Reducing stigma and promoting HIV wellness /mental health of sexual and gender minorities: a novel group-based cognitive behavioural therapy program in Nigeria

Submission date	Recruitment status No longer recruiting	Prospectively registered		
19/02/2024		☐ Protocol		
Registration date	Overall study status Completed Condition category	Statistical analysis plan		
23/02/2024		Results		
Last Edited		[X] Individual participant data		
19/03/2024	Mental and Behavioural Disorders	Record updated in last year		

Plain English summary of protocol

Background and study aims

High levels of stigma due to identifying as a sexual or gender minority (SGM) and living with HIV are increasingly documented in the African setting, and often manifest as internalized stigma (self-stigma). Such stigmas, especially when they are multiple and compounding, impede psychosocial wellbeing as well as HIV prevention and care. Yet there are few, if any, interventions specifically focused on reducing intersecting internalized stigmas tested in Africa. In this study, the researchers will develop and evaluate a novel, group-based cognitive behavioural therapy (CBT) intervention for the MSM and TGW populations.

Who can participate?

Men who have sex with men (MSM) and transgender women (TGW) 18 years of age or older and at risk for or living with HIV in Lagos, Nigeria.

What does the study involve?

Participants are randomly allocated to either the immediate intervention or the delayed intervention group. The intervention comprises four weekly in-person group sessions facilitated by community health workers. The comparison group (i.e., the delayed intervention group) were made aware of the HIV and other services available through partner implementing organizations (e.g., HIV and STI prevention and treatment services). Surveys are carried out before and after the intervention and at a 3-month follow-up (immediate group only).

What are the possible benefits and risks of participating?

Participants benefit from the group sessions, specifically the opportunity to learn and talk about stigma, HIV services, being a sexual and gender minority in Nigeria, and having support from peers and community health workers.

The main risks posed to participants from the study and intervention procedures are psychological distress and breach of confidentiality. Psychological distress could arise from participating in the group sessions and surveys since they touch upon enacted and internalized

stigma, depression and anxiety, HIV risk, living in the HIV, and experiences of violence. Psychological distress could also arise from breaches of confidentiality, particularly given the stigmatized and even criminalized nature of identities and behaviours in question in the country context. While breach of confidentiality is unlikely given protections in place, any potential breach is concerning particularly given same-sex behaviour is criminalized in Nigeria, as well as the stigmatized nature of both same-sex behaviour and HIV.

Where is the study run from? Population Council (USA). The study is conducted in Lagos, Nigeria.

When is the study starting and how long is it expected to run for? October 2021 to October 2023

Who is funding the study? Elton John AIDS Foundation (USA)

Who is the main contact?

- 1. Dr Waimar Tun, wtun@popcouncil.org
- 2. Dr Julie Pulerwitz, jpulerwitz@popcouncil.org

Contact information

Type(s)

Public, Scientific, Principal investigator

Contact name

Dr Waimar Tun

Contact details

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Additional identifiers

Clinical Trials Information System (CTIS)

Nil known

ClinicalTrials.gov (NCT)

Nil known

Protocol serial number

Study information

Scientific Title

Evaluation of an internalized stigma reduction intervention for men who have sex with men, and transgender women, in Lagos, Nigeria via a prospective delayed group randomized controlled trial

Study objectives

The intervention will result in greater decreased internalized stigma related to being a sexual /gender minority, and related to living with HIV, among individuals randomized to the immediate vs delayed arm.

Ethics approval required

Ethics approval required

Ethics approval(s)

approved 15/02/2022, Population Council Institutional Review Board (1230 York Avenue, New York, 10065, United States of America; +1 (0)917 685 7660; ngontarz@popcouncil.org), ref: 988

Study design

Prospective delayed randomized controlled trial

Primary study design

Interventional

Study type(s)

Prevention, Treatment

Health condition(s) or problem(s) studied

Prevention of mental health disorders (depression, anxiety) and increased uptake of HIV prevention and treatment services

Interventions

After recruitment, from separate listings of MSM and transgender women (TGW) enrolled participants, an investigator used a random numbers table and assigned individuals from each list to an intervention group (either the immediate intervention or the delayed intervention group); there were separate groups for MSM or TGW (given different experiences and social networks). Within 10 days of completion of the baseline survey, research assistants called participants to inform them of their randomization assignment.

Group-based affirmative cognitive behavioral therapy: The intervention comprises four weekly in-person group sessions facilitated by community health workers. The evaluation design is a delayed intervention group randomized controlled trial, with pre-post surveys plus 3-month follow-up (immediate group only), as well as qualitative research with participants and program staff.

The comparison group (i.e., the delayed intervention group) were made aware of the HIV and other services available through partner implementing organizations (e.g., HIV and STI prevention and treatment services).

Intervention Type

Behavioural

Primary outcome(s)

Internalized stigma related to being a sexual/gender minority and living with HIV (among those individuals living with HIV), measured using an adaptation of the Internalized AIDS-Related Stigma Scale (IA-RSS) at baseline and three additional points (approximately 1 month after baseline, 2 months after baseline, and 3 months after baseline)

Key secondary outcome(s))

Measured at baseline and three additional points (approximately 1 month after baseline, 2 months after baseline, and 3 months after baseline):

- 1. Depression measured using the Patient Health Questionnaire-8 (PHQ-8)
- 2. Anxiety measured using the General Anxiety Disorder-7 (GAD-7), such as 'trouble relaxing' and 'not being able to control or stop worrying' in the last 2 weeks
- 3. Coping was measured using the 4-item Brief Coping Scale
- 4. Uptake of pre-exposure prophylaxis and antiretroviral therapy (among participants living with HIV). Among those with HIV, adherence was defined as reporting "No days" to the question "During the past 3 days, on how many days have you missed all your pills?". Antiretroviral treatment self-efficacy was assessed based on seven items from a validated scale (HIV-Adherence Self-Efficacy Scale with last month as the reference period.

Completion date

31/10/2023

Eligibility

Key inclusion criteria

- 1. 18 years or older
- 2. Assigned male at birth
- 3. [Among MSM subgroup] Had sex with a biological man in the last year; [Among Transgender women subgroup] Identify as a woman
- 4. Willing to participate in all four sessions of the intervention
- 5. Willing to participate in the study
- 6. Residing in Lagos State for the next 1 year

Participant type(s)

Healthy volunteer

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Upper age limit

99 years

Sex

Male

Total final enrolment

240

Key exclusion criteria

- 1. Do not agree to participate in either the group sessions or study surveys
- 2. Planning to move out of Lagos State within the next one year
- 3. Currently participating in any other stigma reduction intervention
- 4. Not providing consent to participate in the intervention and study
- 5. Not providing contact information for follow-up

Date of first enrolment

26/04/2022

Date of final enrolment

29/09/2022

Locations

Countries of recruitment

Nigeria

Study participating centre

Centre for Population and Health Initiatives (CPHI)

6C Ireti St, opposite Lebanese international school, Sabo yaba

Lagos

Nigeria

100001

Sponsor information

Organisation

Population Council

ROR

https://ror.org/03zjj0p70

Funder(s)

Funder type

Charity

Funder Name

Elton John AIDS Foundation

Alternative Name(s)

Elton John AIDS Foundation, Inc., Elton John AIDS Foundation Inc, EJAF

Funding Body Type

Government organisation

Funding Body Subtype

Trusts, charities, foundations (both public and private)

Location

United States of America

Results and Publications

Individual participant data (IPD) sharing plan

The dataset generated during the study will be stored in a publicly available repository (https://dataverse.harvard.edu/dataverse/popcouncil).

The type of data stored – quantitative de-identified data from the multiple rounds of survey. The process for requesting access (if non-publicly available): Although it will be made available on DataVerse, the researchers require anyone interested in the data to contact one of the two contact investigators (Julie Pulerwitz or Waimar Tun) for permission to use the data. Interested persons must state their affiliation, qualifications, intended purpose, and how they will use the data. Given the sensitive nature of the topic in this environment, the researchers use stringent measures before making the data available.

Dates of availability: 23 February 2024

Whether consent from participants was required and obtained: Yes

Comments on data anonymization: Data has been stripped of any identifiers.

Added 19/03/2024:

Link to dataset: https://doi.org/10.7910/DVN/SG5XLP

IPD sharing plan summary

Stored in publicly available repository

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<u>Dataset</u>		18/03/2024	19/03/2024	No	No
Participant information sheet	Participant information sheet	11/11/2025	11/11/2025	No	Yes