Background. Retention in care is critical for treatment and prevention of HIV. Many HIV care clinics measure retention rate, but data are often incomplete for patients who are classified as lost to follow-up but may be actually in care elsewhere, moved, or died. The Data to Care (D2C) initiative supports data sharing between health departments and HIV providers to confirm patient care status and facilitate retention in care for all HIV positive patients.

Methods. The University of Chicago Medicine (UCM) provided an electronic list to the Chicago Department of Public Health (CDPH) of adult HIV-positive patients whose retention status was not certain. Retention in care was defined as at least 2 visits ≥90 days apart within the prior 12 months. CDPH matched this list of patients with data from the Chicago electronic HIV surveillance database. Matches were based on patient name, including alternative spellings and phonetics, and birth date. CDPH also cross-checked patient names with the CDC’s national enhanced HIV/AIDS Reporting System (DHARS) database. CDPH provided UCM with patient current care status, i.e., patient was in care elsewhere (as verified by lab data), moved out of state, or deceased.

Results. 780 HIV-positive patients received care in the UCM adult HIV clinic from January 1, 2013 to March 31, 2017. Of these, 360 were retained in care as of March 31, 2017. We shared data with CDPH for 492 patients. Of these, 294 (59.8%) were matched, and 168 (34.1%) had a date of last medical care provided. See Table 1 for patient dispositions, before and after data sharing. 24 (13.4%) of patients believed to be lost to follow up according to UCM records were confirmed either transferred care or deceased according to health department data.

Conclusion. Data sharing between the health department and HIV providers can improve data accuracy regarding retention in care among people living with HIV.

Table 1. HIV Patient disposition data before and after data sharing with health department

| Patient disposition | Before data sharing, n (%) | After data sharing, n (%) |
|---------------------|---------------------------|--------------------------|
| Currently in care at UCM | 360 (46.2%) | 360 (46.2%) |
| Transferred | 139 (17.8%) | 143 (18.3%) |
| Deceased | 63 (8.1%) | 63 (10.6%) |
| Incarcerated | 1 (0.1%) | 1 (0.1%) |
| Moved out of jurisdiction | 39 (5.0%) | 39 (5.0%) |
| Lost to follow up | 176 (22.8%) | 154 (19.7%) |

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1364. How Acceptable Is A Wireless Pill Bottle That Monitors and Texts In Response To Missed Doses: Focus Groups With Young African American MSM Living With HIV

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Background. African American MSM (AAMSM) living with HIV are less likely to have viral suppression than other racial groups. Wisepill, a wireless pill bottle, transmits a cellular signal to a server when opened and is designed to measure antiretroviral therapy (ART) adherence. The objective of this study was to explore the acceptability of a proposed intervention in these young AAMSM using the Wisepill device opening data to trigger a real-time text alert that ART may not have been taken during a planned time to either the user, a trusted social contact, or a healthcare worker, depending on the duration of consecutively missed doses (1 dose, 3 doses, 7 doses, respectively).

Methods. From December 2016 – May 2017, AAMSM living with HIV age 18–34 years (N = 25) participated in a study that included five focus groups (n = 23) and one on one interviews (n = 2). We performed theory-based discussion grounded in the Technology Acceptance Model. Specifically, we explored usefulness, convenience, concerns, and intention to use.

Results. Fifty-two percent missed at least one dose in the 4 days prior to the focus group meeting. Almost all participants (94%) favored the idea of a wireless pill bottle monitor and linked text message notification that ART may have been missed. The device was considered convenient for use at home or in a backpack, but too large for a pocket. Stigma and privacy were common concerns. For example, participants did not want to carry the device with them if the pills would “sound like a walking pharmacy” and did not want to get a text message that they had missed your HIV meds. They preferred text message notifications that ranged from emoji icons to cryptic short texts and wanted to receive an email as a backup plan. Most believed that the device appearance would not gain unwanted attention. Thirty percent of the participants identified a partner as a social contact they would not have missed your HIV meds would be sent whereas others designated their mother, aunt, brother, friend, pastor, and case-manager.

Conclusion. An adherence intervention using a wireless pill bottle monitoring device that sends notifications was accepted by most of the AAMSM in this study. Acceptability may be enhanced by personalization of the responsive text messages and a backup email option.

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1365. Improving Appointment Adherence among Young Adults with HIV in Peru with an mHealth Intervention: a Feasibility Study

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Background. Mobile Health (mHealth) interventions, including short message service (SMS) reminders and motivational messages, are associated with improved HIV appointment adherence, though feasibility is context-dependent. We assessed the feasibility of an mHealth intervention to improve appointment adherence among young adults with HIV in Lima, Peru.

Methods. Between November 2016 and April 2017, we implemented a one-way mHealth pilot intervention in an outpatient hospital without electronic medical records. We enrolled young adults (age 18–29) entering HIV care in a 3-component intervention: (i) reminder SMS prior to scheduled appointments (provider, laboratory, pharmacy); (ii) motivational SMS at each visit; and (iii) phone call following a missed visit. Feasibility evaluation included enrollment acceptance, visit tracking (information captured in the study database within 3 days of attendance), and proportion of intervention delivery (threshold >90%). We performed a qualitative assessment to identify implementation challenges reviewing staff field notes and meeting minutes.

Results. We enrolled 89/94 (95%) of eligible participants. The median age was 25 years and 83% were male. The median time of follow-up after enrollment was 115 (interquartile range [IQR]: 84–141) days, and participants had a median of 10 (IQR: 8–14) visits during the study period. Among 850 total participant visits, study personnel tracked 751 (88.4%); most (87.5%) untracked visits were due to technical issues (e.g., dropped call). Among scheduled appointments and 160 (21.3%) were unscheduled walk-ins. Intervention delivery reached 556/591 (94.1%) for reminder SMS, 733/751 (97.6%) for motivational messages, and 169/170 (99.4%) phone calls for missed visits, 127 (73.5%) of which were answered. Qualitative assessment revealed 3 major themes: real-time appointment tracking in a paper-based system consumed most staff time and resources, and meticulous in-person coordination between the implementation and hospital staff was essential for tracking.

Conclusion. An mHealth intervention to improve appointment adherence among young adults with HIV in Peru appears feasible with dedicated staff and a reliable appointment tracking system. Digitalized appointment systems may be needed to address challenges for scale-up.

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1366. Patient-level Factors Associated with HIV Diagnosis at Admission Across a Large Integrated Healthcare System

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Background. Linkage to care is a critical step for achieving HIV viral suppression and improving outcomes in newly diagnosed individuals. An unexpectedly high number of patients who have undiagnosed HIV or ineffective viral suppression are identified upon admission to acute-care. This study aims to understand factors that may be associated with lack of viral suppression for patients who are newly diagnosed at admission.

Methods. Patients with HIV, admitted to one of our eight acute care facilities were identified (n = 1,632) from medical records. Of these, 94 were newly diagnosed and 1,538 had a prior diagnosis of HIV. Factors that may impact the viral suppression status (≥200 copies/mL) prior to admission were examined. Median income and percent-age of individuals living below poverty were inferred from the American Community Survey (U.S. Census) data based on a patient’s zipcode. All other factors were extracted from the medical record at the time of admission. Chi square tests and t-tests were used to compare patients who were virally suppressed to those who were not.

Results. The average age was 48 years old and 60% were male and 82% black race. Patients who presented and were not virally suppressed were more likely to be between the ages of 40–60 and of black race. Distributions of age groups and race were significantly different between those virally suppressed and those who were not suppressed at admission (P < 0.05 for both). Blacks and whites between 40 and 60 years were more likely than other race and age groups to present at admission without viral suppression. Patients from geographic areas associated with high poverty and lower median income were less likely to be virally suppressed. (23.2% vs. 25.2%; P < 0.05 for poverty and $41,183 vs. $43,757; P < 0.05 for income).

Conclusion. These results indicate that age, race and geographically inferred income and poverty are significantly different between patients who are virally suppressed and those who have a detectable viral load. Further investigation is needed to better understand how these patient-level factors, including sociodemographics, impact linkage to care as well as how best to allocate resources to better engage and retain patients in HIV care to improve their long-term outcomes.

Disclosures. All authors: No reported disclosures.