



Ricky K. Hsu^{1,2}, Michael Sension³, Brooke Levis⁴, Jennifer S. Fusco⁴, Carl Millner⁵, Gayathri Sridhar⁶, Vani Vannappagari⁶, Kimberley Brown⁶, Jean Van Wyk⁷, Michael Wohlfeiler⁸, Gregory P. Fusco⁴

¹AIDS Healthcare Foundation, New York, NY, USA; ²NYU Langone Health, New York, NY, US; ³CAN Community Health, Fort Lauderdale, FL, USA; ⁴Epididian, Raleigh, NC, USA; ⁵AIDS Healthcare Foundation, Los Angeles, CA, USA; ⁶ViiV Healthcare, Durham, NC, USA; ⁷ViiV Healthcare, London, England, UK; ⁸AIDS Healthcare Foundation, Miami, FL, USA



Background

CAB+RPV LA

- The first and currently only complete LA ART regimen in the US¹
 - Injections 6 times per year
 - Indicated for ART-experienced individuals with VL <50 copies/mL
 - Nucleoside sparing

BIC/FTC/TAF

- The most common ART regimen in the US-based OPERA Cohort²
 - Single tablet taken orally every day
 - Indicated for ART-naïve or experienced individuals regardless of viral load at initiation
 - Covers chronic hepatitis B

Objective

To assess real world effectiveness after a switch to CAB+RPV LA Q2M injections versus a daily single-tablet oral regimen, BIC/FTC/TAF

Methods

OPERA Cohort

- Prospectively captured, routine clinical data from electronic health records in the US (101 clinics, 23 US states/territories), representing ~14% of people with HIV in the US

Inclusion Criteria

- Adults with HIV
- Treatment-experienced
- Virologically suppressed (VL <50 c/mL)
- Switched to CAB+RPV LA Q2M or BIC/FTC/TAF for the first time between 01FEB2022 and 31MAR2025

Censoring Criteria

- Discontinuation of regimen of interest
- End of study analysis (30SEPT2025)
- Loss to follow-up (no clinic contact for 12 months)
- Death

Outcomes

- Maintenance of virologic suppression over follow up
 - Last VL <50 c/mL
 - All VLs <50 c/mL
- Confirmed virologic failure (CVF): 2 consecutive VL ≥200 copies/mL or 1 VL ≥200 copies/mL followed by discontinuation within 4 months
 - Following regimen
 - Ever re-suppressed to VL <50 c/mL

Statistical Analysis

- Association between regimen and CVF: multivariable logistic regression
 - Adjusted for age, sex, ethnicity, US geographic region, injection drug use, AIDS-defining events, hepatitis C ever, CD4 cell count, comorbid conditions ever, and core agent of prior regimen
- Multiple imputation by chained equations (MICE) for missing data on ethnicity (N = 298) and baseline CD4 cell count (N = 33)
 - Included age, sex, ethnicity, US geographic region, injection drug use, AIDS-defining events, hepatitis C ever, CD4 cell count, comorbid conditions ever, core agent of prior regimen, clinic, race, BMI, VACS score, and prior core agent class
 - Analyses were updated since the abstract submission to exclude individuals with past exposure to regimens of interest

Results

Table 1. Baseline characteristics among individuals switching to CAB+RPV LA Q2M vs. BIC/FTC/TAF

	CAB+RPV LA Q2M (N = 4,090)	BIC/FTC/TAF (N = 2,699)
Age, median years (IQR)	38 (32, 49)	47 (36, 58)
50-64, n (%)	807 (20)	961 (36)
≥65, n (%)	140 (3)	244 (9)
Female sex, n (%)	569 (14)	448 (17)
Black race, n (%) ^a	1,739 (42)	1,112 (41)
Hispanic ethnicity, n (%) ^b	1,372 (34)	725 (27)
Injection drug use, n (%)	231 (6)	289 (8)
Care in Southern US, n (%)	2,166 (53)	1,723 (64)
AIDS-defining events (ever), n (%)	177 (4)	79 (3)
Any comorbid conditions, n (%)	3,065 (75)	2,229 (83)
CD4, median cells/μL (IQR) ^c	714 (532, 918)	694 (506, 914)
Prior core agent class, n (%)		
INSTI	3,433 (84)	1,847 (68)
PI	153 (4)	310 (12)
NNRTI	242 (6)	382 (14)
2 or more core agents	261 (6)	159 (6)
Other	≤5	≤5
Months of prior regimen, median (IQR)	20 (9, 44)	41 (12, 70)

^a Missing race: N = 322
^b Missing ethnicity: N = 298
^c Missing CD4 count: N = 33

Table 2. Persistence

	CAB+RPV LA Q2M (N = 4,090)	BIC/FTC/TAF (N = 2,699)
Months on regimen until first censoring ^a , median (IQR)	15 (8, 25)	22 (14, 31)
On regimen at study end (regardless of any discontinuations), n (%)	2,997 (73)	2,175 (81)
Cumulative months on regimen, median (IQR)	18 (11, 28)	23 (15, 33)

^a Censoring at the first of regimen discontinuation, study end (30SEPT2025), loss to follow-up (no clinic contact for 12 months), or death

Table 3. Virologic re-suppression after CVF

	CAB+RPV LA Q2M (N = 46)	BIC/FTC/TAF (N = 46)
≥1 VL after CVF, n (%)	22 (48)	29 (63)
Re-suppressed to VL <50 c/mL, n (%)	15 (68)	20 (69)
Months to re-suppression, median (IQR)	2 (1, 3)	4 (3, 7)

Figure 1. Maintenance of virologic suppression among individuals with ≥1 follow-up VL

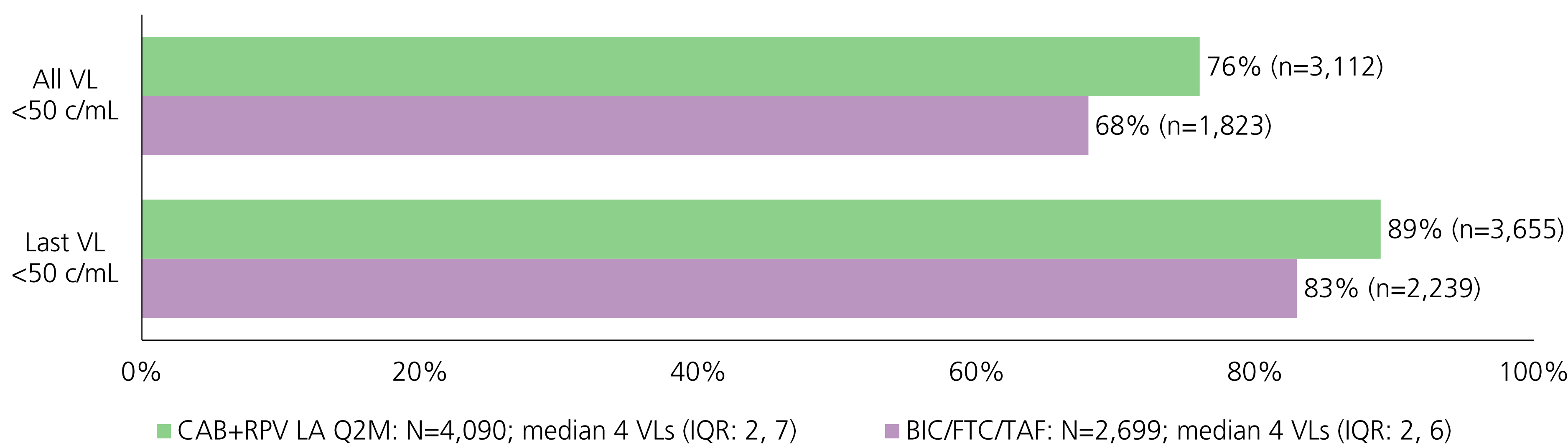


Figure 2. Confirmed virologic failure among individuals with ≥1 follow-up VL

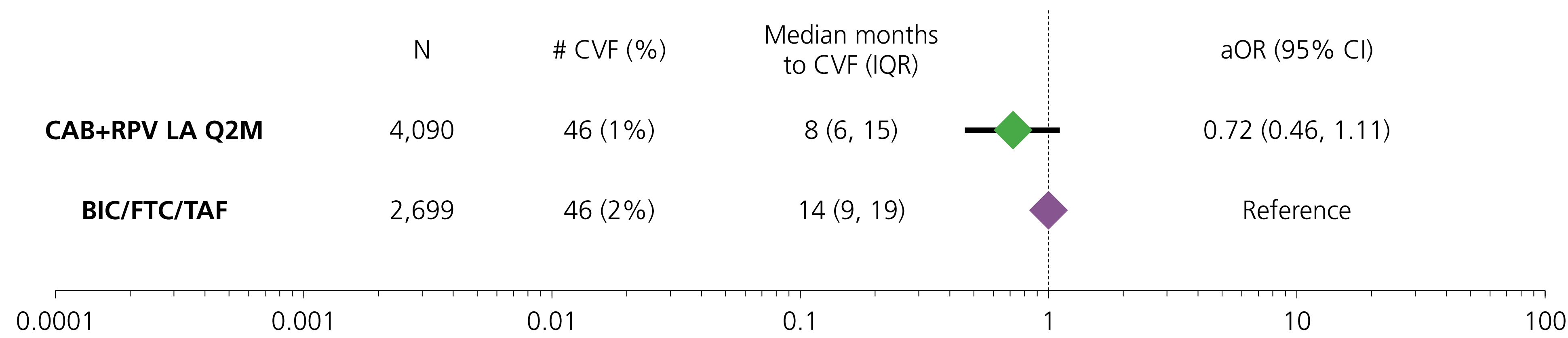
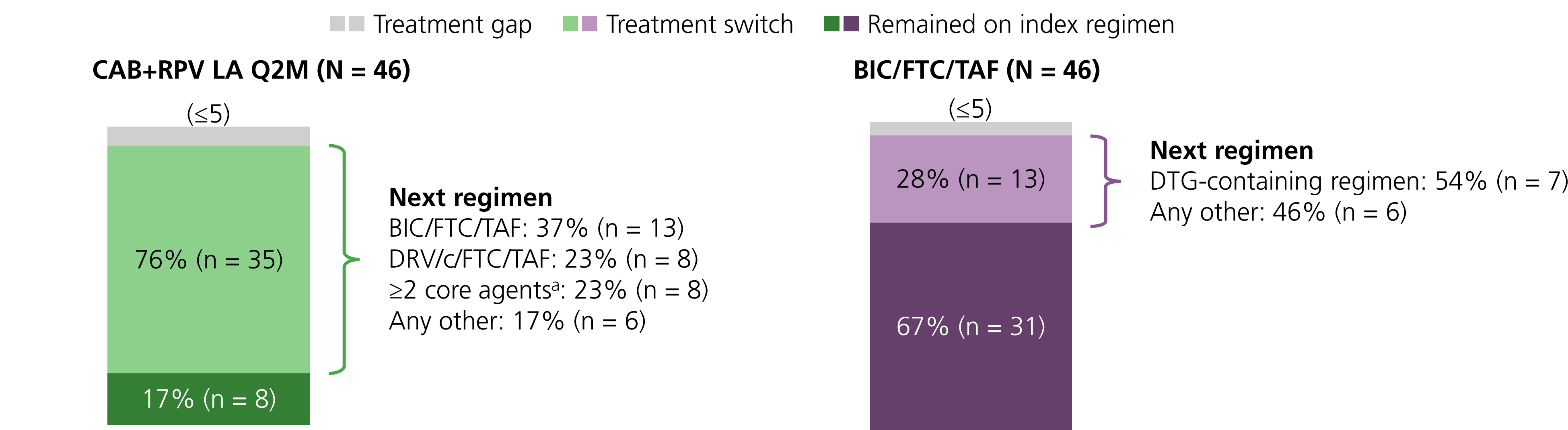


Figure 3. Regimen following confirmed virologic failure



^a Excluding 2 drug regimens (CAB+RPV LA, DTG/RPV)

Abbreviations

AIDS, Acquired immunodeficiency syndrome; **aOR**, adjusted odds ratio; **ART**, antiretroviral therapy; **BIC/FTC/TAF**, bictegravir/emtricitabine/tenofovir alafenamide; **BMI**, body mass index; **CAB+RPV**, cabotegravir + rilpivirine; **CVF**, confirmed virologic failure; **DRV/c/FTC/TAF**, darunavir/cobicistat /emtricitabine/tenofovir alafenamide; **DTF**, dolutegravir; **HIV**, human immunodeficiency virus; **INSTI**, integrase strand transfer inhibitor; **IQR**, interquartile range; **LA**, long-acting; **c/mL**, copies per milliliter; **μL**, microliter; **Q2M**, every 2 months dosing schedule; **US**, United States; **VACS**, Veterans Aging Cohort Study Index; **VL**, viral load.

Discussion

Findings

- Individuals who switched to CAB+RPV LA Q2M differed from those switching to BIC/FTC/TAF (**Table 1**)
 - CAB+RPV LA Q2M users were younger (23% vs. 45% ≥50 years old), Hispanic ethnicity (34% vs. 27%) and more likely to have been on an INSTI prior to the switch (84% vs. 68%)
 - BIC/FTC/TAF users were more likely to have comorbid conditions (83% vs. 75%) and be cared for in the Southern US (64% vs. 53%)
- Median time on CAB+RPV LA Q2M was 15 months vs. 22 months on BIC/FTC/TAF before censoring (**Table 2**)
- CAB+RPV LA Q2M users were more likely than BIC/FTC/TAF users to maintain virologic suppression at last VL (89% vs 83%) and all VLs (76% vs. 68%) (**Fig 1**)
- CVF risk was lower in the CAB+RPV LA Q2M group than BIC/FTC/TAF group (46 [1%] vs. 46 [2%]), but after adjusting for baseline characteristics, the difference was not statistically significant (adjusted odds ratio [95% CI] = 0.72 [0.46, 1.11]) (**Fig 2**)
- Of those who experienced CVF, most of the CAB+RPV LA Q2M group (76%), switched to a new regimen, while most of the BIC/FTC/TAF group (67%) did not (**Fig 3**)
- Of 22 (CAB+RPV LA Q2M) and 29 (BIC/FTC/TAF) individuals with VLs available post-CVF, 68% and 69%, respectively, resuppressed to <50 copies/mL before study end. (**Table 3**)

Strengths

- Real-world clinical experience of CAB+RPV LA and BIC/FTC/TAF
- With 6,794 individuals followed, this comparison is a robust evaluation of the regimens
- Diverse population from 24 US states & territories
- Power to control for many covariates

Limitations

- Electronic health records data may vary in quality across practices and providers; therefore, measurement error is possible
- Accuracy of exposures were not equal
 - CAB+RPV LA injections are directly observed in the clinic
 - BIC/FTC/TAF exposure is captured via prescriptions
- Selection bias may have been introduced when individuals lacking complete ART histories were excluded to avoid unobserved exposure time
- Oral bridging during CAB+RPV LA use is not well documented and may result in over-estimation of discontinuation

References

1. CABENUVA prescribing information; https://www.accessdata.fda.gov/drugsatfda_docs/label/2025/212888Orig1s017lbl.pdf accessed: 19June2026
2. BIKTARVY prescribing information; https://www.accessdata.fda.gov/drugsatfda_docs/label/2026/210251Orig1s026lbl.pdf accessed: 19June2026

Key Findings

- In this real-world analysis of nearly 6,800 virologically suppressed individuals, switching to CAB+RPV LA Q2M consistently maintained high levels of virologic suppression comparable to switching to daily BIC/FTC/TAF
- CVF risk and re-suppression after CVF did not differ between the groups
- These findings support that CAB+RPV LA Q2M is as effective as daily oral therapy in maintaining virologic suppression

Acknowledgements

This research would not be possible without the generosity of people living with HIV and their OPERA caregivers. Additionally, we are grateful for the following individuals: Kelly Oh (SAS programming), Bill Johnson (QA), Bernie Stooks (data management), Lisa Lutz & Nicole Shaw (data management/quality), Judy Johnson & Quateka Cochran (clinical data classification), and Laurence Brunet (poster design)

Support

This research was supported by ViiV Healthcare

