Mpoza, Edward; Rajasingham, Radha; Tugume, Lillian; Rhein, Joshua; Nabaggala, Maria Sarah; Ssewanyana, Isaac; Nyegenye, Wilson; Kushemererwa, Grace Esther; Mulema, Vivienne; Kalamya, Julius; +5 more... Kiyaga, Charles; Kabanda, Joseph; Ssali, Mina; Boulware, David R; Meya, David B; (2019) Cryptococcal Antigenemia in HIV therapy-experienced Ugandans with Virologic Failure. Clinical infectious diseases. ISSN 1058-4838 DOI: https://doi.org/10.1093/cid/ciz1069

Downloaded from: http://researchonline.lshtm.ac.uk/id/eprint/4655311/

DOI: https://doi.org/10.1093/cid/ciz1069

Usage Guidelines:

Please refer to usage guidelines at https://researchonline.lshtm.ac.uk/policies.html or alternatively contact researchonline@lshtm.ac.uk.

Available under license: http://creativecommons.org/licenses/by/2.5/
Cryptococcal Antigenemia in HIV therapy-experienced Ugandans with Virologic Failure

Edward Mpoza¹, Radha Rajasingham², Lillian Tugume¹, Joshua Rhein², Maria Sarah Nabaggala¹, Isaac Ssewanyana⁴, Wilson Nyegenye⁴, Grace Esther Kushemererwa⁴, Vivienne Mulema⁵, Julius Kalamya⁶, Charles Kiyaga⁷, Joseph Kabanda⁶, Mina Ssali⁷, David R Boulware², David B Meya¹²³

1 Infectious Diseases Institute, Makerere University, Kampala, Uganda
2 Division of Infectious Diseases & International Medicine, University of Minnesota, Minneapolis, MN, USA
3 School of Medicine, College of Health Sciences, Makerere University
4 Uganda National Health Laboratory Systems
5 Clinton Health Access Initiative, Uganda
6 Centre for Diseases Control, Uganda
7 Ministry of Health, Uganda

Corresponding Author: Edward Mpoza, Infectious Diseases Institute, Makerere University, Kampala, Uganda. Email: edmypoza@yahoo.com

Summary: In addition to the CD4 threshold of <100 cells/mcL, reflexive CrAg screening should be considered in persons failing ART with viral loads ≥5000 copies/mL.

© The Author(s) 2019. Published by Oxford University Press for the Infectious Diseases Society of America.

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.
Abstract:

**Background:** Detectable serum or plasma cryptococcal antigen (CrAg) precedes symptomatic cryptococcal meningitis. The World Health Organization (WHO) recommends CrAg screening for HIV infected people with CD4<100 cells/mcL initiating antiretroviral therapy (ART). However, an increasing proportion of cryptococcosis patients are now ART-experienced. Whether CrAg screening is cost-effective in those with virologic failure is unknown.

**Methods:** We retrospectively performed nationwide plasma CrAg testing among ART experienced Ugandan adults with virologic failure (≥1,000 copies/mL) using leftover plasma after viral load testing during September 2017–January 2018. For those CrAg-positive, we obtained ART history, meningitis occurrence, and 6-month survival via medical record review.

**Results:** Among 1,186 subjects with virologic failure, 35 (3.0%) were CrAg-positive with median ART duration of 41 months (IQR, 10–84 months). Among 25 subjects with 6-month outcomes, 16 (64%) survived, 7 (28%) died, and 2 (8%) were lost. One survivor had suffered cryptococcal meningitis two years prior. Two others developed cryptococcal meningitis and survived. Five survivors were known to have received fluconazole. Thus, meningitis-free survival at 6-months was 61% (14/23). Overall, 91% (32/35) of CrAg-positive persons had viral load ≥5000 copies/mL compared with 64% (735/1,151) of CrAg-negative (Odds Ratio = 6.0; 95% CI: 1.8-19.8, P = 0.001). CrAg prevalence was 4.2% (32/768) among those with viral loads ≥5000 copies/mL and 0.7% (3/419) among <5000 copies/mL.

**Conclusion:** In addition to the CD4 threshold of <100 cells/mcL, reflexive CrAg screening should be considered in persons failing ART in Uganda with viral loads ≥5000 copies/mL.

**Keywords:** Cryptococcal antigenemia, virologic failure, ART experienced, HIV
INTRODUCTION

Cryptococcal meningitis is responsible for 15% of AIDS-related deaths and is the leading cause of adult meningitis in Sub-Saharan Africa (1-2). Annual global deaths from cryptococcosis are estimated at 181,100, with 75% of these deaths in Sub-Saharan Africa (1). Cryptococcal antigen (CrAg) is an independent predictor of mortality and can be detected in blood by latex agglutination a median of three weeks before developing initial symptoms of clinical cryptococcosis (3-4). CrAg screening is cost effective, has survival benefit, and is recommended by the World Health Organization (WHO) in persons with CD4 count <100 cells/mcL, and can be considered below 200 cells/µL (5-9). Among ART-naïve outpatient populations in low and middle income countries, the CrAg prevalence in 2014 averages 6% with some regional variations (1). Among Ugandan ART-naive outpatients in 2004–2006 with CD4 <100 cells/mcL the CrAg prevalence is 8.8% (10).

There is a substantial proportion of ART-experienced patients with virologic failure presenting with fulminant cryptococcosis (11). During 2013–2017 in Uganda, nearly half of the patients presenting with cryptococcal meningitis were ART-experienced (11). These ART-experienced persons with virologic failure are at risk for cryptococcosis, yet they are not included within WHO CrAg screening guidelines’ high risk category (5, 12). As monitoring of those receiving ART is scaled up from CD4 to virologic monitoring in resource limited settings, CD4-based algorithms for CrAg screening may not be readily applicable to ART-experienced populations. Baseline CD4 counts may be performed at HIV diagnosis but not routinely while monitoring HIV treatment. The Uganda HIV guidelines 2018 recommend CrAg screening in ART naïve or treatment failure patients with CD4 cell counts less than 100 cells/mcL (13). In the absence of CD4 monitoring, it is unknown whether to screen persons with virologic failure.
Given the substantial proportion of ART-experienced patients presenting with cryptococcosis, new viral load-based CrAg screening strategies may be useful. Uganda has also adopted the ‘test and treat’ strategy which will enroll more people on ART irrespective of their CD4 counts. In addition we are now encouraging more virologic monitoring compared to CD4 monitoring. This creates a challenge for CrAg screening which are based on CD4 counts. We conducted this study to determine the CrAg prevalence and clinical outcomes of CrAg-positive persons among ART-experienced people with virologic failure. We evaluated the potential threshold of virologic failure where CrAg screening should be considered, and cost of CrAg screening at this threshold.

METHODS

Study Design, Setting and Participants

This cross-sectional study was conducted at the Uganda National Health Laboratory Services (UNHLS), which performs centralized viral load monitoring in Uganda (~95% of all HIV viral load testing). The UNHLS processed a total of 843,020 viral load tests for adults in 2017 of which 10% (84302/843020) had unsuppressed viremia of \( \geq 1,000 \) copies/mL (14). Dried blood spot and plasma samples from health facilities are delivered to the UNHLS via a hub transport system for HIV viral load testing (15). We retrospectively evaluated stored plasma samples of 1,186 ART-experienced HIV-infected adults (\( \geq 18 \) years) with suspected virologic failure with HIV viral loads above \( \geq 1,000 \) copies/mL. The sample size was estimated using Buderer’s formula (using an absolute precision of 0.1, a confidence Interval of 95%, an estimated prevalence of 8.9%(from a CrAg study in Nigeria which had largely ART experienced people;
(16)) and assumed a sensitivity of 50% giving us 1079 subjects to which we added a 10% margin for possible missing samples making a total of 1186 subjects. We tested left over plasma samples after viral load testing from specimens collected during September 2017–January 2018. The samples from September 2017- January 2018 were the most recent easily accessible and retrievable samples. We didn’t have stored samples of plasma for patients who had dried blood spots. CrAg LFA testing on dried blood spots has lower sensitivity compared to plasma (unpublished data). CrAg testing was performed from March to July 2018 followed by retrospective chart review to determine 6-month outcomes for the CrAg-positive patients.

**Study procedures**

Using the UNHLS database, we identified all patients with virologic failure and left over plasma samples from the months of September 2017 to January 2018. From these we selected a random sample of 1186 patients with left over samples using an auto generated command in Stata 14 (Stata Corp, College Station, TX, USA). The sampling technique used draws observations without replacement from a dataset. Quantitative RNA viral loads on plasma had been done by UNHLS using the COBAS AmpliPrep/ COBASTaqMan system. For this study, we performed CrAg testing on stored plasma samples at UNHLS; CrAg testing was performed using the CrAg lateral flow assay (LFA) (IMMY Inc., Norman, Oklahoma, USA).

Subjects’ demographics and clinical data were obtained from the viral load testing laboratory request forms. Outcome data were obtained by chart review and/or contacting health facility clinicians. Additionally, clinicians were duly informed of the positive CrAg results. Patients who had not had any interaction with their respective clinics in the prior 6 months (from the point of their viral load result) were defined as lost to follow-up. Meningitis free survival was defined as
having not suffered from clinical cryptococcal meningitis during the six month period after the HIV viral load testing.

Ethic Reviews

We obtained ethical review and approval from the Makerere University School of Medicine research and ethics committee, as well as Uganda National Council of Science and Technology, and the University of Minnesota. A waiver for informed consent was obtained as CrAg testing was performed on leftover plasma samples, and the study posed no more than minimal risk to participants.

Statistical methods

Data analysis was primarily descriptive with variables summarized by mean with 95% confidence interval (CI), median with interquartile range (IQR), and number (percentage). We tested continuous variables via nonparametric Wilcoxon Rank-Sum test and categorical data via Chi-square. We additionally used logistic regression to assess whether CrAg-positivity was associated with quantitative HIV viral load thresholds.

Costing

Once CrAg prevalence was determined, we used the inverse to calculate the number needed to test to detect one CrAg+ person. The cost of CrAg screening using the lateral flow assay in Uganda is $3.50 (17). We multiplied the cost of screening by the number needed to test to identify the cost to detect one CrAg positive person.
RESULTS

Prevalence of Cryptococcal Antigenemia in ART-experienced HIV patients with virologic failure

Overall 368,174 adult HIV patients had viral load testing from September 2017 to January 2018. We found 5,348 patients had virologic failure and left over plasma samples. We found 73 patients without left over samples for CrAg testing. Of the randomly selected 1,186 selected patients; 588 were from Central region, 191 from the East, 147 from the North, and 260 from the West. These samples were from 96 of the 127 districts in Uganda. Table 1 shows a summary of demographic and clinical characteristics of the study population by CrAg result. Of the 1,186 samples tested, 61% (724/1,186) were collected from women. Participants’ had a mean (±SD) age of 36 (±10) years and ranged 18 to 75 years. Among 1,186 ART-experienced persons with plasma viral loads ≥1000 copies/mL, we identified 35 CrAg positive persons equating to a CrAg prevalence of 3.0% (95%CI, 2.1% to 4.1%). Among those with virologic failure, the median HIV-1 viral load was 4-fold higher among CrAg-positive persons as compared with CrAg-negative persons (46,000 vs. 12,000 copies/mL; P<0.001).

Distribution of HIV-1 viral load among CrAg positive patients

Among the 1,186 persons, 65% (767/1,186) had viral loads ≥5,000 copies/mL. Overall, 91% (32/35) of CrAg-positive persons had viral loads ≥5,000 copies/mL compared with 64% (735/1,151) of CrAg-negative (Odds Ratio = 6.0; 95% CI, 1.8 to 19.8, P=0.001). CrAg prevalence increased among higher viral loads with 4.2% (32/768) CrAg-positivity among those with ≥5,000 copies/mL versus 0.7% (3/419) CrAg-positivity among those with <5,000 copies/mL. Figure 1 displays prevalence of CrAg positivity with respect to HIV viral load.
Outcomes of CrAg positive patients with virologic failure

Among the 35 CrAg-positive persons, we obtained 6-month survival data for 25 persons. We didn’t get outcome data on the other 10 CrAg positive persons because efforts to contact their respective clinicians or to access their medical files were futile. Of the 25 CrAg positive persons with outcome data, 16 (64%) were known to be alive, 7 (28%) were confirmed dead, and 2 (8%) were lost to follow up.

Of the 25 CrAg-positive, only 1 person had a history of treatment for cryptococcal meningitis 2 years prior and was taking fluconazole for secondary prophylaxis. Of the 25 CrAg positive persons, 17 were getting their first ever CrAg test. Five survivors were known to have been CrAg tested and received pre-emptive therapy fluconazole around the time of their viral load result. Another two patients developed cryptococcal meningitis during these 6 months were treated and survived. Thus, meningitis-free survival at 6-months was 61% (14/23; 95%CI, 35% to 76%). Absent fluconazole receipt (n=19), 6-month meningitis free survival was 42% (8/19; 95%CI, 20% to 67%).

We also did an analysis including those who were lost to follow up. We used 6 month outcome findings from a CrAg+ cohort in Uganda which found 14% mortality rate, 77% meningitis free survival and 9% progression to meningitis(10). If we applied this to the 10 without survival data, we would have 8 who survived meningitis free, 1 who died and 1 who developed cryptococcal meningitis. This would then make a 63% (22/35) meningitis free survival rate. If we were to consider that all the 10 without survival data died then the meningitis free survival would be 40% (14/35).

We determined the likely outcome of those lost to follow up (including those without outcome data) by comparing their sex, median ages and viral loads with those who survived and died. We
used the Fischer’s exact test for the sex comparison across the three groups and found no significant difference (p=0.42). The Kruskal Wallis test was used to compare the medians for age and viral load across the three groups and the test was not significant for both age (p = 0.42) and viral load (p = 0.10). Therefore, the likely outcome of the individuals lost to follow up may be similar to those who were accounted.

**Cost of CrAg screening**

We identified 4.2% CrAg prevalence among persons with a HIV viral load $\geq 5000$ copies/mL. Thus, the number needed to test to detect one CrAg positive person is 25. At a cost of $3.50 for the CrAg lateral flow assay (17), the cost to detect one CrAg positive person among those with virologic failure is $87.50.

**Discussion**

The CrAg prevalence was 3.0% in ART-experienced HIV patients with virologic failure and 4.2% among those with $\geq 5000$ HIV RNA copies/mL. To our knowledge, this is the first study to evaluate CrAg prevalence in ART-experienced patients with virologic failure, irrespective of CD4 count. This CrAg prevalence rate was similar to that in ART-experienced populations in studies in South Africa 2.8% (18), Brazil 3.1% (19), and in Ethiopia 4.1% (20) but lower than that in studies in Nigeria (8.9%) and Ethiopia (8.4%) though all these studies included participants responding to ART or didn’t report viral loads (12, 16). CrAg positivity was shown to significantly increase with higher viral loads indicating a higher risk for cryptococcosis in people with more fulminant virologic failure. We found an all-cause 6-month meningitis-free survival rate of 61% among CrAg-positive persons, of whom at least 6 survivors received
fluconazole therapy. Absent this fluconazole therapy, meningitis-free survival was ~40%, although the 95% confidence interval was wide.

We calculated that it would cost $87.50 to detect one CrAg positive person if screening is performed among persons with viral loads of ≥5000 copies/mL. In 2018, Uganda had 951,692 adult HIV viral load samples tested; 858,426 were suppressed, 6,661 tests were rejected, and 86,603 were virologic failures (VL >1,000)(14). From our sample, one could assume 64% of those with virologic failure had a viral load >5000 copies/mL. Thus 55,426 persons would be screened per year, and 2,327 would be CrAg positive. Thus, of the 951,692 total viral load samples received, the additional cost would be ~$193,991 in total, or $0.20 per viral load sample received.

As HIV programs ramp up test and treat and scale up virologic monitoring, we have observed increasing proportions of persons with cryptococcal meningitis presenting as ART-experienced persons with undetected virologic failure (11, 21). WHO guidelines recommend CrAg screening for persons with CD4<100 cells/µL prior to initiating ART (9, 22), however, for the ART experienced persons no guidelines exist. Given that virologic monitoring is emphasized more than CD4 monitoring in this population, using a viral load threshold to identify who needs CrAg screening becomes more pertinent. The guidelines acknowledge that CrAg screening in ART-experienced persons needs to be further evaluated. Based on our findings, in the absence of CD4 monitoring, we would recommend CrAg screening among persons with virologic failure ≥5,000 copies/mL as this threshold where we found a CrAg prevalence of 4.2%.

The study had limited access to patient medical information, thus missed some clinically relevant data, such as CD4 cell count, which could have been useful in further characterizing the
population of interest. This could also have introduced information bias in characterizing patients as meningitis free. The potential seasonal distribution of cryptococcal infections may have influenced prevalence results. While we only tested plasma specimens, whether dried blood spots would have a different CrAg prevalence is unclear as the CrAg LFA sensitivity is lower in dried blood spots than in plasma due to the extraction process (unpublished data). Viral load testing is being scaled up replacing CD4 monitoring yet absolute CD4 counts provide the risk stratification to target evaluation for opportunistic infections, including CrAg screening. The cost of CD4 testing (~$6) is more than the cost of CrAg testing (~$4), thus from a program implementation perspective, using CD4 testing to then narrow the pool to select for CrAg testing is not efficient, unless a CD4 is already being performed (17, 23). We should consider expanding the threshold for CrAg testing by including those with viral load ≥5,000 copies/mL which would decrease the amount of CrAg testing by 35% (419/1186) yet still detect >90% (32/35) of CrAg+ persons. The limitations of this approach are that it would be dependent on the turn-around time for viral load. Another limitation of the study was that we did not perform CrAg titers. This information could be useful to determine if persons with low plasma CrAg LFA titers ≤1:20 (24), may respond well to prompt switching to second-line ART only. The strengths of the study include generalizability of the results because we had access to samples from all regions of the country. This study also provided an opportunity to pilot implementation of the Uganda CrAg screening guidelines, which now include CrAg screening individuals with virologic failure and those with a positive symptom screen in the advanced HIV disease pathway(13). Further studies in different geographical areas where cryptococcosis is common are warranted in order to characterise the epidemiology of cryptococcal disease among persons failing ART and to evaluate the cost effectiveness of using viral load testing as an entry point to CrAg screening in this population.
Prospective studies on the implementation of such a screening strategy and clinical outcomes with fluconazole preemptive therapy are needed to further characterize this potential strategy to reduce AIDS-related deaths among persons with virologic failure.

This study shows that ART-experienced people failing HIV therapy are at risk for disseminated cryptococcal antigenemia and eventually cryptococcal meningitis or death and demonstrates the feasibility of linking CrAg screening to viral load monitoring. In order to further avert cryptococcosis among individuals with virologic failure, WHO should recommend CrAg screening in persons with virologic failure with viral loads ≥ 5000/mL. Cost effective studies are needed to establish whether CrAg screening based on viral load thresholds (alongside CD4 counts) would be feasible. Studies to determine the association of virologic failure independent of CD4 and CrAg are needed. Prospective studies among ART-experienced populations to further evaluate the optimal viral load threshold for CrAg testing, timing of ART switch, implementation of this strategy, and the potential benefit of CrAg screening are warranted.
Disclaimer:
The views expressed in this publication are those of the author(s) and not necessarily those of AAS, NEPAD Agency, Wellcome Trust or the UK government.

Funding: This research was supported by the NIH Fogarty International Center D43TW009345 (EM) and K01TW010268 (JR) and the National Institute of Allergy and Infectious Diseases via K23AI138851 (RR) and U01AI125003 (DBM, DRB, RR). DBM is supported by DELTAS Africa Initiative grant # DEL-15-011 to THRIVE-2. The DELTAS Africa Initiative is an independent funding scheme of the African Academy of Sciences (AAS)’s Alliance for Accelerating Excellence in Science in Africa (AESA) and supported by the New Partnership for Africa’s Development Planning and Coordinating Agency (NEPAD Agency) with funding from the Wellcome Trust grant #107742/Z/15/Z and the UK government.

None of the authors has a potential conflict of interest
REFERENCES

1. Rajasingham R, Smith RM, Park BJ, Jarvis JN, Govender NP, Chiller TM, et al. Global burden of disease of HIV-associated cryptococcal meningitis: an updated analysis. The Lancet Infectious diseases. 2017 Aug;17(8):873-81.

2. Durski KN, Kuntz KM, Yasukawa K, Virnig BA, Meya DB, Boulware DR. Cost-effective diagnostic checklists for meningitis in resource-limited settings. Journal of acquired immune deficiency syndromes. [Research Support, N.I.H., Extramural]. 2013 Jul 1;63(3):e101-8.

3. Liechty CA, Solberg P, Were W, Ekwaru JP, Ransom RL, Weidle PJ, et al. Asymptomatic serum cryptococcal antigenemia and early mortality during antiretroviral therapy in rural Uganda. Tropical medicine & international health. 2007;12(8):929-35.

4. French N, Gray K, Watera C, Nakiyingi J, Lugada E, Moore M, et al. Cryptococcal infection in a cohort of HIV-1-infected Ugandan adults. AIDS. [Research Support, Non-U.S. Gov't]. 2002 May 3;16(7):1031-8.

5. Meya DB, Manabe YC, Castelnuovo B, Cook BA, Elbireer AM, Kambugu A, et al. Cost-effectiveness of serum cryptococcal antigen screening to prevent deaths among HIV-infected persons with a CD4+ cell count < or = 100 cells/microL who start HIV therapy in resource-limited settings. Clin Infect Dis. [Research Support, N.I.H., Extramural Research Support, Non-U.S. Gov't]. 2010 Aug 15;51(4):448-55.

6. Rajasingham R, Meya DB, Boulware DR. Integrating cryptococcal antigen screening and pre-emptive treatment into routine HIV care. Journal of acquired immune deficiency syndromes. [Research Support, N.I.H., Extramural Review]. 2012 Apr 15;59(5):e85-91.

7. Mfinanga S, Chanda D, Kivuyo SL, Guinness L, Bottomley C, Simms V, et al. Cryptococcal meningitis screening and community-based early adherence support in people with advanced HIV infection starting antiretroviral therapy in Tanzania and Zambia: an open-label, randomised controlled trial. Lancet. [Multicenter Study Randomized Controlled Trial Research Support, Non-U.S. Gov't]. 2015 May 30;385(9983):2173-82.

8. Faini D, Kalinjuma AV, Katende A, Mbwaji G, Mnzava D, Nyuri A, et al. Laboratory-reflex cryptococcal antigen screening is associated with a survival benefit in Tanzania. Journal of acquired immune deficiency syndromes. 2019 Feb 1;80(2):205-13.

9. World Health Organization. Guidelines for the diagnosis, prevention and management of cryptococcal disease in HIV-infected adults, adolescents and children. Geneva2018 [April 1, 2018]; Available from: http://www.who.int/hiv/pub/guidelines/cryptococcal-disease/en/.

10. Meya DB, Kiragga AN, Nalintya E, Morawski BM, Rajasingham R, Park BJ, et al. Reflexive laboratory-based cryptococcal antigen screening and preemptive fluconazole therapy for cryptococcal antigenemia in HIV-infected individuals with CD4 <100 cells/microL: A stepped-wedge, cluster-randomized trial. Journal of acquired immune deficiency syndromes. 2019 Feb 1;80(2):182-9.

11. Rhein J, Hullsiek KH, Evans EE, Tugume L, Nuwagira E, Ssebambulidde K, et al. Detrimental outcomes of unmasking cryptococcal meningitis with recent ART initiation. Open forum infectious diseases. 2018 Aug;5(8):ofy122.

12. Alemu AS, Kempker RR, Tenna A, Smisston C, Berhe N, Fekade D, et al. High prevalence of cryptococcal antigenemia among HIV-infected patients receiving antiretroviral therapy in Ethiopia. PLoS One. 2013;8(3):e58377.

13. http://library.health.go.ug/publications/service-delivery-diseases-control-prevention-communicable-diseases/hiv AIDS/consolidated.

14. https://vldash.cphluganda.org/.

15. Kiyaga C, Sendagire H, Joseph E, McConnell I, Grosz J, Narayan V, et al. Uganda’s new national laboratory sample transport system: a successful model for improving access to
diagnostic services for early infant HIV diagnosis and other programs. PloS one. 2013;8(11):e78609.

16. Oladele RO, Akanmu AS, Nwosu AO, Ogunsola FT, Richardson MD, Denning DW, editors. Cryptococcal Antigenemia in Nigerian Patients With Advanced Human Immunodeficiency Virus: Influence of Antiretroviral Therapy Adherence. Open forum infectious diseases; 2016: Oxford University Press.

17. Rajasingham R, Meya DB, Greene GS, Jordan A, Nakawuka M, Chiller TM, et al. Evaluation of a national cryptococcal antigen screening program for HIV-infected patients in Uganda: A cost-effectiveness modeling analysis. PLoS One. 2019;14(1):e0210105.

18. Vallabhaneni S, Longley N, Smith M, Smith R, Osler M, Kelly N, et al. Implementation and Operational Research: Evaluation of a Public-Sector, Provider-Initiated Cryptococcal Antigen Screening and Treatment Program, Western Cape, South Africa. Journal of acquired immune deficiency syndromes. 2016 Jun 01;72(2):e37-e42.

19. Vidal JE, Boulware DR. Lateral flow assay for cryptococcal antigen: an important advance to improve the continuum of HIV care and reduce cryptococcal meningitis-related mortality. Revista do Instituto de Medicina Tropical de São Paulo. 2015;57:38-45.

20. Beyene T, Woldeamanuel Y, Asrat D, Ayana G, Boulware DR. Comparison of cryptococcal antigenemia between antiretroviral naive and antiretroviral experienced HIV positive patients at two hospitals in Ethiopia. PLoS One. [Research Support, Non-U.S. Gov't]. 2013;8(10):e75585.

21. Flynn AG, Meya DB, Hullsiek KH, Rhein J, Williams DA, Musubire A, et al. Evolving failures in the delivery of Human Immunodeficiency Virus care: Lessons from a Ugandan meningitis cohort 2006-2016. Open Forum Infect Dis. 2017 Spring;4(2):ofx077.

22. World Health Organization. Guideline for managing advanced HIV disease and the timing for initiating antiretroviral therapy. 2017 [31 July 2017]; Available from: http://www.who.int/hiv/pub/guidelines/advanced-HIV-disease/en/.

23. Kahn JG, Marseille E, Moore D, Bunnell R, Were W, Degerman R, et al. CD4 cell count and viral load monitoring in patients undergoing antiretroviral therapy in Uganda: cost effectiveness study. Bmj. 2011;343:d6884.

24. Jarvis JN, Lawn SD, Vogt M, Bangani N, Wood R, Harrison TS. Screening for cryptococcal antigenemia in patients accessing an antiretroviral treatment program in South Africa. Clin Infect Dis. [Evaluation Studies Research Support, N.I.H., Extramural Research Support, Non-U.S. Gov't]. 2009 Apr 1;48(7):856-62.
**Table 1:** clinical characteristic of participants by plasma CrAg status

| Characteristic                  | CrAg Positive (N = 35) | CrAg Negative (N=1151) | P-value |
|--------------------------------|------------------------|-------------------------|---------|
| Age, years                     |                         |                         |         |
|                                | 35 (28 to 45)          | 36 (30 to 43)          | 0.92    |
| Women                          | 20 (57%)               | 704 (62%)              | 0.55    |
| HIV-1 Viral Load, copies/mL    | 46,400 (17,300 to 56,000) | 12,000 (3,020 to 69,800) | <0.001  |
| HIV-1 Viral Load >5000 copies/mL | 32 (91%)                | 735 (64%)              | 0.001   |
| Duration of HIV Therapy        |                         |                         |         |
| 0.5-1 year                     | 4 (12%)                | 69 (6.7%)              | 0.22    |
| 1-2 years                      | 4 (12%)                | 144 (14%)              | 0.77    |
| 2-5 years                      | 12 (36%)               | 400 (39%)              | 0.80    |
| >5 years                       | 13 (39%)               | 424 (41%)              | 0.86    |

Values are median (IQR) or n (%). P-Value from Wilcoxon rank-sum test for medians and Chi Square test for proportions. Abbreviations: CrAg = Cryptococcal Antigen
Figure 1: Cryptococcal Antigen (CrAg) prevalence by HIV-1 plasma viral load among Ugandans undergoing routine virologic monitoring on HIV therapy.