Common Elements Treatment Approach (CETA) for unhealthy alcohol use among persons with HIV in Zambia: Study protocol of the ZCAP randomized controlled trial

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ABSTRACT

Aims: Prevalence of unhealthy alcohol use and co-occurring mental health problems is high among persons living with HIV (PLWH) in sub-Saharan Africa (SSA). Yet, there is a dearth of evidence-based treatment options that can address both unhealthy alcohol use and comorbidities in SSA HIV care settings. Recent studies testing single-session alcohol brief interventions (BIs) among PLWH in SSA have suggested that more robust treatments are needed. This paper describes the protocol of a pilot randomized controlled superiority trial that will test the effectiveness of an evidence-based transdiagnostic multi-session psychotherapy, the Common Elements Treatment Approach (CETA), compared to a control condition consisting of a single session brief alcohol intervention (BI) based on CETA, at reducing unhealthy alcohol use, mental health problems, and other substance use among PLWH in urban Zambia.

Methods: The study is a single-blind, parallel, individually randomized trial conducted in HIV treatment centers in Lusaka. 160 PLWH who meet criteria for unhealthy alcohol use + mental health or substance use co-morbidities and/or have a more severe alcohol use disorder are eligible. Participants are randomized 1:1 to receive the single-session BI or CETA. Outcomes are assessed at baseline and a six-month follow-up and include unhealthy alcohol use, depression, trauma symptoms, and other substance use.

Conclusions: The trial is a first step in establishing the effectiveness of CETA at reducing unhealthy alcohol use and comorbidities among PLWH in SSA. If effectiveness is demonstrated, a larger trial featuring long-term follow-ups and HIV treatment outcomes will be undertaken.

1. Introduction

Unhealthy alcohol use, including heavy and hazardous use, heavy episodic (binge) drinking, and alcohol use disorders (AUD), is a globally recognized public health problem (Saitz, 2005). In sub-Saharan Africa (SSA), alcohol is the most commonly consumed and distributed substance of abuse (Iahn, Dobkin, & Mayanja, 2012), and unhealthy use is attributable for 1% of all deaths among women and 4% of all deaths among men (Pithey & Parry, 2009; Rehm, Samokhvalov, & Neuman, 2009). Unhealthy alcohol use is common among persons living with HIV (PLWH) in SSA (Nouaman, Vinikoor, & Seydi, 2018), who are between two and four times more likely than the general population to have an AUD (Petry, 1999). In Zambia, preliminary studies found that 47% of male and 16% of female PLWH had unhealthy use during their first year on antiretroviral therapy (ART) (Vinikoor et al., 2015). Comorbidity between unhealthy use and mental health problems is very...
common: among Zambians who drank at unhealthy levels, over 50% met criteria for a severe AUD and/or mental health problems and over 30% reported other substance use (Kane et al., 2016, 2017).

Globally, there is recognition that unhealthy alcohol use is significant barrier to ending the HIV/AIDS epidemic (Bedoya et al., 2012; Hahn & Samet, 2010; Kalichman, Simbayi, & Vermaak, 2008). Alcohol has been shown to impact HIV outcomes through both behavioral and biological mechanisms (Hahn & Samet, 2010), including: cognitive impairment (Vagenas et al., 2015), increased sexual risk behaviors (Gerbi, Habtemariam, Tameru, Ngawu, & Robnett, 2009), depression (Leserman, 2008), immunocompromise with increased susceptibility to infections (Hahn & Samet, 2010), and nutritional deficiencies (Fawzi, Msamanga, Spiegelman, & Hunter, 2005; Watzl & Watson, 1992).

Despite the considerable burden of disease, treatment for unhealthy alcohol use is unavailable in many SSA HIV clinics. In high income country HIV clinic settings, brief alcohol interventions (BIs) have been tested for reducing unhealthy alcohol use (Aharonovich, Hatzenbuehler, & Johnston, 2006; Gilbert, Ciccarone, & Gansky, 2008). BIs typically range from 1 to 4 sessions lasting anywhere between 10 and 60 min each. They are often administered in primary care settings by a nurse or physician and include components such as: feedback on use and the harm drinking may be causing, norm referencing, identifying coping strategies, and motivational enhancement for changing behavior (Jonas, Garbutt, & Amick, 2012; Kaner et al., 2007, 2011; Platt, Melendez-Torres, & O’Donnell, 2016; SAMHSA, 2017). Literature from high income countries suggests that BIs in primary care settings can be both cost and clinically effective in reducing unhealthy alcohol use (Kaner, Dickinson, & Beyer, 2007; Platt et al., 2016; Wurzke, Shiel, Gomel, & Conigrave, 2001), including among PLWH (Aharonovich et al., 2006; Chander, Hutton, Lau, Xu, & McCaul, 2015; Gilbert et al., 2008). However, BIs were not designed to treat more severe AUD or comorbid mental health/substance use problems (NIAAA, 2005b; World Health Organization, 2012). Further, in SSA, evidence for the effectiveness of single-session BIs among PLWH is limited. Two studies conducted in Uganda and South Africa among HIV-affected populations found BIs were not superior than a control condition at achieving reduced alcohol consumption (Peltzer, Naidoo, & Louw, 2013; Wandera, Tumwesigye, & Nankabirwa, 2017). In Uganda, authors suggested that a single session may not have been a substantial enough dose and that additional sessions may be required to improve upon standard of care (Wandera et al., 2017); in South Africa, authors posited that the lack of treatment effect may have been due to behavioral reactivity in the control group, actual effectiveness of the active control intervention (a health education leaflet on alcohol use), or natural changes in alcohol use as a result of receiving standard TB treatment (Peltzer et al., 2013).

Based on these results, it seems likely that a more comprehensive treatment approach, beyond the use of BIs alone, is needed to reduce the impact of unhealthy alcohol use on the HIV/AIDS response in SSA. One possible package of care could include a system in which patients receiving HIV treatment are screened for alcohol use, provided with a point-of-care brief intervention (if needed), and a referral to higher level treatment (if indicated). Such screening, brief intervention, and referral to treatment (SBIRT) public health programs in the U.S. have shown promise in treating alcohol and substance use problems and in preventing such problems from becoming more severe disorders (SAMHSA, 2017).

This paper describes the adaptation of an existing evidenced-based, transdiagnostic, 6–12 week psychotherapy treatment approach (CETA) for delivery in HIV care settings in SSA to address unhealthy alcohol use and comorbid mental health problems; the development of a novel single-session BI based on the alcohol treatment component of CETA (BI); and the protocol for testing CETA compared to the BI in a superiority randomized controlled trial in Zambia. The trial aims to evaluate the effectiveness of CETA compared to the BI in reducing unhealthy alcohol use, mental health, and other (non-alcohol) substance use problems among PLWH in Zambia. The study is a Stage 1 pilot/feasibility trial (NIH, 2017). Specific hypotheses include:

1. CETA and the BI can be feasibly delivered at HIV care settings in SSA.
2. Among PLWH with unhealthy alcohol use + comorbidities and/or a more severe AUD, CETA will be clinically superior in improving alcohol, mental health, and substance use outcomes compared to the single-session BI.

The purpose of the trial is to test the comparative effectiveness of CETA versus the BI with a goal of future studies and programs to potentially implement a stepped care system that includes an evidence-based BI (for unhealthy alcohol use alone) and more intensive treatment such as CETA (for unhealthy use + comorbidities and for more severe AUD) as part of an SBIRT program.

2. Methods

2.1. Design overview

The Zambia CETA Alcohol Pilot (ZCAP) is a single-blind, parallel, individually randomized controlled trial comparing the effectiveness of a one-session brief alcohol intervention (BI) to a full course of multi-session CETA treatment (Fig. 1). Participants are adults living with HIV who have high risk alcohol use and possible mental health or other substance use comorbidities. Participants are recruited from two hospitals with large HIV clinics in Lusaka, Zambia during a regular HIV care visit, screened for eligibility, and randomized to receive the BI or CETA. Participants randomized to CETA also receive the BI after screening and before their first CETA session, as would be done in an SBIRT program. Assessment of the primary outcome (alcohol use) and secondary outcomes (mental health, other non-alcohol substance use) is completed at baseline and a six-month follow-up visit.

In addition to the high-risk group of participants in the RCT, we are also enrolling a lower risk group of individuals with HIV who have low-to-moderate risk unhealthy alcohol use and who do not have mental health or other substance use comorbidities (i.e., participants who do not meet symptom threshold criteria [see Measures] for depression, trauma, or substance use, and who have low-to-moderate risk alcohol use but do not meet alcohol use disorder [AUD] criteria). These lower risk participants are enrolled into an uncontrolled ‘cohort’ study and receive the BI; they are also assessed for outcomes at baseline and six-months post-baseline but are not randomized. The purpose of this cohort is to collect preliminary data to begin exploring the utility of the BI for lower risk clients under a hypothesis that, although the BI may not be sufficient for the higher risk participants (such as those with comorbidities and/or severe AUD), it may be sufficient and less costly for lower risk participants than a more intensive intervention such as CETA. It is beyond the scope of the present study to formally test the effectiveness of the BI among this lower risk group.

The study was approved by the Columbia University Medical Center Institutional Review Board (IRB), the Johns Hopkins Bloomberg School of Public Health IRB, the University of Zambia Biomedical Research Ethics Committee, and the National Health Research Authority in Zambia. The trial was registered on ClinicalTrials.gov (NCT03966885; Date of registration 05/29/2019) before participant enrollment. This protocol paper follows the SPIRIT guidelines for reporting of clinical trial protocols (Chan, Tetzlaff, & Altman, 2013).

2.2. Recruitment

Recruitment commenced in June 2019 at two large urban HIV clinics in Lusaka. Before recruitment, a series of meetings between study investigators and clinic personnel was conducted. Study aims and procedures were presented during these meetings and feedback was obtained. The goal was to integrate recruitment and other study
procedures, including intervention provision, into HIV clinic systems with a minimal level of disruption to regular activities. Following these meetings, we conducted a training with clinic staff on eligibility criteria and participant recruitment.

For recruitment, regular health talks are given by lay healthcare workers in the HIV clinic waiting areas. The talks focus on the relationship between alcohol and HIV and also inform patients of the ongoing study. Patients are encouraged to talk to staff if they are interested and also told that their provider may give them some more information about the study. Clinic staff including doctors, nurses, and HIV peer educators (lay counselors from the catchment area) then further introduce the study in private to their patients with HIV who they believe might have unhealthy alcohol use. Potential participants who are interested in hearing more about the study are escorted to our on-site study room where they are met by a research assistant (RA). This strategy represents a likely entry point into alcohol interventions in a real-world (i.e., non-research) HIV care context.

2.3. Screening informed consent

A study RA meets with the potential participant and provides an overview of the study, including aims, purpose, procedures, burden, and risks and benefits. All participants are assured that their participation (or not) would not impact their regular HIV care services. The RA obtains written informed consent from participants for eligibility screening. Participants who are illiterate can provide a thumbprint instead of a signature and an impartial witness cosigns the consent form.

2.4. Eligibility screening and baseline assessment

Participants who provide informed consent are escorted by the RA to a separate room in the clinic to complete eligibility screening, which also serves as the baseline assessment.

Initial eligibility criteria for the overall study include:

- Recruitment from HIV clinics
- Consent and eligibility screening for unhealthy alcohol use: Males: AUDIT ≥ 8; Females: AUDIT ≥ 4
- Secondary screening for eligibility into RCT: Full baseline assessment via ACASI
- Higher risk for AUD OR unhealthy alcohol use WITH comorbidities: Males: (AUDIT ≥ 16) OR (≥ 8 + comorbidities); Females: (AUDIT ≥ 12) OR (≥ 4 + comorbidities)
- Hazardous alcohol use w/o comorbidities: Males: AUDIT ≥ 8 AND <16; Females: AUDIT ≥ 4 AND <12

All participants receive point-of-care one-off session intervention (apprx. 30 minutes) as would occur in an SBIRT program with trained counselor that includes: psychoeducation, advice, engagement, talking about change, goal setting, and skill building.

Fig. 1. Flow diagram.
Once age and HIV status are confirmed, the RA sets up a laptop computer that is pre-loaded with study assessment measures on an audio-computer assisted self-interviewing (ACASI) system. (Kane et al., 2016, 2016, 2017, 2018; Murray et al., 2018) ACASI permits participants to self-complete the questionnaire with the text presented on the laptop screen and also audially read to the participant through headphones. ACASI is programmed in English and the two most commonly spoken languages in Lusaka, Bemba and Nyanja. After the ACASI notebook has been setup, the participant completes the first two (out of five) assessment measures of the ACASI, including demographics (sex, age, education, employment, and marital status) and the AUDIT.

The ACASI automates a total AUDIT score and generates a screen for the RA informing them of the participant’s eligibility status. Participants are eligible if they meet criteria for hazardous drinking defined as a total AUDIT score \( \geq 4 \) among women or \( \geq 8 \) among men. Participants who do not meet this criterion exit the study. Participants who are eligible based on the AUDIT continue with three additional measures in the ACASI evaluating depression (CES-D), trauma symptoms (HTQ), and substance use (ASSIST; Table 1).

The ACASI scores all three measures and generates a summary screen for the RA indicating whether the participant has met clinically relevant cut-off scores for each measure (Table 1). An algorithm built into ACASI then alerts the RA to the participant’s eligibility status with respect to the RCT or the cohort study. If a participant meets criteria for one or more of the CES-D, HTQ, or ASSIST measures and/or meets a threshold for higher risk of AUD according to the AUDIT, the participant is eligible for the RCT. If a participant does not meet symptom criteria for CES-D, HTQ, or ASSIST, and their AUDIT score suggests unhealthy use but not an AUD, then they are eligible for the cohort study. Thus, the RCT includes the highest risk group of participants and the cohort study includes a group with moderate risk.

For all participants, exclusion criteria for the study includes:

- HIV negative status
- Not receiving care at one of the study clinics
- Currently psychotic or on an unstable psychiatric drug regimen
- Actively suicidal and needing immediate hospitalization
- Unable to provide informed consent

The study follows an approved safety protocol where participants who are identified as having safety concerns and/or psychotic symptoms are immediately seen by a CETA clinical supervisor for further assessment. Clinical supervisors then refer any clients in need of additional services to a psychiatrist at the University of Zambia Teaching Hospital (UTH).

2.5. Study informed consent

Following completion of ACASI, the RA obtains informed consent for the remaining study procedures; a separate consent form is used for those eligible for the cohort and those eligible for the RCT.

2.6. Randomization/blinding

Randomization of RCT participants is conducted following completion of the ACASI assessment and informed consent. Lists of randomization ID numbers were generated by a statistician not otherwise involved in the study before trial commencement. Four lists were generated: one each for males and females at the two study clinics. IDs within each list were randomly allocated on a 1:1 basis to BI or CETA.

| Table 1 | Outcome measures in the ZCAP trial. |
|---------|-----------------------------------|
| Outcome                  | Measure                                      | Description                                                                 | Clinically relevant cut-off score for eligibility                                      |
| Unhealthy alcohol use    | Alcohol Use Disorders Identification Test (AUDIT) (Babor et al., 2001; Saunders et al., 1993) | AUDIT is a 10-item measure of hazardous alcohol use. A total score is calculated across the items with a possible range of 0–40 and higher scores indicating more severe alcohol use problems. The AUDIT was previously validated for use in Zambia (Chishinga et al., 2011) | Initial eligibility for unhealthy use: \( \geq 4 \) among women or \( \geq 8 \) among men; eligibility for more severe problem / higher risk of AUD: \( \geq 12 \) among women or \( \geq 16 \) among men (Babor et al., 2001; NIAAA, 2005a) |
| Depression (secondary)   | Center for Epidemiological Studies-Depression (CES-D) (Radloff, 1977) | CES-D is a 20-item measure of depression symptoms. Participants are asked how often they experienced each symptom over the past week (0 = never or less than one day; 1 = 1–2 days; 2 = 3–4 days; 3 = 5–7 days). A total score is calculated with a possible range of 0–60 and higher scores representative of more severe depression symptomatology. The CES-D was previously validated in Zambia. (Chishinga et al., 2011) | CES-D total score \( \geq 16 \) (Vilagut, Forero, Barbaglia, & Alonso, 2016) |
| Trauma symptoms (secondary) | Harvard Trauma Questionnaire (HTQ) (Mollica et al., 1992) | HTQ is a 39-item scale assessing symptoms of post-traumatic stress. Participants are prompted to respond how often each symptom bothered them in the past week (1 = not at all; 2 = a little; 3 = quite a bit; 4 = extremely). An average item score is calculated with a possible range of 1–4 with higher scores indicative of greater trauma symptoms. A previous study in Zambia demonstrated strong internal reliability of the HTQ \((\alpha > 0.90)\) (Kane et al., 2017) | HTQ average item score \( \geq 2.5 \) (Mollica et al., 1992) |
| Substance use (secondary) | Alcohol, Smoking, and Substance Involvement Screening Test (ASSIST) (Humeniuk, Ali, & Babor, 2008) | ASSIST is a 7-item measure that evaluates frequency of use, abuse, and dependence symptoms for a range of substance types, including tobacco, alcohol, marijuana, inhalants, cocaine, sedatives, hallucinogens, methamphetamine, and opioids. A specific substance involvement (SSI) score is calculated for each substance type that a participant reports ever having used in their lifetime. An SSI score can range from 0 to 9. The ASSIST was previously validated in Zambia (Kane, Murray, Bass, Johnson, & Bolton, 2016) | A non-alcohol/tobacco SSI score on the ASSIST of \( \geq 27 \) (Humeniuk, Henry-Edwards, Ali, Poznyak, & Monteiro, 2010) |
CETA was adapted in the ZCAP trial for use in HIV clinic settings by study staff and local counselors and supervisors. CETA training and ongoing supervision use the apprenticeship model for training (Murray et al., 2011). A 10-day live training of local HIV peer educators in Lusaka was conducted by study authors (LKM and SSvW) followed by small group practice (2 h weekly) in which the newly trained counselors practiced the CETA elements with the oversight of a CETA supervisor. Following the eight weeks of practice groups, counselors attend regular weekly supervision for 2–3 h per week for further practice, and review/support of CETA cases. Supervisors meet on a weekly basis with the study authors for clinical supervision for all active study clients (LKM and SSvW). Counselors and supervisors are trained to monitor session activities to ensure fidelity. Counselors document and report session activities to local supervisors, who document details in a clinical database, and this is reviewed with a CETA trainer. If counselors miss steps, they are expected to complete or re-do the component. A similar fidelity process is used for the BI intervention.

CETA is being implemented individually (one-on-one format) with flexibility of elements and number of sessions based on a client’s symptom presentation and severity. Treatment typically lasts from 6 to 12 sessions, usually one hour each, on average. In the ZCAP trial, all clients receive the CBT for Substance Use Reduction elements due to inclusion criteria. Prior or during the first session, the counselor and patient agree on when and where to conduct the subsequent weekly sessions. If sessions are missed, counselors follow up via phone (and home visit if necessary) to reschedule and assure safety.

Participants who are randomized to CETA also receive the single session BI (see description below) (Vinikoor et al., 2015) following screening and before their first CETA session, which typically occurs one week later. The rationale for providing the BI to CETA participants is that this is how we believe an SBIRT program would be administered in a real-world setting. That is, patients would be screened for alcohol use, provided with a BI on the spot (if indicated), and following the BI session.

When possible, the same counselor who provides the BI to a participant also becomes their CETA counselor to facilitate continuity of treatment and build off the therapeutic relationship that was established in the BI session.

### 2.7.2. Control arm: BI

Treatment as usual in Zambia HIV settings for unhealthy alcohol use is currently an alcohol BI. However, in practice, the BI is inconsistently

| Element | Simplified name (Used in training) | Description |
|---------|-----------------------------------|-------------|
| Psychoeducation and Engagement | Introduction and Encouraging Participation | • Focus on obstacles to engagement  
• Linking program to assisting with client’s problems  
• Includes family when appropriate  
• Program information (duration, content, expectations)  
• Normalization/validation of current symptoms/problems  
• Strategies to improve physiological stress  
• Examples: deep breathing, meditation, muscle relaxation, and imagery. Others added by local cultures.  
• Identifying and engaging in pleasurable, mood-boosting, or efficacy-increasing activities |
| Anxiety management strategies | Relaxation |  
• Understand association between thoughts, feelings, and behavior  
• Learn to restructure thinking to be more accurate and/or helpful  
• Facing feared and avoided memories in detail  
• Gradual desensitization/exposure  
• Facing innocuous triggers/reminders in the client’s environment  
• Gradual desensitization/exposure |
| Behavioral Activation | Getting Active (GA) |  
• Develop a plan with the client and client’s family (when appropriate)  
• Additional referral/reporting when needed  
• Utilizes motivation and CBT principles and activities to get client buy-in and alter behavior patterns to change substance use/abuse behavior. |

### Table 2

CETA Elements.
delivered with variable content and is not evidence-based. Preliminary unpublished data by our team found that health workers reported a lack of training and comfort in delivering any kind of alcohol reduction counseling. Given that the standard BI was not consistently delivered, the content varied considerably between and within clinic settings, was not evidence-based, and that providers did not feel comfortable providing it, we believed it was an inappropriately weak control condition to test in an RCT. Further, the variability within the control condition would have made a comparison with the experimental arm very difficult to interpret.

We therefore made the decision to enhance the existing BIs in HIV settings by developing a BI that was adapted from the evidence-based CETA element for Substance Use Reduction. Development of the BI was done by the study authors (CKD, LKM, SSvW) and with input from local partners working in HIV care settings. The BI consists of six components: 1) assessment/screening for alcohol and substance use; 2) understanding the impacts of use; 3) exploring possibilities for change; 4) goal setting; 5) identifying reasons for use; and 6) skill building (Table 3). It is designed to be one session of approximately 20–30 min, in response to local partners input on feasibility in HIV care settings. Counselors work to reach the goals of the session with clients by using a structured guide, the Improving Your Health (IYH) worksheet (included as supplemental material). The IYH worksheet was developed as a user-friendly tool to help counselors structure the BI session, observe the 20–30-minute time goal, and to assist clients with lower literacy rates. It was also hypothesized that the worksheet would reduce stigma for clients by allowing them to point or circle the response options instead of needing to discuss them out loud with the counselor.

The BI training and supervision process is consistent with the apprenticeship model of training used for CETA (Murray et al., 2011). The same HIV peer educators trained as CETA counselors attended a 2-day live training on the BI held by one author (SSvW). Since counselors already knew the CETA element for substance use, this training focused on how to deliver it in a consolidated fashion, using the worksheet.

### Table 3

| Element            | Description                                                                 |
|--------------------|-----------------------------------------------------------------------------|
| Assessment         | • Assessing clients current drinking through completion of a two-week timeline follow back measure |
| Understanding impacts | • Reviewing core ways substance use can impact an individual family and community |
| Exploring Change   | • Identifying the ways substance use impacts the individual and their family directly |
| Goal Setting       | • Exploring possible ways the client would consider changing or reducing their use |
| Identifying the Reasons | • Setting a goal for one way the client could reduce in the next few weeks |
| Skill Building     | • Understanding motivations for using |
|                    | • Teaching a coping skill to help the client combat one of their primary reasons for use |

At follow-up, we will also conduct mixed methods interviews with approximately N = 30 participants. This will include a purposive sample of participants from a range of ages and both sexes who did and did not complete the BI and CETA. Focus group discussions with counselors and clinic staff who participated in the trial will also be conducted. Implementation factors to be explored from clients, counselors, and staff will include: acceptability, appropriateness, and feasibility, as well as the attitudes, thoughts, feelings, and barriers and facilitators related to implementation of SBIRT. We will systematically track throughout the course of the study: (1) the number of participants who successfully complete and decline BI; (2) the number of participants who successfully link to CETA, defined as attending the first CETA session; (3) the amount of time counselors dedicate to client tracking/retention; and (4) the number of clients who successfully complete CETA.

### 2.9. Data and safety monitoring

In addition to the ethical review boards, the trial is monitored by a three-person Data and Safety Monitoring Board (DSMB). All DSMB members reviewed and approved study procedures, as well as procedures for reporting and tracking all adverse events, overall study progress, and identifying any need for premature termination of the protocol. Progress reports are prepared for the DSMB every six months concerning enrollment, attrition, and adverse events and meetings are convened as needed. There are no plans for interim analyses.

### 2.10. Sample size and data analysis

The primary endpoint of the RCT is the difference in mean AUDIT score change from baseline to six-month follow-up between BI and CETA. We believe that an effect size of CETA ≥ 0.5 would be clinically significant and justify further investigation in a subsequent later Stage trial. Further assuming α = 0.05 and β = 80%, we require a total sample size of 128 (n = 64 per arm). To account for possible loss-to-follow-up/drop-out of 20% based on our previous studies with HIV-affected populations in Zambia, we inflated the sample size to 160 (n = 80 per arm). There is no sample size calculation for cohort study participants; however, the ceiling for the cohort is N = 160.

Primary analyses will be intent to treat (ITT). AUDIT score, along with other continuous outcomes (CES-D, HTQ, ASSIST scores) will be evaluated with linear mixed models. As a secondary analysis, we will also estimate generalized linear mixed effects models with binary AUDIT, CES-D, HTQ, and ASSIST outcomes using established cut-off
values. Fixed effects in all models will include treatment arm (0 = -BI; 1 = CETA), time (0 = Baseline; 1 = 6-months post-baseline), and interaction terms between treatment arm and time. Clinic may also be an included variable to account for clustering. Random effects will include client ID and counselor ID. Robust standard errors will be estimated using a sandwich variance estimator (Huber, 1967; White, 1980). For linear models, we will estimate the mean difference in change in score between arms and 95% confidence intervals as well as the estimation of Cohen’s D effect size (Cohen, 1960). For generalized models, we will estimate relative risks. Covariates may be included if they differ meaningfully at baseline or predict significant change in the outcome. In addition to the ITT analysis, we will also conduct a per protocol analysis that includes all participants in the BI arm and only those participants in the CETA arm who completed CETA. We will also explore moderators (e.g., sex) of treatment effectiveness. Paired t-tests will be used to assess within group change among cohort participants. Sensitivity analysis will also be conducted after excluding participants who report abstinence at endline (AUDIT of 0 points) and who have a positive EtG result, indicative of underreporting.

2.11. Trial status

The trial is currently ongoing. All trial activities, including recruitment, treatment, follow-up, and data analysis are expected to be completed by the end of 2020.

3. Discussion

Research has shown that PLWH in SSA are at high risk for unhealthy alcohol use (Hahn et al., 2012) and that, in turn, unhealthy alcohol use can lead to poor HIV care outcomes. (Vagenas et al., 2015) Studies have also suggested that PLWH with unhealthy alcohol use often also have co-occurring mental health or other substance use problems (Hahn & Samet, 2010; Kane et al., 2018; Vagenas et al., 2015). Yet, evidence-based alcohol treatment services are rarely available in SSA HIV settings, particularly within the highly prevalent context of comorbid presentations (Kane et al., 2018). The ZCAP trial is an initial step to fill this gap by testing CETA, a multi-problem, modular, transdiagnostic evidence-based therapy that is designed for addressing unhealthy alcohol use and comorbidities. By developing and testing a brief alcohol intervention based on CETA (BI) as well, we aim to ultimately build an SBIRT system that can accurately and efficiently provide appropriate levels of care to PLWH in Zambia and throughout SSA.

The size, scope, and resource constraints contribute to notable limitations for this ongoing ZCAP trial. First, we do not have the resources to formally test the BI in reducing unhealthy alcohol use among PLWH who do not have comorbidities or a severe AUD; we aim to capture preliminary data on the BI amongst this population through the cohort we are enrolling as part of this project and formally test it in future studies. Second, the study has a risk of underreporting and social desirability bias and this may be stronger in the CETA arm because of desirability bias and this may be stronger in the CETA arm because of the additional time spent with the study counselors, although it is also possible that a therapeutic alliance with counselors may have the reverse effect and increase the likelihood of accurate reporting. To reduce the effect of this bias should it occur, we are using ACASI to assess alcohol use and mental health. Also, we include the EIG test to detect participants who drink but report abstinence. It is possible that the participant’s knowledge of an upcoming EIG test could likewise result in behavioral reactivity and reduced alcohol consumption, however, we believe that an objective biomarker remains superior to self-report alone. Finally, our trial is occurring at two urban facilities in Zambia that are not representative of all HIV care environments in the country or in other SSA countries. The trial does not include PLWH in the community who do not access HIV clinics. Generalizability is also limited by the inability in this study to screen for alcohol among all patients in the HIV clinic. Future studies will be designed to address these generalizability limitations.

If the ZCAP study finds that the BI is feasible to deliver integrated within HIV care and that CETA is effective, we plan to conduct a hybrid implementation-effectiveness study (Curran, Bauer, Mittman, Pyne, & Stetler, 2012) to test an SBIRT system featuring the BI and CETA in several HIV care settings in Zambia and measure downstream HIV outcomes, including ART adherence, retention in care, and viral suppression.

Author Statement

The study was conceived and designed by JCK, MJV, AS, SS, LKM, and GC. LKM, SS, and CKD led intervention design, training, and oversight. JCK wrote the first draft of the manuscript. MEL led drafting of the Introduction. MJV, AS, SS, LKM, GC, MEL, TK, RP, JM, CKD, JC, and CC provided critical feedback on the first draft and contributed to writing of subsequent drafts. All authors contributed significantly to and edited all sections of the manuscript and have approved the final version.

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Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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