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Accessibility
Improving adherence to antiretroviral treatment in Uganda with a low-resource facility-based intervention

Celestino Obua1*, Joshua Kayiwa2, Paul Waako1, Göran Tomson3,4, Hudson Balidawa5, John Chalker6, Dennis Ross-Degnan7 and Rolf Wahlstrom3

1Department of Pharmacology and Therapeutics, School of Biomedical Sciences, College of Health Sciences, Makerere University, Kampala, Uganda; 2Data Department, Joint Clinical Research Centre, Kampala, Uganda; 3Department of Public Health Sciences, Division of Global Health (IHCAR), Karolinska Institutet, Stockholm, Sweden; 4Medical Management Centre (MMC), Karolinska Institutet, Stockholm, Sweden; 5The AIDS Control Programme, Ministry of Health, Kampala, Uganda; 6Centre for Pharmaceutical Management, Management Sciences for Health, Arlington, VA, USA; 7Department of Population Medicine, Harvard Medical School, Boston, MA, USA

Objective: To assess the effects of facility-based interventions using existing resources to improve overall patient attendance and adherence to antiretroviral therapy (ART) at ART-providing facilities in Uganda.

Methods: This was an interventional study which tracked attendance and treatment adherence of two distinct cohorts: experienced patients who had been on treatment for at least 12 months prior to the intervention and patients newly initiated on ART before or during the intervention. The interventions included instituting appointment system, fast-tracking, and giving longer prescriptions to experienced stable patients. Mixed-effects models were used to examine intervention effects on the experienced patients, while Cox proportional hazards models were used to determine the intervention effects on time until newly treated patients experienced gaps in medication availability.

Results: In all, 1481 patients’ files were selected for follow-up from six facilities – 720 into the experienced cohort, and 761 into the newly treated cohort. Among patients in the experienced cohort, the interventions were associated with a significant reduction from 24.4 to 20.3% of missed appointments (adjusted odds ratio (AOR): 0.67; 95% confidence interval (CI): 0.59–0.77); a significant decrease from 20.2 to 18.4% in the medication gaps of three or more days (AOR: 0.69; 95% CI: 0.60–0.79); and a significant increase from 4.3 to 9.3% in the proportion of patients receiving more than 30 days of dispensed medication (AOR: 2.35; 95% CI: 1.91–2.89). Among newly treated patients, the interventions were associated with significant reductions of 44% (adjusted hazard rate (AHR): 0.56, 95% CI: 0.42–0.74) and 38% (AHR: 0.62; 95% CI: 0.45–0.85) in the hazards of experiencing a medication gap of 7 and 14 days or more, respectively.

Conclusions: Patients’ adherence was improved with low-cost and easily implemented interventions using existing health facilities’ resources. We recommend that such interventions be considered for scale-up at national levels as measures to improve clinic attendance and ART adherence among patients in Uganda and other low-resource settings in sub-Saharan Africa.

Keywords: antiretroviral therapy; adherence; management; staff motivation; intervention; Uganda

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*Correspondence to: Celestino Obua, Department of Pharmacology and Therapeutics, School of Biomedical Sciences, College of Health Sciences, Makerere University, P.O. Box 7072, Kampala, Uganda, Email: cobua@chs.mak.ac.ug

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In resource-limited settings, antiretroviral therapy (ART) programs strive to maintain high numbers of patients taking the life-long medications as countries scale-up ART accessibility (1–5). Adherence to treatment is one of the most important predictors of survival for individuals with HIV and AIDS (6, 7). For most HIV regimens, patients need to achieve >95% level of adherence so as to avoid developing drug resistance (8).
For 95% adherence, missing even one pill in a week on a regimen of two pills per day could lead to developing resistance to therapy.

Several factors such as distance to health facilities, availability and affordability of ARVs, lack of money for transport and food, quality of life during ARV, long waiting time, and congestion at health facilities influence adherence to ART (8, 9). Some of these factors require a programmatic approach, which may be difficult to address in a number of resource-limited settings (10, 11). Initiatives implemented at facility level, within facility resources, can affect a large number of patients more efficiently and are thus more sustainable. However, they need to be easy to introduce, implement, and should be affordable in terms of costs. In the past, initiatives designed to improve adherence have included training health personnel on ART supply chain management, HIV prevention, clinical care, counseling, community support programs, and use of experienced ART patients (expert clients) who provide support to other patients who are newly commencing treatment (6, 8). Frequently, such initiatives rely on substantial financial inputs that are often impossible to sustain by the health facilities in resource-limited settings (8, 12).

The current health-care system in Uganda was primarily designed for acute as opposed to chronic care. The system could be adapted to deliver chronic care through interventions targeted at improving clinic attendance and ART adherence. Many health facilities offering ART services lack systematic appointment systems, which results in overcrowding when patients come early hoping to be first to receive care. Subsequently, the congestion that develops negatively affects the quality of services as well as patient satisfaction with care (13). Furthermore, in many clinics, experienced patients already stabilized on ART are still required to present for care and go through the entire clinical process even if the only reason for their visit is to receive medication refills. To relieve congestion, these patients could be fast tracked to the pharmacy and/or dispensed longer durations of therapy.

In this study, we set out to determine the extent to which a set of easily implemented health facility-based interventions that used already existing resources at health facilities, mainly designed to reduce clinic congestion as well as patient waiting time, would improve clinic attendance and adherence among patients seeking ART services at Ugandan health facilities.

Methods

Study setting and design

Six district-level public hospitals previously determined to have poor adherence performance to ART were enrolled in this study based on the results of a previous national survey (8). All of these clinics had poor clinic flow, no or poor appointment systems, congested clinics with long waiting times, medication gaps, or medications not dispensed to patients after prescription (8, 9). These health facilities were supervised by the Uganda Ministry of Health (MoH) with ART funding from the Government of Uganda and donor agencies, including the Global Fund to Fight AIDS, Tuberculosis and Malaria and the U.S. President’s Emergency Plan to Fight AIDS Relief (PEPFAR).

This was designed as an interventional cohort study, which tracked both experienced patients and those newly initiated on antiretroviral drugs (ARVs) using facility-based records. The major aim was to assess the impacts of the interventions on patients’ clinic attendance and adherence. Implemented over a period of 17 months in 2009 and 2010, the study consisted of collecting retrospectively pre-intervention data, an implementation phase of the interventions, and follow-up visits to the health facilities while prospectively collecting post-intervention data. During the pre-intervention period, we also visited each study facility to explain the purpose of the interventions to the clinic staff and to gain consensus on how to conduct the intervention.

The effects of the interventions were assessed by following two groups of patients using their clinical and dispensing records at the health facilities. In each facility, one randomly chosen cohort composed of experienced patients who had been on treatment at least 12 months prior to the interventions, and who were still in treatment 6 months before baseline. The other cohort, consecutively chosen, composed of all newly treated patients over their first 4 months, who initiated treatment during the 6-months prior to the baseline up to 4 months before the end of the follow-up period. After 4 months, these patients were transferred into the experienced cohort. The study was based on patients’ clinical records and facility dispensing records that were selected based on the criteria above. There was no direct involvement between the research team and the patients.

At each health facility, we instituted two facility-level interventions. The impacts were measured at the patient level through time series analysis. These interventions aimed at improving patient flow and decongesting the clinics. We did not include a control group as the interventions implemented at the health facility level affected every patient attending the same facility.

Description of the interventions

The first component of the interventions was to introduce an appointment system and fast-track the ‘refill-only’ patients. Patients categorized as ‘refill only’ were experienced patients, who had been on treatment for at least 12 months from baseline, with a self-reported record of 95% adherence to medication regimen. These were given appointments to come for refill of medications only and
were fast-tracked through the clinic workflow without the need to be seen by clinicians or counselors if they did not report any complaints at triage. Traditionally, all patients would be required to pass through all stages of the clinic workflow, where they would be reviewed by the counselors and clinicians before being dispensed with refill medication. This was in spite of whether they had health complaints or not, and would subsequently lead to clinic congestions. The appointment system consisted of a book where the next review date would be recorded and patients were encouraged to return on that particular date. An appointment diary was also given to the patient to act as a reminder. Thus, the clinicians would be able to ascertain in advance which patients to expect on any particular day, and by the end of the clinic day, which patients had not honored their appointments. Currently in most Ugandan health facilities, patients are not given specific hours of appointment on the review dates. Furthermore, electronic dispensing is rare in the health facilities because of its associated technological challenges.

The second component of the interventions was to recommend that prescribers at the health facilities increase the numbers of days of medicines dispensed (longer prescription) from the traditional 30–60 days or 90 days for patients considered by the clinicians to be 95% adherent to their medication regimens. The aim was for such patients to make less frequent visits to the clinic, thus further reducing clinic congestion.

Health providers were provided with clear instructions to counsel the patients to report any problems during the scheduled visits or at any times without waiting for appointment days. This was to ensure that the patients remained in contact with clinicians at any time. In addition, the patients receiving longer prescription were to be seen by the clinician at least once every 6 months even when they had no complaints.

Apart from the above changes instituted by the research team, clinics continued to operate normally. The introduction of the appointment diaries and the appointment books provided the records of clinic attendance. The facilities also used these data for identification and follow-up of patients who missed appointments. Two weeks prior to implementing the interventions, we visited each study facility to explain the purpose of the interventions to the clinic staff and to gain consensus on how to conduct the intervention. After the intervention began, we made fortnightly supervisory visits, supplemented by telephone calls to ensure proper conduct throughout the post-intervention period. The research team was not directly involved in the practical implementation of the interventions.

**Sample selection**

We aimed to follow a cohort of 100 experienced patients per facility, as recommended in WHO manuals on investigating longitudinal patterns of medication use (14, 15). Anticipating a 20% loss to follow-up during the study period, we thus set out to identify 125 experienced patients per facility. Assuming that a minimum of 80% of the total of 600 patients would complete all scheduled visits, this would mean a 95% confidence interval (CI) of ±4.5% on the sample-wide estimates of monthly rates, which was our primary focus. For newly treated patients, the sample size was determined by the rate of treatment initiation at each facility. We aimed to include all patients who were initiated on treatment during the study period.

**Data sources and data collection**

Four research assistants used in this study were the same ones used in the earlier studies (8, 9). They were trained by the research team to retrieve prescription and clinical records from the patient files and the dispensing records, using methods recommended in the WHO manual on investigating longitudinal patterns of drug use (14, 15). They visited the health facilities with the intervention team to collect data but did not interact with patients. Data were collected from records in the clinic, pharmacy and appointment registers/diaries. During the 11 months of the post-intervention period, we collected data regarding the patients’ appointment dates, actual return dates, and the numbers of pills or days of medication dispensed. From these data, we computed the proportion of missed appointment and the days without medication (medication gaps) for each patient.

**Outcome measures**

The following indicators were used to assess the impact of the interventions among the experienced cohort: the proportion of patients with any scheduled visits missed; the proportion of patients with more than 30 days of dispensed medication days; and the proportion of patients with 3 or more days without medication (medication gap).

A different analysis was performed for the newly treated cohort. This was because of the nature of enrolment of patients into this cohort – where some were enrolled during the post-intervention period. In this cohort, we investigated the event-rates of the patients experiencing a medication gap of seven or more days, and of 14 or more days during the first 120 days of treatment.

**Statistical analysis**

We allowed 2 months after the initiation of the intervention before collecting post-intervention data. In this analysis, we have indicated this period as the ‘during’ period. The rationale was that the effects of the interventions would not accurately be measured immediately after instituting them. The interventions however continued to be implemented until the end of the study duration, overall 2 years.
The baseline characteristics of the two cohorts are presented using frequency tabulations for categorical variables, and descriptive summaries for the continuous variables. To assess the intervention effect, we first constructed graphical representations for each study outcome for the pre-, during and post-intervention periods. Overall aggregates of these outcome data per intervention periods have also been presented in tabular formats. We however excluded the ‘during’ intervention data from the modeling activities. For the experienced cohort, we modeled the intervention effect using mixed-effects models for binary responses with exchangeable correlation structure (16, 17). These models were adjusted for the patient-level demographic and clinical characteristics, including age, gender, initial treatment regimen, and pre-ART weight.

In the newly treated cohort, we used Cox proportional hazards survival models to determine the differences in event-rates of experiencing a medication gap of ≥7 days and ≥14 days after commencing therapy before and after the interventions. Patients were censored at the earliest of the following dates: on experiencing the outcome of interest; on the day of their last visit before reaching 120 days of treatment; or on the date of their last visit before the start of the intervention (for those initiating therapy within the pre-intervention period) or the end of follow-up (for those initiating therapy within the post-intervention period). We constructed the survival curves to depict the fraction experiencing the outcome for ease of interpretation. Statistical significance for the differences in the event-rates was assessed using the log-rank test. Cox proportional hazard models were fitted to determine the adjusted differences in survival between the intervention periods.

All analyses were performed using Stata/IC 10 (Stata Corp., College Station, TX, USA). All statistical inferences were assessed at the 5% type 1 error level, and all p-values presented were based on two-tailed tests.

**Ethical issues**

Ethical clearance for the study was obtained from Makerere University Faculty of Medicine Research and Ethics Committee, while permission to conduct this study was obtained from the Uganda National Council of Science and Technology (UNCST). Consensus meetings were held with the heads of the health facilities before instituting the interventions in the health facilities. During these meetings, the practical implementation of the proposed interventions was discussed, given the workflow and operational settings of the health facilities would have to be adjusted. No written or verbal consent was sought from the patients, since the interventions were implemented by regular clinical staff at the facility from their routine records. No specific data were obtained directly from the patients during the study. The research team did not perform any data collection in the communities. Furthermore, all data were anonymized in all phases of data collection.

**Results**

**Baseline characteristics of patients in the experienced cohort**

In total, 720 patients with an average age of 38.8 years (SD ± 10.2 years) comprising 468 (65%) females and 252 (35%) males were recruited into the experienced cohort. Most patients (482/720, 67%) were aged 31–49, most had commenced therapy with advanced disease (540/720, 75%) at WHO stages 3 or 4. The majority (662/720, 92%) commenced treatment with the stavudine (40)–lamivudine–nevirapine regimen (Table 1).

**Baseline characteristics of patients in the newly treated cohort**

Overall, 761 patients (68% females) with an average age of 35.8 years (SD ± 12.5 years) were included in the newly treated cohort. More than half (60%) commenced

<table>
<thead>
<tr>
<th>Overall study population</th>
<th>Experienced cohort (n = 720)</th>
<th>Newly treated cohort (n = 761)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>407 (65.2%)</td>
<td>315 (68.0%)</td>
</tr>
<tr>
<td>Male</td>
<td>217 (34.8%)</td>
<td>148 (32.0%)</td>
</tr>
<tr>
<td>Age in years, average (sd)</td>
<td>38.8 (10.2%)</td>
<td>35.82 (12.5)</td>
</tr>
<tr>
<td>18–30</td>
<td>113 (19.0%)</td>
<td>147 (32.5%)</td>
</tr>
<tr>
<td>31–49</td>
<td>400 (67.1%)</td>
<td>252 (55.8%)</td>
</tr>
<tr>
<td>50 and older</td>
<td>83 (14.0%)</td>
<td>53 (11.7%)</td>
</tr>
<tr>
<td>WHO stage at ART initiation, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stage 1</td>
<td>31 (5.5%)</td>
<td>32 (8.5%)</td>
</tr>
<tr>
<td>Stage 2</td>
<td>110 (19.5%)</td>
<td>120 (31.8%)</td>
</tr>
<tr>
<td>Stage 3</td>
<td>371 (66.0%)</td>
<td>193 (52.0%)</td>
</tr>
<tr>
<td>Stage 4</td>
<td>51 (9.1%)</td>
<td>32 (8.5%)</td>
</tr>
<tr>
<td>Initial ART regimen, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AZT-3TC-EFV</td>
<td>5 (0.8%)</td>
<td>35 (7.5%)</td>
</tr>
<tr>
<td>AZT-3TC-NVP</td>
<td>35 (5.6%)</td>
<td>274 (58.8%)</td>
</tr>
<tr>
<td>TVD-EFV</td>
<td>2 (0.32%)</td>
<td>12 (2.6%)</td>
</tr>
<tr>
<td>TVD-NVP</td>
<td>1 (0.16%)</td>
<td>18 (3.9%)</td>
</tr>
<tr>
<td>d4T(30)-3TC-NVP</td>
<td>575 (91.9%)</td>
<td>114 (24.5%)</td>
</tr>
<tr>
<td>Othersc</td>
<td>8 (1.3%)</td>
<td>13 (2.8%)</td>
</tr>
<tr>
<td>Average (sd) weight at ART initiation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>52.6 (10.3)</td>
<td>50.63 (11.5)</td>
</tr>
<tr>
<td>Male</td>
<td>54.0 (9.8)</td>
<td>52.81 (14.9)</td>
</tr>
</tbody>
</table>

*p*On ART at least 6 months prior to the index visit.

*Initiated ART within 6 months prior to the follow-up period.

*Other regimens include TDF-3TC-NVP, d4T (30)-3TC-EFV, and d4T (40)-3TC-NVP.

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**Table 1.** Demographic and clinical characteristics of patients by cohort
therapy at WHO stages 3 or 4. The most common initial regimens were zidovudine–lamivudine–nevirapine (59%) and stavudine–lamivudine–nevirapine (25%).

**Impact of the interventions in the experienced cohort**

**Any scheduled visits missed**

The percentage of experienced patients who missed any scheduled appointments increased in the pre-intervention period, and then declined moderately during the post-intervention period (Fig. 1a). The decline was more pronounced during the first 6 months of the intervention before starting to rise again thereafter. The percentages of missed scheduled appointments in the pre- and post-intervention periods were 24.4% and 20.3%, respectively. Overall, the interventions were associated with a significant 33% reduction in the odds of missed visits (adjusted odds ratio (AOR): 0.67; 95% CI: 0.59–0.77) among experienced patients (Table 2).

**Patients with more than 30 days dispensed medication**

After the interventions, there was an increase in the percentage of patients receiving medicines for longer periods at their scheduled refill (Fig. 1b). The percentage of visits where patients received more than 30 days of medication refills increased from 4.3% in the pre-intervention to 9.3% in the post-intervention period (Table 2). Overall, the interventions were associated with a significant 2.4 times increase in the odds of experienced patients receiving more than 30 days of medication refills (AOR: 2.35; 95% CI: 1.91–2.89).

**Patients with medication gaps of 3 days or more**

The percentage of experienced patients who were recorded to have medication gaps of 3 days or more at each visit declined over time (Fig. 1c). The percentages of visits with medication gaps were 20.2% in the pre-intervention, and 18.4% in the post-intervention period (Table 2). Overall, the interventions were associated with a significant 31% reduction in the odds of experiencing a medication gap of three or more days (AOR: 0.69; 95% CI: 0.60–0.79) among experienced patients.

**Impact of the interventions in the newly treated cohort**

Generally, the cumulative probability of newly treated patients experiencing long medication gaps was higher in
the pre-intervention period as compared to the post-
intervention period (Fig. 2a). The rates per 1,000 person-
years of experiencing medication gaps of 7 or more days
was 7.7 (95% CI: 6.5–9.0) in the pre-intervention period
and 4.5 (95% CI: 3.6–5.6) in the post-intervention period
(log-rank p-value < 0.0001). Similarly, the rates per 1,000
person-years of experiencing medication gaps of 14 or
more days was 4.8 (95% CI: 4.0 to 5.8) in the pre-
intervention period and 3.1 (95% CI: 2.4–4.1) in the post-
intervention period (log-rank p-value = 0.0022). From
the adjusted Cox proportional hazards models, the inter-
ventions were associated with a significant 44% reduction
in the event-rates of experiencing medication gaps of
7 and more days (AHR: 0.56, 95% CI: 0.40–0.74), and a significant 38% reduction in the
event-rates of experiencing medication gaps of 14 and
more days (AHR: 0.62; 95% CI: 0.45–0.85) (Table 3).

**Discussion**

Our findings show that clinical quality can be improved
through a strategy whereby providers at facility level are
encouraged to implement simple and feasible interven-
tions to improve clinic patient flow that are adapted to
the resources at their own facility, combined with minimal
support supervision. We demonstrated a modest but
significant 17% relative decrease in the percentage of
patients who missed scheduled appointments. This trans-
slated into more patients keeping scheduled appointments,
and thus improving adherence amongst such patients.
Several studies have shown that appointment keeping is
important for treatment success among patients on ART
(18–20), and this study also demonstrated a decrease in
the number of patients who missed appointments over
time, although there was a tendency towards increase
again during the last few months of follow-up.

One of the interventions was to encourage clinicians
to increase the number of dispensed medication days
for patients who had demonstrated their ability to remain
consistently on therapy as a means to cut down the
number of clinic visits and thereby decongest the clinic.
We showed that the percentage of patients who were
dispensed medication for more than 30 days more than
doubled after the intervention. This practice has also
been implemented elsewhere with considerable success
(21–24). The greater than twofold increase in the pro-
portion of patients who had thirty and more days of
medication dispensed achieved in this study would help
reduce crowds at the clinic and make the care process

Table 2. Changes in adherence-based indicators over time for experienced\(^a\) patients

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Pre-intervention (%)</th>
<th>During intervention (%)</th>
<th>Post intervention (%)</th>
<th>Odds ratio (95% CI)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any scheduled visits missed</td>
<td>24.4</td>
<td>21.4</td>
<td>20.3</td>
<td>0.67 (0.59, 0.77)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Patients with ≥30 days dispensed medication days</td>
<td>4.3</td>
<td>8.0</td>
<td>9.3</td>
<td>2.35 (1.91, 2.89)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Patients with ≥3 days medication gaps</td>
<td>20.2</td>
<td>15.0</td>
<td>18.4</td>
<td>0.69 (0.60, 0.79)</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

\(^a\)On ART at least 6 months prior to the index visit.

![Fig. 2](image-url). Probabilities of experiencing a medication gap of (a) ≥7 days or (b) ≥14 days during the first 120 days of antiretroviral treatment in the newly treated patient cohort.

Citation: Glob Health Action 2014, 7: 24198 - http://dx.doi.org/10.3402/gha.v7.24198
for these patients more convenient. However, dispensing greater quantities of medication may be limited by low stock levels or stock-outs in some resource-poor health facilities. Although we did not determine the stock levels of medicines at the facilities, this study has shown that increased medication days can be an effective way to decongest clinics and thereby improve adherence to ART even in health facilities with limited resources.

The percentage of experienced patients, who had medication gaps of three or more days, decreased significantly following the interventions. Among newly treated patients, the interventions significantly reduced the likelihood of experiencing a medication gap of seven or more days by about one-half, and the risk of a medication gap of 14 or more days by over one-third. The literature suggests that three or more days without medication may predict treatment failure among ART patients, while medication gaps of 14 or more days have been reported to be related to drug resistance development (25–27). The demonstrated reduction in the risk of experiencing medication gaps therefore shows that introducing a patient-appointment system, fast-tracking experience stable patients through the clinic, and dispensing larger prescriptions, makes the clinic less congested improving overall attendance and may be effectively executed as a way of improving patient adherence and guarding against treatment failure.

Strengths and limitations

By engaging local health workers and empowering them to implement simple systems changes in their clinics, this study has shown that it is possible to improve patient attendance and adherence to ART without any increase in external resources. Introducing an appointment system, fast-tracking experienced patients, and dispensing larger prescriptions as part of routine service were intended to improve clinic flow, reduce congestion, and shortened clinic waiting time. These changes resulted in modest but significant improvements in adherence indicators. Other workers have shown that integrating interventions into routine service delivery does not only improve uptake of services but ensures sustainability (28).

The main strengths of the study are the design, the low-resource interventions and the robust analysis. The interventions were actively implemented by the staff at the respective facilities and accepted into their routine. They could be contextually adapted and institutionalized to promote sustainability. The relatively modest effects on attendance and adherence should be seen in light of the minimal input of extra resources and the huge benefits for those individuals, who maintain treatment. If the same type of interventions would be implemented at a national level with similar effects, another 15,000–20,000 of the 200,000 patients on treatment (29) would keep their appointments and thereby experience a better change to stay adherent.

Despite the success of this approach, there are many challenges inherent in implementing real-world interventions in low-resource settings. Unfortunately, changes in policy and programs during the study period were critical events that may have affected the success of the interventions described in this study. For example, during the implementation period, the guidelines for first-line ARVs were changed. Many facilities were consequentially unable to consistently stock ARVs to support the new regimens resulting in an increased rate of stock-outs, and due to the retrospective nature of the study we were not able to collect data on daily stock availability. In addition, and most devastating, some ART support programs curtailed their activities when funding from PEPFAR was withdrawn (30), resulting in disruption of supply, thus directly influencing access to medicines and attendance. During the study, some facilities also introduced a nutrition program, which competed for resources and staff attention. However by reducing patient load using the recommended interventions, this may have made more time available for these other important tasks without the need for more resources. Given all of these external changes, it is remarkable that the simple measures resulted in meaningful and statistically significant improvements in attendance and adherence.

Conclusions and recommendations

This study has demonstrated that easily implemented interventions designed to reduce clinic crowding, such as appointment systems, patient fast-tracking, and increasing medication days, and where existing resources are

Table 3. Probabilities of experiencing medication gaps of ≥7 days or ≥14 days during the first 120 days of treatment among newly treated patients

<table>
<thead>
<tr>
<th>Period</th>
<th>Number of events</th>
<th>Rate (95% CI) estimates</th>
<th>Log-rank p-value</th>
<th>Adjusted proportional hazard estimates</th>
<th>Intervention effect (95% CI) p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥7 days</td>
<td>Pre 145</td>
<td>7.68 (6.52, 9.04)</td>
<td>&lt;0.0001</td>
<td>[ref]</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td></td>
<td>Post 78</td>
<td>4.48 (3.58, 5.59)</td>
<td>0.62 (0.45, 0.85)</td>
<td></td>
<td>0.56 (0.42, 0.74)</td>
</tr>
<tr>
<td>≥14 days</td>
<td>Pre 102</td>
<td>4.81 (3.96, 5.83)</td>
<td>0.0022</td>
<td>[ref]</td>
<td>0.004</td>
</tr>
<tr>
<td></td>
<td>Post 58</td>
<td>3.14 (2.43, 4.07)</td>
<td></td>
<td>0.62 (0.45, 0.85)</td>
<td></td>
</tr>
</tbody>
</table>

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mobilized, are effective ways of improving adherence to ART among patients in Uganda. Key to the positive outcomes was the willingness of health workers to participate in the interventions to improve service quality. This way of implementing feasible interventions could be recommended for roll-out to all ART programs in Uganda and beyond. Success will largely depend on well-motivated staff members in the clinics, who are willing to make operational and workflow changes.

Authors’ contributions

CO, PW, GT, RW, and JC conceptualized, designed, and planned the research. CO, PW, and HB conducted the study. CO, JK, and DRD analyzed the data. CO and JK drafted the manuscript. All authors commented on the article drafts and approved the final version.

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Conflict of interest

The authors declare no conflict of interest.

References


