Oral Abstract Session-12

Wednesday, March 12, 2025

195 - Performance of HIV RNA Screening in the Context of Long-Acting Injectable Cabotegravir in HPTN 084

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Disclosure: Dr Delany-Moretlwe has no financial relationships with ineligible companies to disclose.

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Background



- HPTN 084 demonstrated the effectiveness of long-acting injectable cabotegravir (CAB-LA) compared to daily oral TDF/FTC for PrEP in individuals born female.
- CAB-LA may delay the detection of early HIV infection using conventional diagnostics, leading to the emergence of resistance
- In retrospective analyses in HPTN 083, HIV RNA testing detected HIV infection prior to the emergence of resistance
- HIV RNA testing may not be feasible in many settings
- We evaluated the performance of HIV RNA screening in the HPTN 084 openlabel extension (OLE)

Attributes of a good screening test





When selecting a screening test, there is a need to balance
the benefits of early treatment for those with undetected infection
vs the harm to those that do not need treatment

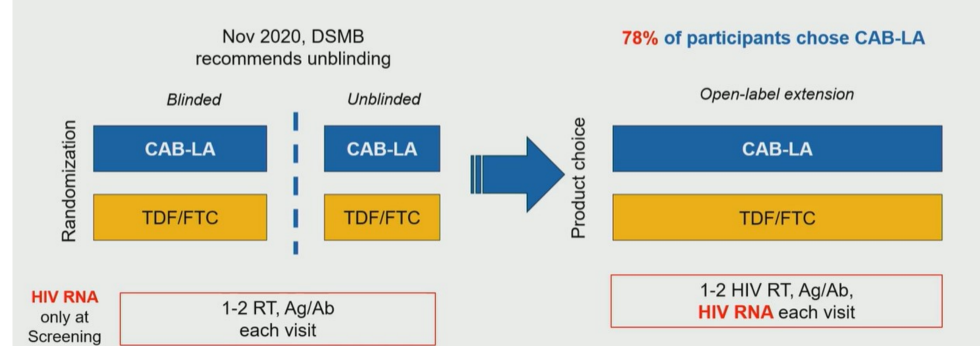
Ideally a screening test should

- Should be capable of detecting infection at an early stage
- accurately identify those with disease i.e. <u>high sensitivity</u>
- Have a <u>high positive predictive value</u> i.e. it accurately predicts the presence of infection
- Results should be easy to interpret with <u>clear cut-off for what constitutes a positive test</u>
- · Should be reasonably priced
- Should be widely available



HPTN 084 study design





Methods



- Site based testing in OLE (all visits)
 - 1-2 HIV rapid tests (RT), antigen/antibody testing (Ag/Ab)
 - Added HIV RNA testing (LLOQ 50 copies/ml)
- Retrospective testing at central laboratory
- Final HIV status adjudicated by external committee
 - Site testing data AND retrospective testing results
- All tests included from OLE entry through Nov 30, 2023
 - Entry into OLE varied by site, starting Jan, 2022
- Estimated the positive predictive value (PPV) and false positive rate (FPR) of isolated positive HIV RNA, and sensitivity of HIV RNA screening with other tests



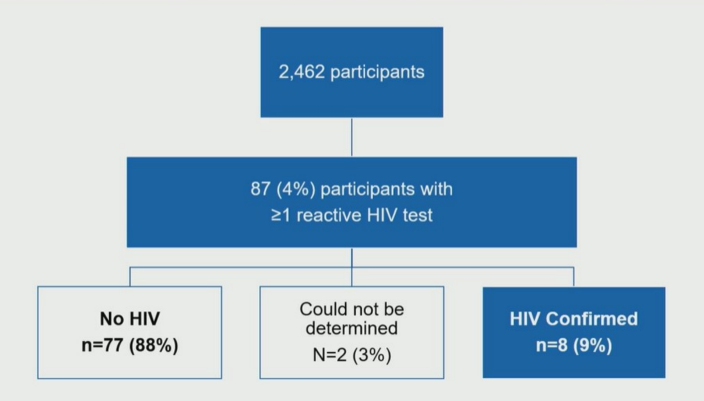
Participant characteristics



	Participants	No. of visits with RNA screening	Person-years
Overall	2,462	24,244	3,229
Country			
Botswana	71	810	108
Kenya	63	733	96
Malawi	157	1,517	200
South Africa	997	9,641	1,329
Eswatini	118	1,155	164
Uganda	419	3,881	509
Zimbabwe	637	6,507	823
PrEP choice			
CAB	1,927	20,262	2,697
TDF/FTC	535	3,982	532

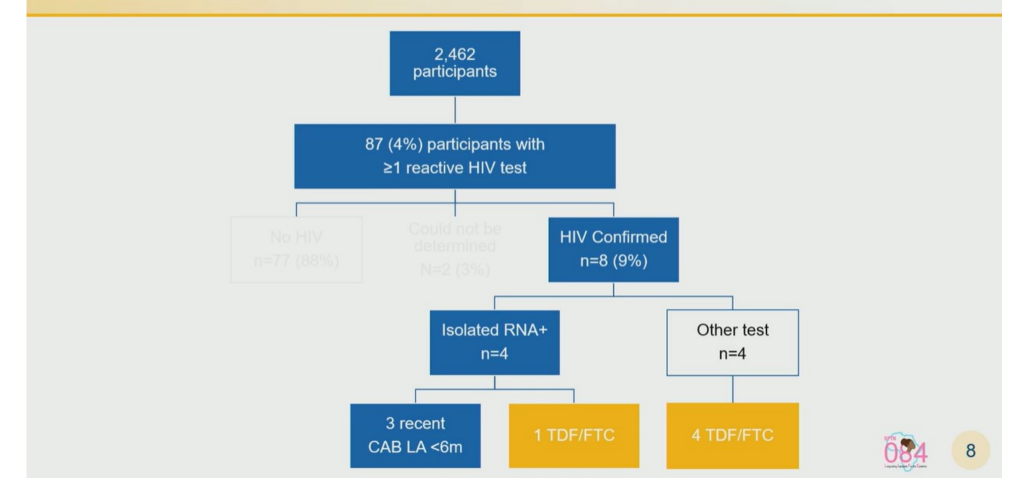
Results – HIV final adjudicated status





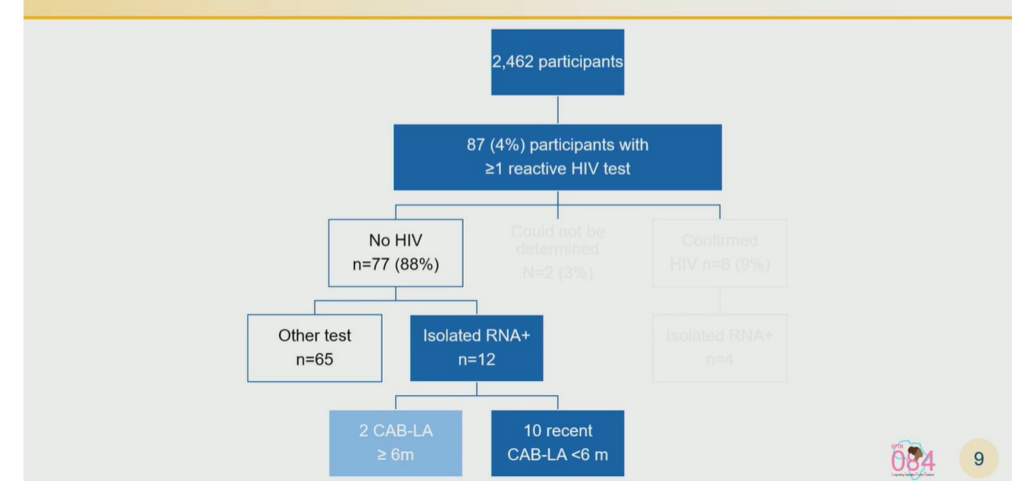
Results – true positive





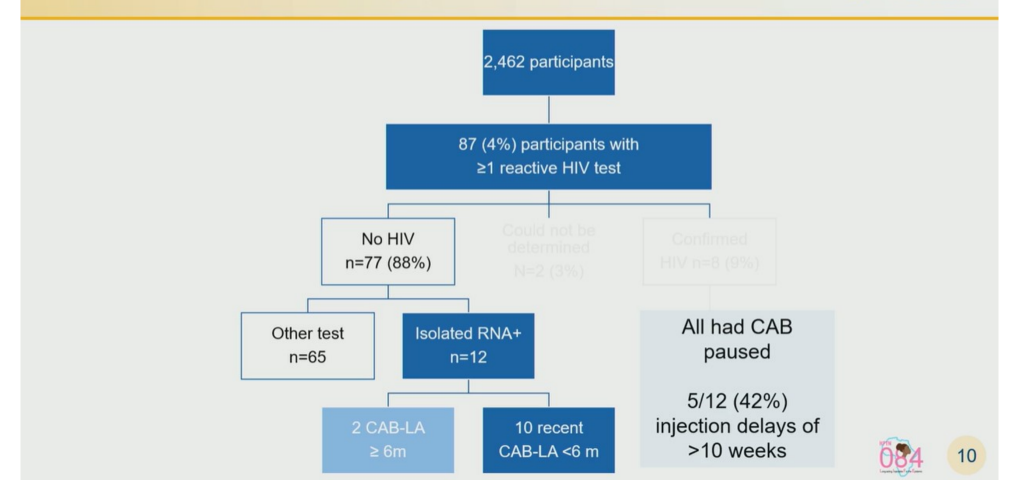
Results – false positive





Results – false positive





HIV RNA performance characteristics



	FPR	PPV	Sensitivity*
	(95% CI)	(95%)	(95% CI)
Overall	75%	25%	62.5%
	(47.6%, 92.7%)	(7.3%, 52.4%)	(24.5%, 91,5%)
CAB-LA use < 6 m	76.9%	23.1%	100.0%
	(46.2%, 95.0%)	(5.0%, 53.8%)	(29.2%, 100.0%)
CAB-LA use ≥ 6m	100% (15.8%, 100.0%)	0% (0%, 84.2%)	0%



^{*}Sensitivity is based on HIV RNA with other screening tests

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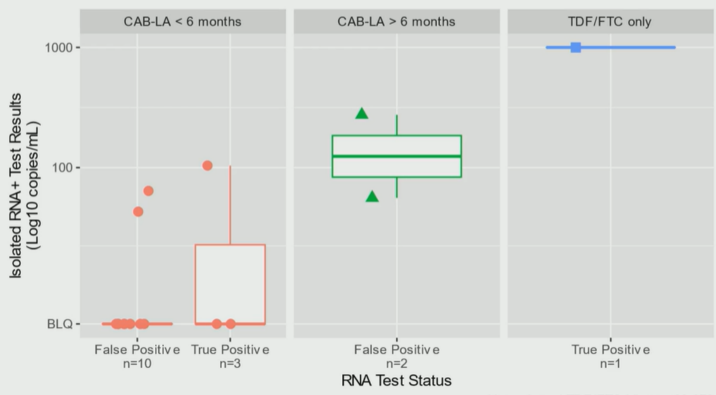


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HIV viral load at isolated HIV RNA positive cases





Note: Actual TDF/FTC VL was 93,873



Conclusions



- Single isolated HIV RNA tests performed poorly for detecting HIV infections in the context of CAB-LA PrEP use.
 - Able to detect early infection,
 - But insufficient accuracy (low sensitivity and specificity)
 - Difficult to distinguish true from false positives based on viral load
- Although infrequent, 75% of isolated positive HIV RNA tests were false positive
 - potential for negative clinical consequences, including prolonged PrEP interruptions.
 - High CAB-LA effectiveness in this population and subsequent low prevalence of true infection may explain the low PPV for HIV RNA screening.
- Future HIV testing algorithm guidelines should carefully consider the costs and risks in addition to any benefits of HIV RNA screening, particularly in resource-constrained settings.

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HIV Prevention Trials Network

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- Laboratory Centre (Johns Hopkins)
- Statistical Center for HIV/AIDS Research and Prevention, Fred Hutchison Cancer Research Center
- HPTN Leadership

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... and our study participants!











