the surveillance area, death, etc. will be detailed in the statistical analysis plan. Contamination between arms will be tracked by detection of patients with VL assessments at multiple providers in the year.

It should be noted that the existing surveillance procedures include extensive efforts to link cases across jurisdictions. Individuals moving outside the jurisdiction will be censored at last measure prior to relocation.

4.8.2 Sample Size and Power

The top 20 provider sites in Washington, D.C. and the Bronx account for approximately 5,000 HIV patients in Washington, D.C. and 16,000 in the Bronx (exclusive of incarcerated patients in HIV care). It is assumed that the number of cases in care during the 24-month study duration will be similar to these past case loads on an annual basis. In the Bronx, VL was suppressed in 57% of 8,316 patients with VL assessed in 2008. The mean number of patients per clinic with a VL assessment in 2008 in the top 20 care sites was 400, ranging from 50 to 3,300. The ICC of proportion virally suppressed estimated from the Bronx 2008 data was 0.07. In Washington, D.C., 67% of 2,926 patients with VL assessments in 2008 achieved viral suppression at their most recent assessment. The mean number of patients per clinic with VL assessments was 245, ranging from 35 to 1,400 in the top 20 clinics. The ICC of proportion virally suppressed estimated from the Washington, D.C. 2008 data was 0.104.

Table 3 shows the mean number of patients with VL assessments needed per clinic, assuming 20 clinics in each arm (40 clinics in total), to detect a change in proportion of virally suppressed for a range of effect, power and ICC. If the clinics had a mean of 219 patients with VL assessments in the 12 months of evaluation, we would have 90% power
to detect an increase in viral suppression from 60% to 66% between the SOC and intervention clinics.

Sample size calculation for clustered designs use the approach described by Thomson, Hayes and Cousens (Thomson, Hayes et al. 2009).
Table 3.  Mean number of patients in care at a site required to achieve sufficient power to detect differences in proportion with suppressed VL. Number of clusters fixed at 20 per arm.

<table>
<thead>
<tr>
<th>Percent in care virally suppressed</th>
<th>Standard Deviance of percent virally suppressed</th>
<th>Intra-class correlation</th>
<th>Mean number of patients in care</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Standard of care</td>
<td>Intervention</td>
<td>Standard of care</td>
</tr>
<tr>
<td>60%</td>
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</table>
4.8.3 Randomization Scheme

In the viral suppression study component, facilities that provide HIV care services (HIV care sites) will be randomized to use FIs or the SOC.

HIV care sites will be assigned to one of the following two arms:

- Sites offering FIs for VL suppression: FIs offered to all those who are on ART, upon the confirmation of each suppressed VL measurement (<400 copies/mL)
- Sites continuing with SOC to encourage VL suppression: Each person on ART will be offered support via the site’s SOC procedures to attend HIV care site visits and remain adherent to their ART regimen in order to achieve and maintain VL suppression

The site randomization will be balanced by the following two baseline characteristics:

- The size of the site’s HIV-positive patient case load
- The proportion of HIV-positive patients with VL suppression at each site

4.8.4 Data Analysis

Data to be analyzed for this study component will be obtained from routine HIV surveillance data as well as from the participating sites. Participating sites will maintain tracking logs for FIs provided to participants maintaining suppressed VLs. Information from the tracking logs, with all participant-identifying information removed, will be provided to the study team.

The data for the effectiveness of an FI intervention for high ART adherence to achieve viral suppression compared to the SOC will be analyzed as described below. Analysis of secondary endpoints will be detailed in the statistical analysis plan.

4.8.4.1 Primary Effectiveness Analysis

The assessment cohort includes all HIV cases identified from HIV surveillance in the intervention city DOH jurisdiction that have a VL assessment in the 24 months of trial duration. The assessment period for each site begins 12 months after the onset of FIs in each city. An intention-to-treat approach will be used to assign patients to the care site assignment existing at the trial onset. For the 12-month assessment period (i.e. from 12 through 24 months), all VLs in the surveillance data assessed by the 40 clinics in the study are used to assess differences in percent achieving viral suppression by study arm. Details about each site’s contribution of patients who enter into care during the study will be provided in the statistical analysis plan. As described in 4.8.1.2, an imputation scheme for assessing VL suppression during the 12 months of the assessment that avoids the bias of missing assessments will be applied to all patients in care.
The estimate of intervention effect is the odds ratio of VL suppression in the FI intervention compared to SOC arms, assuming both a within-person and within-clinic correlation, stratified by city, using GEE methods. Sensitivity analysis will be conducted to assess the impact of patients who switch providers or establish care at multiple providers during the trial.

4.8.4.2 Feasibility Endpoints

Descriptive statistics will be used to summarize the mean uptake and overall cost of FIs and its variation between sites by site and population characteristics.

4.9 Human Subjects/Ethical Considerations

The Viral Suppression component of the study involves an intervention using FIs to encourage HIV-positive individuals on ART to be adherent to their medications and to maintain HIV VL suppression. This component is public health research. In this component, HIV care sites will be randomized, not individuals, and no de novo individual-level data will be collected. Instead, (de-identified) HIV surveillance data routinely collected by the DOH in the two communities will be analyzed. Because this research involves minimal risk and would be impracticable with informed consent, a waiver of patient informed consent will be requested under 45 CFR 46.116 (c) or (d).

The protocol will be submitted to appropriate IRBs (a central and/or local site IRBs) for ethical review prior to study initiation. Any subsequent modifications to the protocol will be submitted to appropriate IRBs, and, at a minimum, the protocol will be submitted annually for continuing review and approval by these same ethics boards.

No individual data, other than what exists in the surveillance databases, will be collected from study participants in the viral suppression component of this study. To assess the impact of the study on viral suppression in the communities of Washington, D.C. and the Bronx, surveillance data routinely collected by the DOHs will be analyzed. Data from the DOHs will be provided to the study team in a de-identified fashion, as per usual HIV surveillance procedures.

No study-specific laboratory testing will be conducted under this protocol. Therefore, no additional study-related test results will need to be reported to authorities. HIV testing and care data collected during routine clinical care will be reported per local HIV and AIDS reporting requirements.

4.10 Safety Monitoring and Adverse Event Reporting

The study team will not collect or report Adverse Events because there is no biomedical intervention. However, the team will collect and report all social harms that are brought to the attention of study staff members, using a study-specific incident report form. This form will be anonymous and will query common social harms such as altered personal relationships, forced change in housing, and physical violence. The form will also include space for a written narrative to document additional details of any social harm experienced. All research staff will be trained to properly complete the form. As a part
of study training, research staff will also be trained on the provision of referrals to counseling and social service support. Reports of social harms will be reviewed quarterly or more often, if indicated, and reported to the medical officer together with any actions that are taken. Social harms will be summarized and reported to appropriate IRB(s) on an annual basis.

5.0 PREVENTION FOR POSITIVES

5.1 Study Objectives for Prevention for Positives

The primary objective of the Prevention for Positives study component is to evaluate the effectiveness of a computer-delivered counseling intervention (“CARE+ Prevention”) containing Prevention for Positives messages in addition to SOC compared to SOC alone. Both intervention and control groups will have access to available SOC support and counseling services available to HIV-positive patients at the participating HIV care sites. All participants will have regular assessment of risk behaviors. The objective for this study component will be measured by comparing key self-reported HIV-transmission sexual risk behaviors.

5.2 Design for Prevention for Positives

The Prevention for Positives component of the study is a two-arm, individually randomized, prospective, effectiveness trial to be conducted at 12 HIV care sites (6 within each intervention community). This component will compare CARE+ containing Prevention for Positives messages in addition to SOC with SOC alone. All consenting individuals will be randomly assigned to one of the two arms of the study. In this component of the study, participant informed consent will be obtained for all participants in both of the study arms. Limited medical records data pertaining to HIV disease and individual data will be abstracted and analyzed in this study component.

5.3 Study Population for Prevention for Positives

The Prevention for Positives component of the study will include HIV-positive patients at a select number of participating HIV care sites. These patients will include individuals newly diagnosed with HIV and linked to care, and those established in care. ART use is not required for participation.

5.3.1 Inclusion Criteria

The inclusion criteria for this study component are as follows:

- All individuals who are permitted to consent for HIV care according to New York State or Washington, D.C. law
- Receiving care at the selected HIV care sites in the Bronx or Washington, D.C.
- Have attended the clinic one or more times in the last seven months
• Able to understand either spoken English or Spanish
• Able and willing to provide informed consent

Subjects enrolled into the Prevention for Positives component of the study will participate in the patient survey (see Section 6.0).

5.3.2 Exclusion Criteria

The exclusion criteria for this study component are the following:

• Not seen in the clinic in the last seven months
• History or evidence of altered mentation, inebriation or substance use that would interfere with participation in the study
• Unable or unwilling to provide informed consent
• Participation in another study focusing on HIV prevention for positives

5.4 Study Sites for Prevention for Positives

Participants will be recruited for the Prevention for Positives component of the study from six HIV care sites (three FI sites and three SOC of care sites) in each intervention city.

The three sites among the FI sites and the three sites in the SOC sites in each city will be selected based on the following criteria:

• The highest volume of HIV-positive patients in care
• Site agreement to participate in this component of the study

5.5 Intervention for Prevention for Positives

A modified version of CARE+ for HIV-positives will be used with Audio Computer Assisted Self-Interviewing (ACASI) to measure behavioral risk and to compare a full computer counseling intervention (assesses self-efficacy and motivation, provides tailored feedback on specific risk behaviors, shows skill-building videos, and helps the user make a risk-reduction plan) to a control session consisting only of ACASI ascertainment of risk behavior.

An additional component will be the utilization of the computer-based system described above to measure factors that impact both the acceptability of the interventions utilized in this study and the feasibility of future TNT strategies. Other data, beyond those that can be obtained from routine HIV surveillance data, will be captured including the following:

• HIV testing history and prior experience with linkage/drop out from care (if applicable)
• CD4 cell count (via chart abstraction, not routinely available from HIV surveillance), at study enrollment, and then every 3 months up to and including month 12

• VL (via chart abstraction, not routinely available from HIV surveillance), at study enrollment, and then every 3 months up to and including month 12

CD4 cell counts and HIV RNA levels obtained from month 0 through month 12, will be extracted from participant’s medical records to enable individual-level subgroup analyses of adherence and virologic responses among CARE+ participants who are receiving versus not receiving FIs for virologic suppression, as well as other subgroup analyses, such as:

• Behaviors likely to transmit HIV (sexual and parenteral)

• Receipt of and experience with services to promote retention and adherence

• Knowledge of, attitudes towards, and the feasibility of early initiation of ART

• Knowledge of and attitudes towards use of ART for prevention in partners

5.6 Study Procedures for Prevention for Positives

5.6.1 Recruitment Process

Study staff at the HIV care sites will be trained for the Prevention for Positives study procedures and on human subject requirements. Clinic or study staff will systematically approach potential subjects in the waiting room or during consultation and inform them of the study. Brief, anonymous demographic data will be noted for those individuals refusing study participation, to assess comparability to study participants. Persons who are interested in the study will be assessed for eligibility and consented in a private area of the clinic. The study staff will provide information to the participant on the use of the tablet computer and headphones, and assist the participant with the anonymous ID log-in and beginning of the session, after which the staff person will remain nearby only to provide assistance if the participant requests it.

5.6.2 Co-Enrollment Guidelines

Prevention for Positives study participants will be allowed to be enrolled in other studies except those focused on reducing the risk of HIV transmission.

5.6.3 Participant Retention

Once a participant enrolls in the Prevention for Positives component of this study, the research staff will make every effort to retain him/her for 12 months of follow-up in order to minimize possible bias associated with loss to follow-up.

5.6.4 Participant Withdrawal
Participants may voluntarily withdraw from the study for any reason at any time. Participants also may be withdrawn by the study sponsor, site IRBs, government or regulatory authorities, the Investigator of Record (after consultation with the Protocol Chair), or Division of AIDS (DAIDS) Medical Officer.

Participants who elect to withdraw from the study prior to month 12 will be asked to complete one last computer session, however, participants will not be required to do so. Study staff will record the reason(s) for all withdrawals from the study in participants’ study records.

5.7 **Study Duration for Prevention for Positives**

Duration of the Prevention for Positives study component is 12 months.

Patients enrolled in the Prevention for Positives component of the study will complete the CARE+/ACASI tool at baseline (month 0) and every three months thereafter for one year (at months 3, 6, 9, and 12).

5.8 **Safety Monitoring and Adverse Event Reporting**

Physical and psychological risks of this study are expected to be minimal, as no medical/surgical or pharmacologic procedures are planned. The Prevention for Positives study component addresses sensitive behaviors around sex and social interactions. Questions regarding sex may cause embarrassment or discomfort. Participants will be informed in writing and verbally that they may skip any questions and/or drop out of the study at any time without repercussion. Mental stress identified in the course of the study, such as depression, suicidality and intimate partner violence (IPV) will be followed-up appropriately, with counseling services available through the clinics. Counseling services for any other issues may be requested by participants during their routine appointment at the HIV care site.

Researchers will do everything possible to emphasize and maintain the confidentiality of participants, and safeguards for protecting confidentiality of data will be strictly enforced.

The study team will not collect or report Adverse Events because there is no biomedical intervention. However, the team will collect and report all social harms that are brought to the attention of study staff members, using a study-specific incident report form. This form will be anonymous and will query common social harms such as altered personal relationships, forced change in housing, and physical violence. The form will also include space for a written narrative to document additional details of any social harm experienced. All research staff will be trained to properly complete the form. As a part of study training, research staff will also be trained on the provision of referrals to counseling and social service support. Reports of social harms will be reviewed quarterly or more often, if indicated, and reported to the medical officer together with any actions that are taken. Social harms will be summarized and reported to appropriate IRB(s) on an annual basis.

5.9 **Statistics and Data Analysis for Prevention for Positives**
5.9.1 Endpoints

The primary endpoint for the effectiveness of the Prevention for Positives component of the study is the following:

- The proportion of participants reporting any unprotected vaginal or anal sex the last time they had sex, evaluated for all partners and also separately for primary and non-primary partners.

The secondary endpoints for the effectiveness of CARE+ Positive Prevention component of the study are the following:

- Proportion of those who had unprotected vaginal or anal sex the last time they had sex with negative or unknown HIV status partners, evaluated for all partners and also separately for primary and non-primary partners.

- The frequency of participants reporting any unprotected vaginal or anal sex in the previous three months with non-primary partners.

- The number of different persons with whom the participant shared needles or works (including cookers and cottons) in the previous three months.

5.9.2 Accrual, Follow-up, and Sample Size

The primary behavioral endpoint of this study component is the proportion of patients reporting any unprotected vaginal or anal sex the last time they had sex. In the Strategies for Management of Antiretroviral Therapy (SMART) study, at 11.2% of visits participants reported any unprotected anal or vaginal sex in the previous two months. Using post-enrollment data; 5.4% of patient visits included a report of unprotected anal or vaginal sex with an HIV-uninfected partner in the previous two months.

Individual randomization will occur, potentially stratified by gender.

Using a 2-sided alpha of 5%, and calculating the sample size required to achieve 90% power to detect differences in percentages of participant visits where high-risk behaviors are reported in each arm, Table 4 presents a number of assumptions. A within-person, ICC of 0.3-0.4 is assumed, based on data from the SMART trial behavioral endpoints.

Assuming at least one follow-up visit for 90% of enrollees, and an average number of visits per participant of 3.75 amongst retained participants, the study will enroll 660 per arm (1320 in total) to allow for loss due to attrition and loss to follow-up. It is estimated that around 110 participants will be recruited per HIV care site, with a target enrollment of two subjects/day per site. Target enrollment should be reached within eight-12 months of initiation, and accrual will be terminated at 1320 participants or after approximately 12 months of accrual, whichever occurs first.

It is anticipated that a total of approximately 1320 individuals will be randomized. Data will be collected from each person an average of 4 times during the study (out of four possible visits following enrollment at month 0, i.e. months 3, 6, 9, and 12). Allowing for
loss to follow-up, with 522 people available for assessment per arm, the study has 90% power to detect a decrease from 11% to 8% in proportion of patients reporting any unprotected vaginal or anal sex at a given visit during the study.
Table 4:  Per-Arm Sample Size Required for 90% Power to Detect Difference in Proportion Reporting High Risk Behavior at a Visit, with Two-sided Alpha of 5%, Assuming Four Visits per Participant

<table>
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<tr>
<th>Control (%)</th>
<th>Intervention (%)</th>
<th>Mean Number of Visits</th>
<th>Intra-class Correlation</th>
<th>Number of People per Arm</th>
<th>Intra-class Correlation</th>
<th>Number of People per Arm</th>
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<tr>
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5.9.3 Randomization Scheme

Randomization to intervention (SOC and full CARE+ Prevention counseling session) or control arm (SOC alone) is done automatically within the software application using a pseudo-random number generator.

5.9.4 Data Analysis

Data to be analyzed for this study component will be obtained from the CARE+ electronic storage system and abstracted from participant’s medical records. As participants proceed through the CARE+ intervention, their answers will automatically be stored electronically in a secure system. Access to the storage system will be limited to essential personnel. CD4 cell counts and HIV RNA levels obtained from month 0 through month 12, will be extracted from participant’s medical records to enable individual-level subgroup analyses of adherence and virologic responses among CARE+ participants who are receiving versus not receiving FIs for virologic suppression, as well as other subgroup analyses.

The primary behavioral endpoint of this study component is the proportion reporting any unprotected vaginal or anal sex the last time they had sex.

Effectiveness of the CARE+ Prevention for Positives component will assess differences in proportion reporting self-reported risk behavior for participants in the CARE+ Prevention arm compared to the control arm. GEE methods for repeated measures will be used to assess the difference in proportion reporting high-risk behavior between the two study arms for participants. Major subgroup analyses for those on ART and not on ART will be conducted.

5.10 Human Subjects/Ethical Considerations

In the Prevention for Positives component of the study, a subset of patients enrolled at select HIV care sites in the two intervention communities will be randomized either to an intervention arm (receiving SOC prevention activities plus a computer-delivered intervention) or to the control arm (receiving only the SOC prevention activities at the care site). Individual-level data will be collected and analyzed. This component is a public health research requiring informed consent. Written informed consent will be obtained from participants.

The protocol and supporting study materials (the informed consent form and any advertising materials) will be submitted to appropriate IRBs (a central and/or local site IRBs) for ethical review prior to study initiation. Any subsequent modifications to these materials will be submitted to appropriate IRBs, and, at a minimum, they will be submitted annually for continuing review and approval by these same ethics boards.

Written informed consent will be obtained from each study participant (or a mark for those who are illiterate, which will be witnessed by a third party) prior to study enrollment. Each study site is responsible for developing an informed consent form for local use, based on the template provided with this protocol that describes the purpose of...
the study, the study procedures and the risks and benefits of participation, in accordance with all applicable regulations. Participants will be provided with a copy of their informed consent form if they wish to receive it.

A small incentive will be provided to study participants in the Prevention for Positives component to compensate them for transportation and time.

All study-related information will be stored securely at the study site in areas with access limited to study staff. To maintain participant confidentiality, a coded number will identify all study data and administrative forms. All local databases will be secured with password-protected access systems. Forms, lists, logbooks, appointment books and any other listings that link participant ID numbers to other identifying information will be stored in a separate area with limited access. A participant’s study information will not be released without the written permission of the participant, except as necessary to: authorized medical care providers; monitors for the National Institute of Allergy and Infectious Diseases (NIAID) and/or its contractors; representatives of the HPTN Coordinating and Operations Center (CORE) and/or the (Statistics and Data Management Center) SDMC; other government and regulatory authorities; and/or the site IRB.

No study-specific laboratory testing will be conducted under this protocol. Therefore, no additional study-related test results will need to be reported to authorities. HIV care data collected during routine clinical care will be reported per local HIV and AIDS reporting requirements.

6.0 SURVEY OF PATIENTS AND PROVIDERS

6.1 Study Objectives for the Survey of Patients and Providers

The objectives of the survey of patients and providers study component are the following:

- To assess knowledge, attitudes and practices with regard to ART initiation, the potential of starting ART at higher CD4 cell counts, and the use of ART for prevention of HIV transmission

- To ascertain key socio-demographic and behavioral characteristics of study participants that cannot be obtained from routine HIV surveillance data

- To assess the acceptability and attitudes towards FIs

6.2 Design for the Survey of Patients and Providers

All participants in the Prevention for Positives component of the study will complete a patient survey module prior to and at the end of that study component. The survey will assess knowledge and attitudes regarding the use of ART in HIV disease and other key factors related to the feasibility and acceptability of the study’s interventions.

Clinical providers who prescribe ART at HIV care sites in the intervention communities will be invited to complete a survey regarding their knowledge, attitudes and practices about the use of ART in HIV patients as well as FIs. Clinical providers will be surveyed.
prior to and following the viral suppression intervention to assess trends in the provider knowledge, attitudes and practices.

6.3 Study Population for the Survey of Patients and Providers

The patient survey component of the study will include HIV-positive patients at select HIV care sites that are participating in the Prevention for Positives component of the study.

Prescribing clinical providers (e.g., physicians, nurse practitioners and/or physician assistants) at select HIV care sites in the Bronx and Washington, D.C. will participate in provider survey.

6.3.1 Inclusion Criteria

The inclusion criteria for the patient survey are identical to those for the Prevention for Positives component of the study.

Prescribing clinical providers (e.g., physician, nurse practitioner/nurse-midwife, or physician assistant) at select HIV care sites are eligible for the provider survey.

6.3.2 Exclusion Criteria

The exclusion criteria for the patient survey are identical to those for the Prevention for Positives component of the study.

There are no exclusion criteria for the provider survey.

6.4 Study Sites for the Survey of Patients and Providers

The patient survey will be conducted at those HIV care sites involved in the Prevention for Positives component. The provider survey will survey providers from participating HIV care sites in the Bronx, NY and Washington, D.C.

6.5 Study Procedures for the Survey of Patients

Once consented and enrolled in the Prevention for Positives component, the clinic study staff will train participants how to use the computer tablet. Randomization to intervention (SOC plus full CARE+/ACASI counseling session) or control arm (SOC alone) is done automatically within the software application following the user’s login with an anonymous study ID. Up to 110 individuals will be assigned to each arm, per site. At the end of the first (baseline, month 0) session, both arms will be presented, via the computer tool, with a series of questions (the patient survey). This patient survey module will be repeated only one more time, at the end of the intervention (month 12).
6.6 **Study Procedures for the Survey of Providers**

Providers at participating HIV care sites in both cities will be recruited by an introductory letter followed by up to two e-mail reminders, then two phone calls by study staff, asking them to take the brief survey on a secure, anonymous website.

Providers who go to the Web site will read and click on a brief consent, then complete a short survey assessing attitudes and practices.

At the completion of the survey, a printable coupon, compensating them for their time will be generated.

6.7 **Study Duration for the Survey of Patients and Providers**

Participants in the patient survey will complete the survey upon enrollment in the Prevention for Positives study and at the end of that study component.

The provider survey will also take place twice, once at the time of initiation of the overall HPTN 065 study and at the end of the viral suppression component.

6.8 **Statistics and Data Analysis for Surveys of Patients and Providers**

Data for analysis of this study component will be derived from answers to survey questions provided by both the patients and providers. Patient surveys will be administered electronically, after the CARE+ intervention at the beginning and at the end of the intervention. All patient responses will be stored in an electronic and secure database. Limited personnel will have access to this database. Provider surveys will be administered over the Internet. The survey will be secure and responses will also be stored securely. Survey data will be provided to the Statistical Center for HIV/AIDS Research and Prevention (SCHARP) for the following analyses:

- To assess baseline knowledge, attitudes and practices regarding the use of ART and other key factors related to the feasibility and acceptability of the study interventions in the subset of survey participants in the Prevention for Positives study component

  Descriptive statistics will be used to summarize the survey participants’ attitudes regarding the use of ART and other key factors related to the feasibility and acceptability of the study interventions by site and combined.

- To compare the providers’ knowledge, attitudes and practices about the use of ART, assessing these at study baseline and at study end at HIV care sites in the intervention communities

  Descriptive statistics will be used to summarize the prescribing providers’ knowledge, attitudes and practices about the use of ART, at study baseline and study end by site and combined. Chi-squared statistics will be used to compare the baseline data with study end data.
All Prevention for Positives data collected on the tablet computer will be password security-protected (anonymous study ID assigned timed to the millisecond of initial user log-in, with no way for patients to access other user’s data). All study data will be accessible only to investigators, study staff and biostatisticians assisting with the analysis. The CARE+ Prevention application will reside on each tablet computer with encrypted data transferred to a secured server at each clinic. The session database will be backed up by study staff. The 12 HIV care sites will send their study data to a SCHARP server on a routine basis. Only authorized study staff members are able to log onto the tablet computers or to access the master study database.

6.9 Human Subjects/Ethical Considerations

All patients participating in the Prevention for Positives study component will be administered the Patient Survey, regardless of whether or not they are randomized to the CARE+/ACASI intervention. Individual-level data will be analyzed. This component is human subjects research. Written informed consent will be obtained from participants at the time of consenting for participation in the Prevention for Positives study component.

Providers from participating HIV care sites will be invited to complete a Web-based survey regarding their knowledge, attitudes and practices concerning ART for treatment, frequency of HIV testing, and limited sociodemographic information. The survey will be anonymous, and will collect no identifying information. This component is public health research. Informed consent will be obtained from the provider’s completing the survey. Before the survey will display, providers will read the consent and if they agree to participate, will indicate their agreement by clicking a button labeled “I agree.”

The protocol and supporting study materials (the informed consent form and any advertising materials) will be submitted to appropriate IRBs (a central and/or local site IRBs) for ethical review prior to study initiation. Any subsequent modifications to these materials will be submitted to appropriate IRBs, and, at a minimum, they will be submitted annually for continuing review and approval by these same ethics boards.

Written informed consent (or a mark for those who are illiterate, which will be witnessed by a third party) will be obtained from each study participant willing to complete the patient survey prior to study enrollment. The informed consent for the survey is incorporated into the consent form for the Prevention for Positives component. Each study site is responsible for developing an informed consent form for local use for the patient survey, based on the template provided with the protocol, that describes the purpose of the survey, the survey procedures, and the risks and benefits of participation, in accordance with all applicable regulations. Participants completing the patient survey will be provided with a copy of their informed consent form if they wish to receive it.

No physical or psychosocial risks are anticipated from participation in the patient or provider surveys as they involve no invasive procedures or reports of illegal or socially stigmatized behaviors. Further, all data will be anonymized once merged into the study database, and no names will be used in publications.

A small incentive will be provided to survey participants, both providers and patients, to compensate them for their time. For the patients, this compensation is included in
the incentive they receive for participating in the Prevention for Positives component of the study.

For patient surveys, all study-related information will be stored securely at the study site in areas with access limited to study staff. To maintain participant confidentiality, a coded number will identify all study data and administrative forms. All local databases will be secured with password-protected access systems. Forms, lists, logbooks, appointment books and any other listings that link participant ID numbers to other identifying information will be stored in a separate area with limited access. Survey information will not be released without the written permission of the participant, except as necessary to: authorized medical care providers; monitors of the NIAID and/or its contractors; representatives of the HPTN CORE and/or the SDMC; other government and regulatory authorities; and/or the site IRB.

6.10 Safety Monitoring and Adverse Event Reporting

The study team will not collect or report Adverse Events because there is no biomedical intervention. However, the team will collect and report all social harms that are brought to the attention of study staff members, using a study-specific incident report form. This form will be anonymous and will query common social harms such as altered personal relationships, forced change in housing, and physical violence. The form will also include space for a written narrative to document additional details of any social harm experienced. All research staff will be trained to properly complete the form. As a part of study training, research staff will also be trained on the provision of referrals to counseling and social service support. Reports of social harms will be reviewed quarterly or more often, if indicated, and reported to the medical officer together with any actions that are taken. Social harms will be summarized and reported to appropriate IRB(s) on an annual basis.

7.0 HIV SURVEILLANCE, ROUTINELY-COLLECTED AND OTHER SURVEY DATA

Use of HIV case and behavioral surveillance data and HIV testing data will be used for site selection in intervention communities and for process and outcome measures. Other data, including site provided data, may be utilized for these purposes in certain circumstances.

We will assess completeness, accuracy, timeliness and performance of the surveillance and any other data.

7.1 HIV Testing Data

HIV testing data will be used to select HIV test sites for randomization to the intervention vs. SOC arm, and as a process measure for the testing intervention in ED and inpatient facilities (Table 5). For selection of HIV test sites for interventions, information on HIV testing is available for HIV test sites that are publicly supported through federal funds in each of the six communities included in this study; additional data are available for other
HIV test sites supported in some local jurisdictions, such as NYC’s AIDS Institute Reporting System. This includes the majority of HIV testing conducted by CBOs.

Table 5. Data Sources and Definitions

<table>
<thead>
<tr>
<th>Information needed</th>
<th>Description</th>
<th>Data source</th>
<th>Variables</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HIV Testing Data</strong></td>
<td><strong>Site selection, intervention communities</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Volume of testing</td>
<td>Number of persons tested in XX year, by testing site</td>
<td>Local PEMS, local testing data</td>
<td>Name of testing facility, testing date</td>
</tr>
<tr>
<td></td>
<td>Number of new diagnoses, by testing site</td>
<td>HIV surveillance data</td>
<td>Name of facility of diagnosis, date of diagnosis</td>
</tr>
<tr>
<td></td>
<td>Where available, number of case reports to surveillance, by testing site</td>
<td>HIV surveillance data</td>
<td>Name of facility of diagnosis, date of diagnosis</td>
</tr>
<tr>
<td><strong>Process measures for HIV testing interventions, intervention communities</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of persons eligible</td>
<td>Number of persons eligible</td>
<td>Project specific collection at intervention sites</td>
<td>Data may or may not already be collected at selected sites. If not collected, collection will be implemented for this study</td>
</tr>
<tr>
<td>Number of persons tested</td>
<td>Number of persons tested</td>
<td>Project specific collection at intervention sites</td>
<td>Data may or may not already be collected at selected sites. If not collected, collection will be implemented for this study</td>
</tr>
<tr>
<td>Number of new diagnoses</td>
<td>Number of new diagnoses</td>
<td>HIV surveillance data, and project specific collection at intervention sites</td>
<td>Surveillance data: name of testing facility, testing data. Data may or may not be collected by selected sites, and if not collected, can be implemented at site or use surveillance data.</td>
</tr>
<tr>
<td><strong>HIV Surveillance Data</strong></td>
<td><strong>Outcome Measures</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number/proportion of persons newly diagnosed who entered care, by site</td>
<td>CD4 or VL within 3 months of diagnosis (CD4/VL date not equal to diagnosis date), by site</td>
<td>HIV surveillance data</td>
<td>HIV diagnosis date, name of testing site, VL date, VL result, CD4 date, CD4 result, demographics (DOB, sex, race/ethnicity, transmission category)</td>
</tr>
</tbody>
</table>
## Data Sources and Definitions

<table>
<thead>
<tr>
<th>Information needed</th>
<th>Description</th>
<th>Data source</th>
<th>Variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number/proportion of previously diagnosed persons not in care entered into care, by site</td>
<td>CD4 or VL within 3 months of diagnostic test, among persons with previous diagnostic test and no CD4/VL within the past year, by site</td>
<td>HIV surveillance data</td>
<td>HIV diagnosis date, name of testing site, VL date, VL result, CD4 date, CD4 result, demographics (DOB, sex, race/ethnicity, transmission category)</td>
</tr>
<tr>
<td>VL suppression</td>
<td>Probability of undetectable VL amongst people established in care with previous VL/CD4, by site</td>
<td>HIV surveillance data</td>
<td>Name of treatment site, VL date, VL result, CD4 date, CD4 result, demographics (DOB, sex, race/ethnicity, transmission category)</td>
</tr>
</tbody>
</table>

### Behavioral Data

<table>
<thead>
<tr>
<th>Information needed</th>
<th>Description</th>
<th>Data source</th>
<th>Variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number/proportion of persons who had an HIV test in the past 12 months</td>
<td>See BRFSS 2009 questionnaire: 20.1 Have you ever been tested for HIV? Do not count tests you may have had as part of a blood donation. Include testing fluid from your mouth. (213) 1 Yes 2 No [Go to Q20.5] 7 Don’t know / Not sure [Go to Q20.5] 9 Refused [Go to Q20.5] 20.2 Not including blood donations, in what month and year was your last HIV test? <a href="http://www.cdc.gov/brfss/questionnaires/pdf-ques/2009brfss.pdf">http://www.cdc.gov/brfss/questionnaires/pdf-ques/2009brfss.pdf</a></td>
<td>Local surveys (see descriptions below)</td>
<td>Testing, testing date</td>
</tr>
<tr>
<td>Special HPTN 065 emphasis population: number/proportion of MSM who had an HIV test in the past 12 months</td>
<td>NHBS questions</td>
<td>National HIV Behavioral Surveillance (NHBS) system</td>
<td>Testing, testing date</td>
</tr>
</tbody>
</table>

The CDC has developed a data entry and reporting system, the PEMS, to strengthen monitoring and evaluation of HIV prevention programs. PEMS is used by health departments and CBOs funded through CDC HIV-prevention cooperative agreements. PEMS allows grantees to collect agency data, community planning data, program plan data, and client-level data. This assures a comprehensive set of standardized variables are available.

The client-level data include information on testing site, client demographics, risk factors, test information and testing history. This allows a description, by site, of the number...
tested and number positive, and a description of the population tested or positive. Additional variables that can be used to describe sites include agency characteristics (budget, sites, workers, contracts, network agencies), program plans (program models, target populations, interventions (CTR, Health Communications/Public Information Outreach, settings, sessions, activities), service delivery (service activities, recruitment, referrals), and community planning activities (target populations, priority interventions).

Some areas have testing report systems in addition to PEMS. For example, New York State collects testing data through the AIDS Institute Reporting System. Florida has PEMS data as well as a counseling and testing data base that includes all the variables needed to report an HIV case as well as testing history questions; these two data sources cover all clients tested in the public sector, e.g., registered counseling and testing sites (AIDS Service Organizations (ASOs), CBOs, faith-based initiatives), and county health department clinics (STD, TB, Family planning, etc.).

Another testing measure for selection of intervention sites that can be derived from surveillance data is the number of new diagnoses by site.

Process measures (Table 5) to monitor testing at the testing intervention sites will be collected directly from the participating facilities.

### 7.2 HIV Surveillance Data

The CDC’s national system for the surveillance of HIV infection is based on mandatory name-based reporting of all HIV and AIDS cases in every state and Washington, D.C. Health departments maintain case records of every new and established HIV case according to the HIV case definition. Laboratory reports of positive HIV test results and provider reports of new confirmed diagnoses are tracked by the health departments. In most jurisdictions, data from mandatory reporting of CD4 cell count and VL from all laboratories are linked via case names to individuals maintained in the eHARS data entry and reporting system (not all states have mandatory reporting of all VL and CD4 cell count). Across the United States, and within each community, the system captures key variables for this study. The key data elements from HIV surveillance that will be used are listed in Table 6.

#### Table 6. Key Date Elements and Primary Use

<table>
<thead>
<tr>
<th>Key Data Elements</th>
<th>Primary Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD4 result</td>
<td></td>
</tr>
<tr>
<td>CD4 date</td>
<td></td>
</tr>
<tr>
<td>VL result</td>
<td></td>
</tr>
<tr>
<td>VL date</td>
<td></td>
</tr>
<tr>
<td>Name of facility of diagnosis</td>
<td>Study intervention outcome measures</td>
</tr>
<tr>
<td>Name of treatment facility (lab order)</td>
<td></td>
</tr>
<tr>
<td>Type of HIV test</td>
<td></td>
</tr>
<tr>
<td>HIV test date</td>
<td></td>
</tr>
<tr>
<td>HIV test result</td>
<td></td>
</tr>
<tr>
<td>HIV diagnosis date</td>
<td></td>
</tr>
<tr>
<td>AIDS diagnosis date</td>
<td></td>
</tr>
</tbody>
</table>
De-duplication (removing double-reported cases) of cases between jurisdictions occurs at least annually, with case tracing on an ongoing basis as individuals relocate and new diagnostic labs are investigated in a new jurisdiction. Cases are identified through active follow-up of positive Enzyme Immunoassay (EIA), positive WB and detectable VL reports from names not matched in the local eHARS database, as well as through reporting by providers. Communities selected for this study (both intervention and non-intervention communities) have quality data collected on these parameters.

HIV surveillance data can describe treatment success based on reported VL measurements. However, specific information on type of treatment received is not collected.

HIV surveillance data have been evaluated on the national level and for individual state/local program areas for completeness, timeliness, and quality of individual data items (Buehler, Berkelman et al. 1992; Rosenblum, Buehler et al. 1992; Greenberg, Hindin et al. 1993; Meyer, Jones et al. 1994; Klevens, Fleming et al. 1998; Schwarcz, Hsu et al. 1999; Solomon, Flynn et al. 1999; Jara, Gallagher et al. 2000; Doyle, Glynn et al. 2002; Hall, Song et al. 2006) and through annual progress reports [unpublished]). For the accuracy of case counts, all surveillance programs conduct annual linkage of HIV cases to death records, intrastate duplicate reviews, and interstate duplicate resolution based on potential duplicate listings received from CDC.

CDC sets standards for the completeness and timeliness of case reporting as well as the completeness and accuracy of individual data elements, against which national and individual program data are evaluated each year (e.g., 2008 diagnosis data are currently evaluated using data reported through December 2009) (Hall and Mokotoff 2007; Hall, Song et al. 2008). For 2007 diagnoses, New York City, Philadelphia, and Florida case counts were estimated to be >95% complete by December 2008 (note, while the other cities were transitioning to the new data system and completeness was not formally assessed, they closely monitor expected case reports from laboratories and providers to assure complete reporting). More than 80% of cases are reported to the health departments within 6 months of diagnosis. Critical variables include sex assigned at birth, race/ethnicity, and age; completeness in these variables is very high (near 100% for sex and age; >90% for race/ethnicity) and active follow up is conducted to obtain these data elements; all cases must have a diagnosis date (test result or physician diagnosis,
according to the CDC case definition). Additional variables of importance include risk factor information, and CD4 and VL test results. Risk factor information is available for more than 60% of cases (some areas achieve better than 80%); CDC has developed methods to adjust for missing risk factor information in local or national analyses using multiple imputation.

The completeness of CD4 and VL results, critical to this study, depends on 1) state laboratory reporting laws and regulations; 2) reporting/abstracting from providers; and 3) whether patients entered care. Most important to the completeness of these data is the assessment of whether all laboratories routinely report to the health departments to assure that this information is reported for any patients entering care. For example, both New York State and DC have laws requiring reporting of all values of CD4 and VL test results. The New York State Health Department has identified all labs conducting testing for New York residents and monitors all labs to assure that reporting is complete and there are no missing data. Accuracy is also assessed by comparing reports from labs against what is obtained from chart review and re-abstractation. Regarding lag times, the mean time for New York City receiving Western Blot lab results is 21 days from diagnosis and CD4 and VL results 30 days from test date. (Please see Appendix for NYC example of additional indicators and evaluation of reporting completeness and timeliness). In DC, similar processes have been implemented. Regarding lag times in DC, approximately 85% of District laboratory reports are received within 2 weeks of the test date via electronic reporting. The remaining 15% are reported via the US mail, averaging about a 3 week lag in reporting. (Please see Appendix for DC indicators).

Data quality is continuously monitored during surveillance activities. Any notification (e.g., HIV diagnostic lab report) initiates data abstraction by field staff, with assignments within days of report. Abstracted data is checked for quality and the data entry system includes data quality checks. In order to assist in the timely input and generation of surveillance data during this study, resources will be provided for additional personnel in the DOHs in all participating cities (intervention and non-intervention). As part of TLC Plus, both process and outcome evaluations for data quality (accuracy, completeness, and lag times) will be conducted for intervention and non-intervention cities on a routine schedule.

CDC funds the Morbidity Monitoring Project (MMP) (McNaghten, Wolfe et al. 2007), an interview and medical-record-abstraction project, to obtain information on the following questions: Are patients receiving care and treatment in accordance with United States Public Health Service (USPHS) guidelines? Are patients receiving care in Ryan White-funded facilities receiving the same quality of care as patients in private facilities? What are the barriers to receiving care and services? Included is a locally and nationally representative sample of HIV-positive adults in care with assessments of: adherence; sexual behavior; drug use; care-seeking; clinical outcomes; treatment; CD4 cell count and VL; opportunistic illnesses; type and quality of care received; and met and unmet needs for HIV care and prevention services. This information is limited in scope for the purposes of this study (funded locations, sites included, etc). However, an assessment will be made to determine the information that can be gained from MMP.
7.2.1 Human Subjects/Ethical Considerations

HIV case surveillance data are collected as part of routine HIV surveillance as mandated by state or local laws or regulations. Similarly, other data are routinely collected by DOHs.

According to CDC’s Guidelines for Defining Public Health Research and Public Health Non-Research (CDC 1999) and Title 45 Part 46 of the Code of Federal Regulations (DHHS 2005), the CDC has determined that HIV surveillance is not a research activity, and, therefore, does not require review by an IRB.

State and local HIV surveillance programs must comply with federal security and confidentiality guidelines for collecting, storing and releasing data (CDC 2006).

Data are sent to the CDC without personal identifiers.

7.3 Behavioral Data

Information on population HIV-testing rates is generally available for the United States as a whole and for select communities that have implemented behavioral surveys that are representative for their community. The primary measures are the proportion of persons who ever tested for HIV and the proportion of persons who tested for HIV within the past year. This information is critical to monitor trends of HIV testing in the future.

One behavioral survey conducted in all states and Washington, D.C. is the Behavioral Risk Factor Surveillance System (BRFSS) funded by CDC (CDC, http://www.cdc.gov/BRFSS/). BRFSS is an on-going, telephone, health survey system, tracking health conditions and risk behaviors in the United States yearly since 1984. Currently, data are collected monthly in all 50 states, Washington, D.C., Puerto Rico, the United States Virgin Islands and Guam. State and local areas may add questions to the standard questionnaire. However, oversampling of local areas is needed to make inferences for areas smaller than statewide, and representativeness/response rates have diminished with the widespread use of cell phones. Surveys conducted in the local areas are described in Table 5. One limitation of BRFSS surveys is the diminishing population reached using land-lines; therefore, any BRFSS-type survey used for this project should include cell phones.

Our approach for this study will be to support currently planned surveys to obtain representative HIV testing data from the communities.

Accuracy of recall of HIV testing will not be assessed as part of this study.

This study also includes a special emphasis population, MSM. The National HIV Behavioral Surveillance (NHBS) system is a CDC-funded project conducted in 25 cities in the United States, including Washington, D.C. The purpose of this serial cross-sectional study is to yield information about what people do that puts them at risk for HIV. NHBS has three cycles focusing on different risk groups: NHBS-MSM, injecting drug users (NHBS-IDU), and heterosexuals at risk of HIV infection (NHBS-HET).
Areas conducting NHBS will be able to obtain HIV testing data among MSM in the MSM cycle years.

7.3.1 Behavioral Data Collection in Washington, D.C.

Washington, D.C. implements HIV testing questions through the BRFSS and will include cell phones in 2009. Washington, D.C. also collects behavioral data via NHBS.

7.3.2 Behavioral Data Collection in New York City

NYC conducts the NYC Community Health Survey (CHS), a telephone survey conducted annually. CHS provides robust data on the health of New Yorkers, including neighborhood, borough and citywide estimates on a broad range of chronic diseases and behavioral risk factors.

The CHS is based on the BRFSS. The CHS is a cross-sectional survey that samples approximately 10,000 adults aged 18 and older from all five boroughs of NYC—Manhattan, Brooklyn, Queens, Bronx and Staten Island.

A computer-assisted telephone interviewing (CATI) system is used to collect survey data from respondents accessed by random-digit dialing of household-based land lines.

Interviews are conducted in a variety of different languages.

All data collected are self-report.

Questions on HIV testing are included. NYC also collects behavioral data via NHBS.

7.3.3 Behavioral Data Collection in Miami

In Miami, behavioral data is collected via NHBS and BRFSS. BRFSS collected county data in 2007, but not in 2008 and 2009. County data will be collected again in 2010. However, representativeness for Miami and inclusion of cell phones need to be determined.

7.3.4 Behavioral Data Collection in Philadelphia

In Philadelphia, behavioral data is collected via NHBS and BRFSS/Southeastern Pennsylvania Household Health Survey (PHHS).

Public Health Management Corporation’s (PHMC) Southeastern Pennsylvania Household Health Surveys are extensive health surveys that provide timely information on more than 13,000 residents, both children and adults, living in Bucks, Chester, Delaware, Montgomery and Philadelphia counties. The survey targets key information about health status, personal health behaviors, and access to and utilization of area health services. These data are available at the census tract, ZIP code, county and regional level. The Household Health Survey provides primary data on a broad range of health topics such as health status, access to care, utilization of services, personal health behaviors, health screening information, health insurance status, women's health, child health, and older adult health and social support needs. The survey asks whether persons
have had an HIV test. The survey includes responses from more than 10,000 households in the region, representing more than 13,000 adults and children.

Interviews are conducted by telephone using a random-digit dial methodology. Adult and child respondents are selected randomly using the "last birthday" method.

7.3.5 Behavioral Data Collection in Chicago

In Chicago, behavioral data is collected via NHBS. The state conducts BRFSS and data is available for Chicago. However, representativeness for Chicago and inclusion of cell phones need to be reviewed.

7.3.6 Behavioral Data Collection in Houston

In Houston, behavioral data is collected via NHBS. The state conducts BRFSS. However, representativeness for Houston and inclusion of cell phones need to be determined.

8.0 ADMINISTRATIVE PROCEDURES AND OPERATIONAL CONSIDERATIONS

This study differs from other HPTN studies to date. In fact, it differs from most other studies conducted through HIV clinical trials networks. It is an ambitious effort that aims at identifying a strategy to tackle the entrenched HIV epidemic in some communities in the United States. It differs from traditional clinical trials in the following manners:

- The study will be a collaborative effort among the HPTN, the CDC, and the DOHs in the intervention and non-intervention communities.

- Many study outcomes will be based on data collected through surveillance systems in the public health programs of local health departments in the intervention and non-intervention communities.

- The study will include two site randomizations (one of HIV test sites and one of HIV care sites) as well as individual randomization in the Prevention for Positives cohort.

- The laboratory outcomes in this study will be based on standardized assays conducted in certified laboratories in the United States. No laboratory testing will be conducted at the HPTN Network Laboratory (NL).

The SDMC at SCHARP will assist in the design of the study, determine the sampling plan and coordinate the data collection from various sources. The SDMC will play the key role in communicating with staff responsible for surveillance in the jurisdictions where the study will be conducted. The SDMC will also lead data analysis efforts and generation of data reports.

The NL will assist in the design of the study as well as in guiding the conceptualization and selection of appropriate laboratory objectives and endpoints. The NL will provide expert input into the methodology for various assays, as needed. However, no assays will
be conducted at NL in the main part of the study. All laboratory data will be collected through surveillance systems or medical records abstraction (for the Prevention for Positives study component).

The HPTN CORE at Family Health International (FHI) will coordinate the development of the protocol and facilitate the implementation of the study in the intervention communities.

8.1 Study Activation

For this study, traditional DAIDS site activation and protocol registration procedures will not be conducted for the following study components: Expanded HIV Testing, Linkage-to-Care, Viral Suppression, and Provider Surveys. The study team will define a start date for the Expanded Testing component. The HPTN CORE will notify all participating HIV test sites when they may begin to distribute coupons to patients for the Linkage-to-Care study component and when participating HIV care sites may begin to distribute FIs for the Viral Suppression component.

Protocol registration and modified DAIDS site activation will be conducted for the following study components: Prevention for Positives and Patient Surveys. Traditional participant informed consent will be obtained for the Prevention for Positives component and for the Patient Survey. Operationally, the Prevention for Positives intervention and the Patient Survey will be administered together during Month 0 and Month 12 of the Prevention for Positives study component. Therefore, a single comprehensive informed consent form was designed (see Appendix IIA). Following ethical review and approval, study sites will submit required administrative documentation — as listed in the study-specific procedures manual — to the HPTN CORE. CORE staff will work with study site staff and complete DAIDS protocol registration in accordance with the current DAIDS Protocol Registration Policy and Procedure Manual. Included in this step will be CORE review of each site-specific study informed consent form. Site-specific informed consent forms (ICFs) WILL NOT be reviewed and approved by the DAIDS PRO and sites will receive an Initial Registration Notification from the DAIDS PRO that indicates successful completion of the protocol registration process. A copy of the Initial Registration Notification should be retained in the site's regulatory files. Pending successful protocol registration and submission of all required documents, CORE staff will “activate” the site to begin study operations. Study implementation may not be initiated until a study activation notice is provided to the site.

The HPTN CORE will also notify SCHARP and other network partners of dates that study components will start.

8.2 Study Coordination

Study implementation will be directed by this protocol as well as the SSP manual. The SSP manual will outline procedures for conducting study visits, data collection and processing, management and reporting, and other study operations.
Data for this study will be collected in three ways:

- The DOHs in Washington, D.C. and in the Bronx of NYC will provide aggregate data from their routine surveillance systems to the data management group for the study, SCHARP. SCHARP will conduct all data analyses.
- Data will also be collected from patients and providers using study-specific surveys. These data will also be submitted to SCHARP for analysis.
- For the Prevention forPositives study component, data will be collected by patient self-report and medical records abstraction. These data will be submitted to SCHARP.

Close coordination between protocol team members will be necessary to track study progress, respond to questions about proper study implementation and address other issues in a timely manner. Rates of accrual, adherence and follow-up will be monitored closely by the team as well as an HPTN study-monitoring committee (SMC) for the Prevention for Positives and survey components of the study.

8.3 Study Monitoring

This study will not undergo traditional HPTN monitoring by PPD Research Associates. However, the study will be carefully reviewed by an HPTN SMC and/or external monitoring group on a regular basis. Study elements that will be monitored include data quality, achievement of milestones, and magnitude of utilization of study test and care sites, as well as migration of patients between test and care sites based on their randomization assignments.

8.4 Protocol Compliance

The study will be conducted in full compliance with the protocol. The protocol will not be amended without prior written approval by the Protocol Co-Chairs and NIAID Medical Officer. All protocol amendments must be submitted to and approved by the relevant IRB(s) prior to implementing the amendment. Upon receiving final IRB approval for all protocol amendments, including Letter of Amendments [LoAs] and full protocol amendments, sites should implement the amendment immediately.

For the Prevention for Positives and Patient Survey study components, sites are required to submit an amendment (LoA or full amendment) registration packet to the DAIDS PRO at the RSC. HPTN CORE staff will work with study site staff to complete this procedure. Site-specific ICF(s) WILL NOT be reviewed and approved by the DAIDS PRO and sites will receive an LoA or Amendment Registration Notification from the DAIDS PRO that indicates successful completion of the amendment protocol registration process. A LoA or Amendment Registration Notification from the DAIDS PRO is not required prior to implementing the LoA or full amendment. A copy of the final LoA or Amendment Registration Notification issued by the DAIDS PRO should be retained in the site's regulatory files.
For additional information on the protocol registration process and specific documents required for initial and amendment registrations, refer to the current version of the DAIDS Protocol Registration Manual.

8.5 Human Subjects/Ethical Considerations

Prior to implementation of this protocol, and any subsequent full version amendments, each site must have the protocol and the protocol consent form(s) approved (for Prevention for Positives study component only), as appropriate, by their local institutional review board (IRB).

Expanded HIV testing. This component is public health practice. Social mobilization and emergency department testing is already taking place in the two intervention communities (Bronx and Washington DC) and is intended to be specific to the needs of those populations. While some observations from the overall TLC Plus project with regard to social mobilization and expanded testing may be applicable elsewhere in the United States, these activities were initiated originally for the specific benefit of the respective communities, and do not constitute research.

Linkage-to-Care. This component of the study involves a randomized intervention using financial incentives (FIs) to link HIV-positive individuals to HIV care after diagnosis. This component is public health research. In this component, HIV test sites will be randomized, not individuals, and no de novo individual-level data will be collected. Instead, (de-identified) HIV surveillance data routinely collected by the Departments of Health (DOH) in the two communities will be analyzed. Because this research involves minimal risk and would be impracticable with informed consent, a waiver of patient informed consent will be requested under 45 CFR 46.116 (c) or (d).

Viral Suppression. This component of the study involves an intervention using FIs to encourage HIV-positive individuals on ART to be adherent to their medications and to maintain HIV VL suppression. This component is public health research. In this component, HIV care sites will be randomized, not individuals, and no de novo individual-level data will be collected. Instead, (de-identified) HIV surveillance data routinely collected by the DOH in the two communities will be analyzed. Because this research involves minimal risk and would be impracticable with informed consent, a waiver of patient informed consent will be requested under 45 CFR 46.116 (c) or (d).

Prevention for Positives. In this component of the study, a subset of patients enrolled at select HIV care sites in the two intervention communities will be randomized either to an intervention arm (receiving computer-delivered intervention for sexual risk reduction plus SOC prevention activities) or to the control arm (receiving only the SOC prevention activities at the care site). Individual-level data will be collected and analyzed. This component is public health research requiring informed consent. Written informed consent will be obtained from participants.

Patient survey. All patients participating in the Prevention for Positives study component will be administered the Patient Survey. Individual-level data will be analyzed. This component is public health research. Written informed consent will be obtained from
participants at the time of consenting for participation in the Prevention for Positives study component.

**Provider survey.** Providers from participating HIV care sites will be invited to complete a Web-based survey regarding their knowledge, attitudes and practices concerning ART for treatment, frequency of HIV testing, and limited sociodemographic information. The survey will be anonymous, and will collect no identifying information. This component is public health research. Informed consent will be obtained from the provider’s completing the survey. Before the survey will display, providers will read the consent and if they agree to participate, will indicate their agreement by clicking a button labeled “I agree.”

**Investigator's Records**

Study-specific records will only be located at HIV care sites participating in the Prevention for Positives study component. At these sites, all study source documents (such as informed consent forms and patient ID linkage logs) will be maintained in a locked cabinet or room. Access to these records will be restricted to appropriate study staff members, NIAID, OHRP, government or regulatory authorities, and/or IRB.

### 8.6 Use of Information and Publications

Publication of the results of this study will be governed by HPTN policies. Any presentation, abstract or manuscript will be submitted to the HPTN Manuscript Review Committee for review prior to submission. Similar review may be necessary by other collaborating organizations.

### 8.7 Study Discontinuation

The study may be discontinued at any time by NIAID, the HPTN, the Office for Human Research Protections (OHRP), government or regulatory authorities, and/or an IRB. However, should the study be stopped early, all sites and community partners would be notified of such a decision, as well as the reasons behind it, prior to any termination activities.
REFERENCES


Royal, S. W., D. P. Kidder, et al. (2009). "Factors associated with adherence to highly active antiretroviral therapy in homeless or unstably housed adults living with HIV." AIDS Care 21(4): 448-55.


APPENDICES
Appendix I: Schedule of Study Visits and Procedures
### Schedule of Study Visits and Procedures
Prevention for Positives Computerized Patient Intervention and Survey

<table>
<thead>
<tr>
<th>Administrative and Regulatory Procedures</th>
<th>HIV Care Site Visit #1 Enrollment</th>
<th>HIV Care Site Visit #2-Visit #5 (Months 3, 6, 9 &amp; 12)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-screening</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Administer Informed Consent</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Confirm Eligibility</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Collect Locator Information</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Administer CARE+/ACASI</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Administer Survey</td>
<td>X</td>
<td>X (Month 12 only)</td>
</tr>
<tr>
<td>Provide Compensation</td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>
Appendix IIA: Patient Computer-Delivered Intervention and Survey Informed Consent Form
Computera-delivered Intervention and Survey Subject Information and Consent Form and Authorization to Use and Disclose Personal Health Information for Research:

Title of the Research Study: TLC-Plus: A Study to Evaluate the Feasibility of an Enhanced Test, Link-to-Care, Plus Treat Approach for HIV Prevention in the United States

Protocol #: HPTN 065, Version 3.0, 14 January 2014
DAIDS ID: 11685

Sponsor: National Institute of Allergy and Infectious Diseases (NIAID), National Institute on Drug Abuse (NIDA), National Institute of Mental Health (NIMH), National Institutes of Health (NIH)

Investigator of Record: (insert name)

Research Site Address(es) (insert address)

Daytime Telephone Number: (insert number)

24-hour Contact Number: (insert number)

Purpose of the Subject Information and Consent Form

This Subject Information and Consent Form may contain words you do not understand. Please ask the study investigator or the study staff to explain any words or procedures that you do not clearly understand.

The purpose of this form is to give you information about the research study and, if signed, will give your permission to take part in the study. The form describes the purpose, procedures, benefits, risks, discomforts and precautions of the research study. You should take part in the study only if you want to do so. You may refuse to take part or withdraw from this study at any time without penalty or loss of benefits to which you are otherwise entitled. Please read this Subject Information and Consent Form and ask as many questions as needed. You should not sign this form if you have any questions that have not been answered to your satisfaction.

Your study investigator will be paid by the sponsor to conduct this research.

Introduction

You have been asked to take part in a study that is testing a new computer program to help HIV-positive people. This research will use a computer to privately ask you questions. You will be
asked to use the computer a total of 5 times. You will use the computer every three months when you come in for your clinic visits for twelve months.

Around 1320 people who are HIV-positive in Washington, D.C. and in the Bronx will participate in the study. You do not have to know how to use a computer or be able to read to be in this study.

What will happen during this study?

If you agree to take part in this study, you will first sign this Subject Information and Consent Form before any study-related procedures are performed.

We will use a computer to talk about what is going on for you with HIV. During your first and 12-month visit, we will also ask you questions about your knowledge of HIV, your knowledge of medications for treating HIV and your feelings about medical care for people with HIV.

You will be asked to answer all of the questions openly and honestly, but you may refuse to answer any of the questions or stop at any time if you feel uncomfortable. You will also be provided with contact and referral information if any of the questions raises issues that you would like to talk about further, at this or some later time.

Computer sessions are anticipated to take you approximately 15 - 40 minutes to complete.

For your time and effort, we will reimburse you $10 per visit. There is no cost to you to participate in this part of the study.

Subjects who choose to join the study will be randomly assigned to a study group. There are only two groups:

- One study group will be asked questions by the computer program.
- The second study group will be asked the same questions and will also be shown some videos. The videos will be short and will include HIV risk reduction topics. After the videos are shown, the computer then will help people create a health plan.

You will have a 50% chance of being in the group that is asked questions by the computer program. You will also have a 50% chance of being in the group that is asked questions and shown videos. The group assignment will be made randomly by the computer program. Staff at the site where you get your HIV care cannot assign you to a group and the staff will not know which group to which you are assigned.

Computer sessions for both study groups are anonymous. No names or identifying information will be attached to the computer. For both study groups, we would like permission to access medical records at the HIV clinic. We will use this information to evaluate HIV Viral Load, CD4 cell count, and other information relevant to your health, at each computer session. We would only like to access your information for as long as you are enrolled in the study. Once you have completed the study we will no longer access your medical records at any HIV clinic. We will not share information with the study clinic staff about your answers to the questions on the computer. If you decide to allow us to access your health information for the study, you will
need to sign an authorization form at the end of this consent form document giving permission to let us see your records.

For the study group that is asked questions and shown videos, an anonymous health plan will print out at the end of the computer session. If you are assigned to this group, you can decide whether to share the health plan print out with your provider. You do not have to show your provider the print out.

For both groups there are questions about depression, suicide and domestic violence. If your answers to these questions show that you may be depressed, suicidal or are currently in an abusive relationship, a health worker here at the clinic will follow-up with you to offer support and/or referral but will not know what you may be having trouble with. For example, the healthcare worker will not know whether you indicated that you are suicidal or whether you indicated that you are in an abusive relationship. We will not share your actual answers with study staff at the clinic. You can decide what information you want to share with the healthcare worker.

**What are the possible risks or discomforts?**

It is possible that answering the questions on the computer may make you embarrassed or upset. You may refuse to answer any of the questions or stop answering at any time. The greatest risk may involve your privacy. The steps that the study team has taken to protect your privacy are described in this form.

**What are the potential benefits?**

There may be no direct benefits to you. We hope the information we collect will help us find better ways to provide HIV care in your community. You may feel a benefit from sharing your experiences with someone who is interested in your opinions.

**Are there any alternatives to participation?**

The study coordinator will explain other programs at this site that can help HIV-positive people change their behavior so that they do not pass HIV on to others.

**How will my confidentiality and privacy be protected?**

We cannot guarantee absolute confidentiality. However, we will do everything possible to protect your confidentiality if you join this study.

To protect your privacy, you will meet with a healthcare provider in a private area where others cannot overhear conversations with you. While you are participating in the computerized session, you will be given headphones and a place to sit where no one can look over your shoulder to see what you are doing.

Every effort will be made to keep your personal information confidential. Your personal information (name, address, phone number) will be protected by the research clinic. This information will not be used in any publication of information about this study.
In addition to the efforts made by the study staff to help keep your personal information confidential, we have obtained a Certificate of Confidentiality from the U.S. Federal Government. This certificate protects researchers from being forced to tell people who are not connected with this study, such as the court system, about your participation. Any publication of this study will not use your name or identify you personally.

People who may review your records include: the U.S. Food and Drug Administration (FDA), [insert name of site IRB], National Institutes of Health (NIH), study staff, study monitors, and drug companies supporting this study. Also, the Certificate of Confidentiality does not prevent you from releasing information about yourself and your participation in the study. The eCertificate cannot be used to resist a demand for information from personnel of the United States Government that is used for auditing or evaluation of Federally funded projects or for information that must be disclosed in order to meet the requirements of the federal Food and Drug Administration (FDA).

Even with the Certificate of Confidentiality, if the study staff learns of possible child abuse and/or neglect or a risk of harm to you or others, we will tell the proper authorities.

What happens if I am injured by participating in this study?

Because this study only involves answering questions, reading messages and viewing videos, it is very unlikely that you could be injured. However, if you are injured as a result of joining this study, you will be given immediate treatment for your injuries. You may have to pay for this care. There is no program for compensation either through this institution or the United States NIH.

What are my legal rights?

The above section does not restrict your right to seek legal assistance. You will not be giving up any of your legal rights by signing this Subject Information and Consent Form.

Your participation is voluntary.

You are not required to join this study. You do not have to participate in the computer sessions for us. If you decide to participate, you may refuse to answer any of the questions or stop at any time without reducing or affecting any care that you receive at this site. If you do decide to leave the study we will ask you to complete one final computer session, however, you will not be required to do this.

What are some reasons why I may be withdrawn from this activity without my consent?

You may be withdrawn from the study without your consent for the following reasons:

- The research study, or this part of the study, is stopped or canceled.
- The study staff feels that completing the study or this part of the study would be harmful to you or others.

Persons to Contact for Problems or Questions
If you have any questions about your participation in this research study, your rights as a research subject, or if you feel that you have experienced a research-related injury, contact:

- Investigator of Record Name: *(site insert name of the investigator or other study staff)*
- Research Site Address: *(site insert physical address of above)*
- Daytime Telephone Number(s): *(site insert telephone number)*
- 24-hour contact number(s): *(site insert telephone number)*

If you have any questions or concerns about your rights as a research subject or want to discuss a problem, get information or offer input, you may contact: Institutional Review Board: *(site insert name or title of person on the IRB or other organization appropriate for the site)*
- Address of Institutional Review Board: *(site insert physical address of above)*
- Daytime Telephone Number(s): *(site insert telephone number of above)*
SUBJECT’S STATEMENT OF CONSENT

TLC-Plus: A Study to Evaluate the Feasibility of an Enhanced Test, Link to Care, Plus Treat Approach for HIV Prevention in the United States

- I have been given sufficient opportunity to consider whether to participate in this study.
- My taking part in this research study is voluntary. I may decide not to take part or to withdraw from the research study at any time without penalty or loss of benefits or treatments to which I am entitled.
- The research study may be stopped at any time without my consent.
- I have had an opportunity to ask my study investigator questions about this research study. My questions so far have been answered to my satisfaction.
- I have been told how long I may be in the research study.
- I have been informed of the procedures and tests that may be performed during the research study.
- I have been told what the possible risks and benefits are from taking part in this research study. I may not benefit if I take part in this research study.
- I do not give up my legal rights by signing this form.
- I have been told that prior to any study related procedures being performed, I will be asked to voluntarily sign this Study Information and Consent Form.
- I will receive a signed and dated copy of this Subject Information and Consent Form.

If you have either read or have heard the information in this Subject Information and Consent Form, if all of your questions have been answered, and if you agree to take part in the computer assisted interview and subject survey questionnaire, please sign and print your name on the line below.

I voluntarily agree to take part in this research study.

Subject’s Name (print)   Subject’s Signature and Date

I certify that the information provided was given in a language that was understandable to the subject.

Study Staff Conducting Consent Discussion (print)   Study Staff Signature and Date

Witness’ Name (print)   Witness’ Signature and Date
(As appropriate)
Authorization to Use and Disclose Personal Health Information for Research

The United States government has issued a privacy rule to protect the privacy rights of patients. This rule was issued under a law called the Health Insurance Portability and Accountability Act of 1996 (HIPAA). The Privacy Rule is designed to protect the confidentiality of your personal health information. The document you are reading, called an “Authorization,” describes your rights and explains how your health information will be used and disclosed (shared).

In working with the sponsor, the study investigator, (insert the name of site’s study investigator), will use and share personal health information about you. This is information about your health that includes information in your medical record and information created or collected during the study. This information may include laboratory test results. Some of these tests may have been done as part of your regular care. The study investigator will use this information about you to complete this research.

The study investigator will assign a code number to your information that is shared with the sponsor. The sponsor and its representatives may review or copy your personal health information at the study site. Your IRB, (insert name of your IRB), may also review or copy your information to make sure that the study is done properly or for other purposes required by law.

By signing this Authorization, you allow the study investigator to use your personal health information to carry out and evaluate this study. You also allow the study investigator to share your personal health information with:

- the sponsor and its representatives
- (insert name of site’s IRB)

Your personal health information may be further shared by the groups above. If shared by them, the information will no longer be covered by the Privacy Rule. However, these groups are committed to keeping your personal health information confidential.

You have the right to see and get a copy of your records related to the study for as long as the study investigator has this information. However, by signing this Authorization you agree that you might not be able to review or receive some of your records related to the study until after the study has been completed.

You may choose to withdraw this Authorization at any time, but you must notify the study investigator in writing. Send your written withdrawal notice to [insert study investigator’s name & address].

If you withdraw from the study and withdraw your Authorization, no new information will be collected for study purposes unless the information concerns a social harm (a bad effect) related to the study. If a social harm occurs, your entire medical record may be reviewed. All information that has already been collected for study purposes, and any new information about a social harm related to the study, will be sent to the study sponsor.

If you withdraw from the study but do not withdraw your Authorization, new personal health information may be collected until this study ends.
This Authorization does not have an expiration date. If you do not withdraw this Authorization in writing, it will remain in effect indefinitely. Your study investigator will keep this Authorization for at least 6 years.

If you do not sign this Authorization, you cannot participate in this research study. If you withdraw this Authorization in the future, you will no longer be able to participate in this study. Your decision to withdraw your Authorization or not to participate will not involve any penalty or loss of access to treatment or other benefits to which you are entitled.

AUTHORIZATION

I authorize the release of personal health information from my medical records related to this study to the sponsor and its representatives, and (insert name of site’s IRB), as described above. I have been told that I will receive a signed and dated copy of this Authorization for my records.

__________________________________________
Printed Name of Subject

__________________________________________ Date
Signature of Subject

__________________________________________
Printed Name of Person Obtaining Authorization

__________________________________________ Date
Signature of Person Obtaining Authorization
Appendix IIB: Provider Survey Online Informed Consent Text
The NIH-funded HIV Prevention Trials Network (HPTN) is conducting a study to assess the feasibility of increasing HIV testing and facilitating linkage to care for HIV-infected-positive patients with the eventual goal of reducing incident HIV infection in the U.S. For the study to successfully impact HIV testing and treatment, the study team first needs to understand current practices of front-line providers.

You are invited to participate in a brief survey. This usually takes about *XXX* minutes to complete. You were selected because of your experience and geographic location. The questionnaire includes questions about your knowledge of HIV, your knowledge of medications for treating HIV, and your feelings/opinions about medical care for people with HIV. No personally identifying information will be collected about you, only some basic demographics. We hope that you will feel comfortable answering all of the questions openly and honestly, but you may refuse to answer any of the questions or stop completing the questionnaire, at any time. For your time and effort in completing this survey, we are providing you with a $50 Amazon.com gift certificate. You may access the certificate by printing the web coupon attached to this survey.

If you have read the information in this consent form, and if you agree to take part in the questionnaire, please CLICK on the link below. This will constitute your informed consent to participate in this survey.

I CONSENT
Appendix IIIA: Adult HIV/AIDS Confidential Case Report

(Currently under revision, Office of Management and Budget expiration February 2010)
ADULT HIV/AIDS CONFIDENTIAL CASE REPORT
(Patients ≥13 years of age at time of diagnosis)

II. HEALTH DEPARTMENT USE ONLY

<table>
<thead>
<tr>
<th>REPORT STATUS</th>
<th>Report</th>
<th>Update</th>
</tr>
</thead>
<tbody>
<tr>
<td>REPORTING HEALTH DEPARTMENT</td>
<td>State:</td>
<td>City/County:</td>
</tr>
<tr>
<td>DATE FROM COMPLETED</td>
<td>Mo. Day Yr:</td>
<td></td>
</tr>
<tr>
<td>REPORT SOURCE</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

III. DEMOGRAPHIC INFORMATION

<table>
<thead>
<tr>
<th>HIV Infection (not AIDS)</th>
<th>AIDS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at Diagnosis:</td>
<td>yrs</td>
</tr>
<tr>
<td>Date of Birth:</td>
<td></td>
</tr>
<tr>
<td>Current Status: Alive</td>
<td>Dead</td>
</tr>
<tr>
<td>Date of Death:</td>
<td></td>
</tr>
<tr>
<td>State/Territory of Death:</td>
<td></td>
</tr>
</tbody>
</table>

SEX:
- Male
- Hispanic
- Not Hispanic or Latine

ETHNICITY:
- Select one

RACE:
- Select one or more
- American Indian
- Alaska Native
- Asian
- Native Hawaiian or Other Pacific Islander
- White
- Unk.

RESIDENCE AT DIAGNOSIS:
- City: |
- State:
- Country: |
- Zip Code: |

IV. FACILITY OF DIAGNOSIS

Facility Name: |
City: |
State/Country: |

FACILITY SETTING:
- Public
- Private
- Federal
- Unk.

FACILITY TYPE:
- Physician, HMO
- Hospital, Inpatient
- Other (specify): |

V. PATIENT HISTORY

AFTER 1977 AND PRECEDING THE FIRST POSITIVE HIV ANTIBODY TEST OR AIDS DIAGNOSIS, THIS PATIENT HAD
<table>
<thead>
<tr>
<th>Respond to ALL Categories</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
</tr>
<tr>
<td>No</td>
</tr>
<tr>
<td>Unk.</td>
</tr>
</tbody>
</table>

- Sex with male |
- Sex with female |
- Injected nonprescription drugs |
- Received clotting factor for hemophilia/coagulation disorder |
- Factor VIII |
- Factor IX |
- Other (specify): |
- HETEROSEXUAL relations with any of the following: |
- Intravenous/injection drug user |
- Bisexual male |
- Person with hemophilia/coagulation disorder |
- Transfusion recipient with documented HIV infection |
- Transplant recipient with documented HIV infection |
- Person with AIDS or documented HIV infection, risk not specified |
- Received transfusion of blood/blood components (other than clotting factor) |
- Received transfusion of blood/blood components (other than clotting factor) |
- Worked in a health-care or clinical laboratory setting |
- Worked in a health-care or clinical laboratory setting |
- Worked in a health-care or clinical laboratory setting |
- Worked in a health-care or clinical laboratory setting |
- Worked in a health-care or clinical laboratory setting |

VI. LABORATORY DATA

1. HIV ANTIBODY TESTS AT DIAGNOSIS:
<table>
<thead>
<tr>
<th>(Indicate first test)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HIV–1 ELISA</td>
</tr>
<tr>
<td>HIV–1/HIV–2 combination ELISA</td>
</tr>
<tr>
<td>HIV–1 Western blot/IFA</td>
</tr>
<tr>
<td>Other HIV antibody test (specify):</td>
</tr>
</tbody>
</table>

2. POSITIVE HIV DETECTION TEST: (Record earliest test) |
   - culture |
   - antigen |
   - PCR, DNA or RNA probe |
   - Other (specify): |

3. DETECTABLE VIRAL LOAD TEST: (Record most recent test) |
   - Test type (specify): |
   - Dupe/SmL |
   - Other (specify): |

4. IMMUNOLOGIC LAB TESTS:
<table>
<thead>
<tr>
<th>AT OR CLOSEST TO CURRENT DIAGNOSTIC STATUS</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD4 Count</td>
</tr>
<tr>
<td>CD4 Percent</td>
</tr>
<tr>
<td>First &lt;200 µL or &lt;14%</td>
</tr>
<tr>
<td>CD4 Count</td>
</tr>
<tr>
<td>CD4 Percent</td>
</tr>
</tbody>
</table>

CD 50.42A REV. 02/2007 (Page 1 of 2) — ADULT HIV/AIDS CONFIDENTIAL CASE REPORT —

14 January 2014
### VIII. CLINICAL STATUS

<table>
<thead>
<tr>
<th>AIDS INDICATOR DISEASES</th>
<th>Initial Diagnosis</th>
<th>Initial Date</th>
<th>AIDS INDICATOR DISEASES</th>
<th>Initial Diagnosis</th>
<th>Initial Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>C. bronchi, trachea, or lungs</td>
<td>1 NA</td>
<td></td>
<td>Lymphoma, Burkitt’s (or equivalent term)</td>
<td>1 NA</td>
<td></td>
</tr>
<tr>
<td>C. esophagus</td>
<td>1 2</td>
<td></td>
<td>Lymphoma, immunoblastic (or equivalent term)</td>
<td>1 NA</td>
<td></td>
</tr>
<tr>
<td>C. invasive cervical</td>
<td>1 NA</td>
<td></td>
<td>Lymphoma, primary in brain</td>
<td>1 NA</td>
<td></td>
</tr>
<tr>
<td>C. disseminated or extrapulmonary</td>
<td>1 NA</td>
<td></td>
<td>Mycobacterium avium complex or M. kansasii, disseminated or extrapulmonary</td>
<td>1 2</td>
<td></td>
</tr>
<tr>
<td>C. extra pulmonary</td>
<td>1 NA</td>
<td></td>
<td>M. tuberculosis, pulmonary*</td>
<td>1 2</td>
<td></td>
</tr>
<tr>
<td>C. chronic intestinal (&gt;1 mo. duration)</td>
<td>1 NA</td>
<td></td>
<td>M. tuberculosis, disseminated or extrapulmonary*</td>
<td>1 2</td>
<td></td>
</tr>
<tr>
<td>C. other than liver, spleen, or nodes</td>
<td>1 NA</td>
<td></td>
<td>Mycobacterium, other species or unidentified species, disseminated or extrapulmonary</td>
<td>1 2</td>
<td></td>
</tr>
<tr>
<td>C. reitinitis with loss of vision</td>
<td>1 2</td>
<td></td>
<td>Pneumocystis carinii pneumonia</td>
<td>1 2</td>
<td></td>
</tr>
<tr>
<td>HIV encephalopathy</td>
<td>1 NA</td>
<td></td>
<td>Pneumonia, recurrent, in 12 mo. period</td>
<td>1 2</td>
<td></td>
</tr>
<tr>
<td>Herpes simplex: chronic ulcer(s) (&gt;1 mo. duration) or blisters, pneumonia or sores</td>
<td>1 NA</td>
<td></td>
<td>Progressive multifocal leukoencephalopathy</td>
<td>1 NA</td>
<td></td>
</tr>
<tr>
<td>H. disseminated or extrapulmonary</td>
<td>1 NA</td>
<td></td>
<td>Salmonella septicemia, recurrent</td>
<td>1 NA</td>
<td></td>
</tr>
<tr>
<td>I. chronic intestinal (&gt;1 mo. duration)</td>
<td>1 NA</td>
<td></td>
<td>Toxoplasmosis of brain</td>
<td>1 2</td>
<td></td>
</tr>
<tr>
<td>Kaposi’s sarcoma</td>
<td>1 2</td>
<td></td>
<td>Wasting syndrome due to HIV</td>
<td>1 NA</td>
<td></td>
</tr>
</tbody>
</table>

Det. = definitive diagnosis  Pres. = presumptive diagnosis  * RVCT CASE NO.:  

### IX. TREATMENT/SERVICES REFERRALS

Has this patient been informed of his/her HIV infection? 1 Yes 0 No 9 Unk.

This patient’s partners will be notified about their HIV exposure and counseled by:

1 Health department 2 Physician/provider 3 Patient 9 Unknown

If this patient received or is receiving:

- Anti-retroviral therapy 1 Yes 0 No 9 Unknown
- PCP prophylaxis 1 Yes 0 No 9 Unknown

This patient has been enrolled at:

1 Clinical Trial 2 NIH-sponsored 3 Other-sponsored 9 Unknown

This patient’s medical treatment is primarily reimbursed by:

1 Medicaid 2 Private insurance/HMO 3 No coverage 4 Other Public Funding 7 Clinical trial/ government program 9 Unknown

### FOR WOMEN:

- This patient is receiving or has been referred for gynecological or obstetrical services: 1 Yes 0 No 9 Unknown
- Is this patient currently pregnant? 1 Yes 0 No 9 Unknown
- Has this patient delivered live-born infants? 1 Yes (if delivered after 1977, provide birth information below for the most recent birth) 0 No 9 Unknown

### CHILD’S DATE OF BIRTH:

Mo. Day Yr.  

Hospital of Birth: ________________________________  

City:  

State:  

Child’s Surname:  

Child’s State Patient No.:  

### X. COMMENTS:

Public reporting burden of this collection of information is estimated to average 70 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing this collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information, unless it displays a currently valid OMB control number. Consult with the agency having jurisdiction if you have any questions. Do not send the completed form to this address.
Appendix IIIB: Behavioral Risk Factor Surveillance System (BRFSS)

(Section 20: HIV/AIDS)
The next few questions are about the national health problem of HIV, the virus that causes AIDS. Please remember that your answers are strictly confidential and that you don’t have to answer every question if you do not want to. Although we will ask you about testing, we will not ask you about the results of any test you may have had.

20.1 Have you ever been tested for HIV? Do not count tests you may have had as part of a blood donation. Include testing fluid from your mouth.

1 Yes 7 Don’t know / Not sure [Go to Q20.5]
2 No [Go to Q20.4] 9 Refused [Go to Q20.5]

20.2 Not including blood donations, in what month and year was your last HIV test?

NOTE: If response is before January 1985, code “Don’t know.”

CATI INSTRUCTION: If the respondent remembers the year but cannot remember the month, code the first two digits 77 and the last four digits for the year.

_ _ / _ _ _ _
Code month and year
77/7777 Don’t know / Not sure
99/9999 Refused

20.3 Where did you have your last HIV test — at a private doctor or HMO office, at a counseling and testing site, at a hospital, at a clinic, in a jail or prison, at a drug treatment facility, at home, or somewhere else?

0 1 Private doctor or HMO office
0 2 Counseling and testing site
0 3 Hospital
0 4 Clinic
0 5 Jail or prison (or other correctional facility)
0 6 Drug treatment facility
0 7 At home
0 8 Somewhere else
7 7 Don’t know / Not sure
9 9 Refused

CATI note: Ask Q20.4; if Q20.2 = within last 12 months. Otherwise, go to Q20.5.

20.4 Was it a rapid test where you could get your results within a couple of hours?

1 Yes 7 Don’t know / Not sure
2 No 9 Refused

20.5 I’m going to read you a list. When I’m done, please tell me if any of the situations apply to you. You do not need to tell me which one.

- You have used intravenous drugs in the past year.
- You have been treated for a sexually transmitted or venereal disease in the past year.
- You have given or received money or drugs in exchange for sex in the past year.
- You had anal sex without a condom in the past year.

Do any of these situations apply to you?

1 Yes 7 Don’t know / Not sure
2 No 9 Refused
Appendix IIIC: 2008 PEMS Variable Requirements
PEMS Required Variables
Quick Reference Guide
February 8, 2008

This document provides a summary of the variable requirements for the January 1 and July 1, 2008 data collection periods, excluding variable requirements for HIV Testing and Partner Counseling and Referral Services (PCRS). HIV Testing variable requirements are currently specified in the HIV Testing Form and Variables Manual and the CDC HIV Testing Variables Data Dictionary. Requirements for PCRS will be released later in 2008. Since this document only provides a summary of the requirements, please refer to the PEMS DVS for a more detailed description of definitions and value choices.

<table>
<thead>
<tr>
<th>Variable Number</th>
<th>Variable Name</th>
<th>HD &amp; CBO Reported Required</th>
</tr>
</thead>
<tbody>
<tr>
<td>A01</td>
<td>Agency Name</td>
<td>Required</td>
</tr>
<tr>
<td>A01a</td>
<td>PEMS Agency ID</td>
<td>Required</td>
</tr>
<tr>
<td>A02</td>
<td>Community Plan Jurisdiction</td>
<td>Required</td>
</tr>
<tr>
<td>A03</td>
<td>Employer Identification Number (EIN)</td>
<td>Required</td>
</tr>
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<td>A04</td>
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**Intervention Plan Characteristics (Table F)**

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**Client Characteristics (Table G)**

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***Note: The recall period for client risk factors is 12 months.

**Note: Additional value choices for risk factors added:**
- Sex without using a condom
- Sharing drug injection equipment
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Referral (Table X7)

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Required Variables for Data Collection 2008  
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</tr>
<tr>
<td>HC21</td>
<td>Site Name/ID</td>
<td>Required</td>
</tr>
</tbody>
</table>

| **Community Planning Level (Table CP-A/B/C)** |                                                     |                            |
| CP-A01         | Name of HIV Prevention CPG                          | HD only                    |
| CP-A02         | Community Plan Year                                 | HD only                    |
| CP-B01         | Priority Population                                 | HD only                    |
| CP-B02         | Rank                                                | HD only                    |
| CP-B03         | Age                                                 | HD only                    |
| CP-B04         | Gender                                               | HD only                    |
| CP-B05         | Ethnicity                                            | HD only                    |
| CP-B06         | Race                                                 | HD only                    |
| CP-B07         | HIV Status                                           | HD only                    |
| CP-B08         | Geo Location                                         | HD only                    |
| CP-B09         | Transmission Risk                                   | HD only                    |
| CP-C01         | Name of the Prevention Activity/Intervention        | HD only                    |
| CP-C02         | Prevention Activity/Intervention Type               | HD only                    |
| CP-C04         | Evidence Based                                      | HD only                    |
| CP-C05         | CDC Recommended Guidelines                          | HD only                    |
| CP-C06         | Other Basis for Intervention                        | HD only                    |
| CP-C07         | Activity                                             | HD only                    |
Appendix IIID: NYC HIV Surveillance Performance Indicators
New York State requires named reporting of all diagnoses of HIV and AIDS, all HIV-related illness, all positive Western Blot (WB) tests for HIV antibody, all VL and CD4 lymphocyte values, and all HIV genotypes. The NYC HIV/AIDS Reporting System (HARS) is a population-based registry that since 1981 has been continuously updated with new, de-duplicated diagnoses and laboratory results. All incoming provider and laboratory reports that do not match an existing registry record initiate a field investigation with medical record review to confirm the case, date and disposition of diagnosis and collect all other data required for surveillance and partner notification. HARS also obtains data through regular matches with other disease registries, the NYC Death Registry, the National Death Index and the Social Security Death Master File. Because of its comprehensive nature, long history (AIDS reporting since 1981, HIV reporting since 2000) and location (AIDS epicenter, largest in the west), the system can reasonably be characterized as the largest longitudinal community HIV/AIDS database in the world. It is therefore an ideal source of population-level outcome data for communities mounting interventions to improve early diagnosis and uptake of care.

Because of the size, history, continuous feed of new information (>35,000 new laboratory results received per month), and reliance on outside entities (laboratories, medical records) of the surveillance system, control of its quality, completeness, accuracy and timeliness is an ongoing challenge. The program follows a set of monthly performance indicators to track data quality and staff and provider performance, and meets on a quarterly basis to review indicators, identify problem areas and take steps toward corrective action. The following points summarize the volume and periodicity of electronic laboratory reporting, the volume, timeliness and outcome of field investigations, the periodic registry matches, internal matching and deduplication procedures, and the interstate deduplication activities of the program.

**NYC HIV Core Surveillance Monthly Performance Indicators July 1, 2008-June 30, 2009**

**Electronic laboratory reporting from laboratory to NYS and from NYS to NYC in calendar year 2008:**

- Western Blot file received weekly (total N = 15,308)
- VL file received every two weeks (total N = 337,418)
- Low CD4 (<200) received every four weeks (total N = 95,479)
- High CD4 (200-499) received every 4-5 weeks (total N = 192,494)
- Very High CD4 (500+) received periodically (total N = 134,019)
- Total laboratory reports excluding genotype received 2008= 640,699
- Nucleotide sequences received monthly (total cumulative N 2008-2009= 71,074)

**Total number of laboratory reports received and processed July 1, 2008-June 30, 2009:** 453,791

- Mean 37,816 laboratory reports were received per month
- Mean 7,003 unique individuals were represented by these reports
- Mean 1,045 reports were potential new cases and initiated field investigations
- Mean 18,999 reports matched to previous cases
• Range 158-240 reports could not be assigned because of missing name, DOB, provider, or other critical matching data (specific labs flagged by city and state for corrective action)

**NYS electronic laboratory reporting lag indicators 2008-9:**

- **Western Blot**
  - Time from draw date to submission by laboratory (varies by lab, negotiated by state, range = 1-30 days, mean = 14 days)
  - Mean time from state to city = 7 days
  - Total time to city mean = 21 days

- **VL**
  - Time from draw date to submission (varies by lab, negotiated by state, range = 1-30 days, mean = 7 days)
  - Mean time from state to city = 21 days
  - Mean time from draw date to city = 30 days

- **CD4**
  - Varies by test result, transmissions every 3 weeks
  - Mean time from draw date to submission to state = 7 days
  - Mean time from state to city = 30 days

- **Genotype**
  - Mean time from draw date to submission by laboratory 90 days
  - Mean time from state to city = 30 days

**Field investigations:**

- 12,539 field investigations initiated on potential new cases (non-matches to registry July 1, 2008-June 30, 2009
- 4,472 (35.7%) dispositioned as new diagnoses
  - 20.9% previously reported
  - 40.1% non-cases
  - 2% patient not at site/chart not found
  - 1.5% out of jurisdiction

**Process:**
- 93% of new case investigations returned to surveillance within 2 months of initiation
- 50% of new diagnoses had provider report form (PRF) submitted as required by NYS Public Health Law Article 21
Vital Status Ascertainment

- Quarterly matches of HIV Registry with NYC vital registry file on HIV-related cause of death
- Semiannual match of HIV Registry with NYC death registry for all causes of death
- Annual match of HIV Registry against Social Security Death Master File
- Annual or biannual match of HIV presumed living cases in HIV Registry against National Death Index

Internal Duplicates and Merges

- Program algorithm (“DupMerge”) identified potential duplicate pairs:
  - Same last name, first name, DOB (16 pairs 2008-2009, 130 cumulative)
  - Same SSN (1 pair 2008)
  - Same death certificate and date of death (1 pair 2008, 400 cumulative)
  - Same NYSID number (500 cumulative pairs)
  - Other categories (300 cumulative pairs)

- Corrections
  - Identified, entered, and investigation initiated N=379
  - Investigation completed, fix in progress N=70
  - Investigation completed, fix accomplished N=253
  - Duplicates eliminated: 116 cases deleted

- New matching product in test mode now – IBM Quality Stage:
  - Uses reference dataset to standardize key variables, e.g., first name, last name, date of birth, Social Security number, street address, city and zip
  - Divides matching variables into blocks, sequentially matches and rates blocks in multiple data passes, and creates match weight
  - Test file: 4.6 million records in pre-eHARS laboratory database
    - 672,257 duplicate tests identified and removed
    - 3.7/3.9 million laboratory records rated “high scoring” matches by QS
    - Generated 3,500 cases for manual review
      - 780 duplicate pairs, many of which had already been identified by standard program algorithm and “DupMerge” facility
      - 2,700 cases with mismatched lab documents (represents 30,588 laboratory reports)

Interstate Duplicates – Routine Interstate Duplicate Review

- Backlog of 17,000 cases dating from 2005 was cleared in 2009, entered into Duplicate Review tab in eHARS and submitted to CDC
• Review completed of 2,795 (89%) potential duplicate pairs out of 3,148 potential interstate duplicates assigned for resolution by CDC in 2008.

Completeness and Timeliness of Reporting

• NYC used least squares regression to estimate number of new diagnoses expected in 2007

• Of 3,862 cases expected to be reported in 2007, 3,829 (99.2%) had been reported to the NYCDOHMH by December 31, 2008 (within one calendar year of the date of diagnosis)
Appendix III E: Washington, D.C. Core HIV Surveillance System
The District has had AIDS reporting since 1985 and transitioned to names based HIV reporting in November, 2006. Since the transition to names based HIV reporting, the District has worked to develop policies and procedures to control the quality, completeness, accuracy and timeliness of HIV reporting data. The District of Columbia requires named reporting of all diagnoses of HIV and AIDS, all HIV-related illness, all positive Western Blot (WB) tests for HIV antibody, all VL and CD4 lymphocyte values, and all HIV genotypes. The District’s enhanced HIV/AIDS Reporting System (eHARS) is a population-based registry that since 1981 has been continuously updated with new, de-duplicated diagnoses and laboratory results. All incoming provider and laboratory reports that do not match an existing registry record initiate a field investigation with medical record review to confirm the case, date and disposition of diagnosis and collect all other data required for surveillance and partner notification. The District’s eHARS also obtains data through routine matches with other disease registries; Sexually Transmitted Disease Surveillance system (STD*MIS), the Districts electronic Death Registry (eDeath Registry), electronic Birth Registry (eBirth) and HIV client level data from HRSA and AIDS Drug Assistance Program(ADAP) as well as the Social Security Death Master File. The surveillance program follows a set of monthly performance indicators to track data quality and staff and provider performance, and meets on a monthly basis to review indicators, identify problem areas and take appropriate action steps.

Laboratory Reporting Summary
According to the District’s Laboratory Licensing Administration, there are 27 laboratories licensed to do HIV testing in the District. Annually these laboratories are surveyed to assess their testing volume and the types of testing they perform. In the District, positive western blots and all CD4 and VL values are reportable. In 2008, only 2 smaller testing laboratories were found not to be in compliance with HIV laboratory reporting requirements, accounting for less than 1% of testing in the District. Completeness of reporting has improved significantly since the Districts transition to names based HIV reporting in 2006 and has continued to improve annually as more labs in the District begin electronic laboratory reporting (ELR). To improve on timeliness of laboratory reporting, in August, 2009, laboratories were notified of the legal requirements for reporting and received detailed instructions on tests they are required to report including data from 2007-present. Below is a summary of the laboratory reports received during 2008 and 2009.

The following points summarize the volume and periodicity of electronic laboratory reporting, the volume, timeliness and outcome of field investigations, the periodic registry matches, internal matching and deduplication procedures, and the interstate deduplication activities of the program.

Electronic laboratory reporting from laboratory to DC-DOH

Estimated total number of laboratory reports received and processed in the District of Columbia, 2008-2009

<table>
<thead>
<tr>
<th>Lab Test</th>
<th>2008</th>
<th>2009</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>WBLOT</td>
<td>2,175</td>
<td>1,138</td>
<td>3,028</td>
</tr>
<tr>
<td>CD4CNT</td>
<td>5,029</td>
<td>5,532</td>
<td>9,178</td>
</tr>
<tr>
<td>CD4PCT</td>
<td>4,870</td>
<td>5,378</td>
<td>8,903</td>
</tr>
<tr>
<td>VL</td>
<td>11,634</td>
<td>12,428</td>
<td>20,955</td>
</tr>
</tbody>
</table>
- Western Blot file received monthly (total N = 126)
- VL file received monthly (total N = 873)
- Number and Proportion of CD4, Low CD4 (<200), monthly (total N = 191 (37.1%))
- Number and Proportion of CD4, High CD4 (200-499) monthly (total N = 166 (32.2%))
- Number and Proportion of CD4, Very High CD4 (500+) monthly (total N = 158 (30.7%))
- Number and Proportion of VL, Undetectable (<400 C/ML), monthly (total N = 491 (46.3%))
- Number and Proportion of VL, Detectable (400-10,000 C/ML) monthly (total N = 264 (24.9%))
- Number and Proportion of VL, Very High (>10,000 C/ML) monthly (total N = 305 (28.8%))
- Mean 2,005 laboratory reports were received per month
- Mean 176 reports were potential new cases and initiated field investigations. These reports on average were for 53 unique people
- Mean 1,174 reports matched to previous cases or to 387 unique people

**De-duplication of eHARS**

All “newly reported” HIV cases must undergo thorough de-duplication within the District and with other jurisdictions. Monthly de-duplication of HIV/AIDS is done prior to sending records to CDC and completed using version SAS 9.0 and Linkplus. The key data elements for matching include the following:

- Name
- Date of Birth
- Gender
- Social Security Number

**Intrastate/Interstate De-Duplication**

If a duplicate exists, the record with the earliest date of diagnosis is retained and any additional information is added to this case record. In addition to reconciling Routine Interstate De-Duplication lists generated by CDC, any cases under investigation that are reported with an indication of being an out of state resident are referred to the appropriate jurisdiction.

**Review of other Documents to ensure completeness of reporting:**

The District employs the following routine database matches to assess the completeness, timeliness and quality of HIV reporting data.

**Vital Status Ascertainment**

- Death certificates are received monthly from Vital Records. Quarterly matches with DC vital statistics database
- Annual match of HIV Registry against Social Security Death Master File

**AIDS Drug Assistance Program (ADAP)**

- Quarterly matches conducted to update existing case data and to identify new cases to eHARS surveillance database
Medicaid

- Claims for persons with ICD9/10 codes indicative of HIV/AIDS are reviewed quarterly to update existing case data and to identify new cases

Timeliness

Approximately 85% of District laboratory reports are received within 2 weeks of the test date via electronic reporting. The remaining 15% are reported via the US mail, averaging about a 3 week lag in reporting.