

NON-INTERVENTIONAL STUDY PROTOCOL

STUDY DETAILS

UNIQUE IDENTIFIER	224047
TITLE	CABFIDENCE CAB+RPV LA Study: Global Real-World Evidence Cohort Study to evaluate Utilization, Clinical and Patient- Reported Outcomes among CAB+RPV LA users
STUDY ACCOUNTABLE PERSON	PPD
SCIENTIFIC LEAD	PPD
CONTRIBUTING AUTHORS	PPD
ASSET ID	GSK4071318
GSK or ViiV ASSET	Cabotegravir + Rilpivirine LA
EFFECTIVE DATE	30 April 2026
INDICATION	HIV-1

DATA COLLECTION TYPE	PRIMARY AND SECONDARY
SAFETY OBJECTIVE	YES
ASSET INVOLVEMENT	YES
TSS/PASS ASSESSMENT PERFORMED	Yes
STUDY CLASSIFICATION	TSS, Voluntary PASS
EVALUATING A PRODUCT (TIER TYPE)	Tier 1
REGULATORY COMMITMENT	No

TITLE PAGE

Study ID: 224047

Division: Global Medical

Information Type: Non-Interventional Study Protocol

Title: CABFIDENCE CAB+RPV LA Study: Global Real-World Evidence Cohort Study to evaluate Utilization, Clinical, and Patient-Reported Outcomes among CAB+RPV LA users

Effective Date: 30 April 2026

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STUDY INFORMATION

Title	CABFIDENCE CAB+RPV LA Study: Global Real-World Evidence Cohort Study to evaluate Utilization, Clinical and Patient-Reported Outcomes among CAB+RPV LA users
Protocol version identifier	Amendment 2
Date of last version of protocol	19 October 2025
EU PAS (ENCEPP) register number	EUPAS1000000616
Active substance	Cabotegravir Rilpivirine
Medicinal product	VOCABRIA® 30 mg film-coated tablets Edurant® 25 mg film-coated tablets Cabenuva VOCABRIA® 400 mg/ 600 mg prolonged-release suspension for injection REKAMBYS® 600 mg / 900 mg prolonged-release suspension for injection
Product reference	NDA 212887 VOCABRIA (cabotegravir) tablets NDA 202022 EDURANT (rilpivirine) tablets NDA 212888 CABENUVA (cabotegravir extended-release injectable suspension; rilpivirine extended-release injectable suspension), copackaged, for intramuscular use
Procedure number	NA
Marketing authorisation holder(s)	ViiV Healthcare
Joint PASS	No

Research question and objectives	Using global real-world clinical data, this study will describe utilization, clinical and patient-reported outcomes among people with HIV on CAB+RPV LA regimen
Country(-ies) of study	Anticipated countries include United Kingdom, Italy, Portugal, France, Spain, Japan, Australia, and the United States
Author	PPD [REDACTED]

MARKETING AUTHORISATION HOLDER(S)

Marketing authorisation holder(s)	ViiV Healthcare 410 Blackwell St. Durham, NC 27701
MAH contact person	PPD [REDACTED]

REVISION CHRONOLOGY

Date	Version	Change(s) since last version
01 May 2025	Original	N/A
19 October 2025	Amendment 1	Study design/study population and setting: - Updated inclusion criteria for prospective participants (new initiators) that study enrollment should occur prior to the first CAB+RPV LA injection, either during the OOLI phase, if applicable, or directly before their first CAB+RPV LA injection for those on the DTI pathway - Clarified the study design and population for retrospective and prospective participants - Anticipated country scope revised to include Canada, Japan, South Korea, and Taiwan. Data sources: - Added that the baseline survey will be administered prior to the first CAB+RPV LA injection Primary objective: - Updated adherence outcome definition to include both early as well as delayed injections. - Added clarifying text to outcome definitions including persistence and effectiveness Secondary objective: - Added additional subgroups for BMI(<30, ≥30-40, ≥40 kg/m²) and sex/race category - Added treatment satisfaction and treatment preference to be collected as part of the participant questionnaire - Added clarification that for data collected beyond the two-year follow-up period,

		outcomes will be analyzed at the extended time points Outcome definitions: • Updated adherence outcome definition to include both early as well as delayed injections. • Added clarifying text to outcome definitions including persistence, discontinuation and effectiveness • Added references to the HIVTSQs, HIVTSQc and treatment preference to be collected as part of the participant questionnaire • Added assessment of quality of life and treatment related worries and stigma at discontinuation in addition to baseline and month 12 Confounder and effect modifiers: • Added additional variables to be collected if available (i.e., CD4/CD8 ratio, CD4 nadir prior to initiation of CAB+RPV LA, Type of viral load assay for testing, VACS Mortality Index, ASCVD risk score, Framingham Risk Score, additional comorbidities for mental health disorders Study size: • Sample size increased from 1,000 to up to 2,000 Data management: • Clarified that baseline survey will be completed at the site and follow-up survey either on participant's own device or tablet provided to sites by RTI-HS • Added a sentence to allow comparison of study data to data from other protocols or studies Data analysis: • Added clarification in the main analytical approach and secondary analysis Ethics approval and subject consent: • Removed text outlining country specific ethics and regulatory requirements; this information will be captured in a separate document
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		Milestones: Simplified the milestones table Updated ‘individuals’ to ‘participants’ throughout the protocol as appropriate
30 April 2026	Amendment 2	Rationale and background - Added description of the CABFIDENCE platform program with the shared data lake and use of a master protocol of which CABFIDENCE CAB+RPV LA study would be a sub-protocol Study design/study population and setting: - Clarified that the date of first CAB+RPV injection is considered index date - Anticipated country scope revised to remove Canada, Germany, South Korea, and Taiwan. - Added exclusion criteria for excluding individuals with prior resistance if data is available and individuals who are unable to complete the consent process Outcome definitions: - Added 6 month data collection of select patient-reported outcomes Data sources: - Added 6 month data collection of participant surveys and participant medical information via site-completed eCRF: Data management: - Language updated to be inclusive of newly added 6 month data collection timepoint - Added platform program language Ethical approval and subject consent - Added platform program language Management and reporting of adverse events - Added some clarifications to this section Milestones: - Updated the milestones table

SUMMARY OF CHANGES

Amendment 1:

“CABFIDENCE: Global Real-World Evidence Cohort Study to evaluate Utilization, Clinical and Patient-Reported Outcomes among CAB+RPV LA users” (Study #224047) is a ViiV-sponsored three-year observational cohort study using data from individual medical records from participating clinical sites and participant-reported data from electronic questionnaires (effective date of original protocol: 01 May 2025).

Changes to study design/ study population and setting:

- Updated inclusion criteria for prospective participants (new initiators) that study enrollment should occur prior to the first CAB+RPV LA injection, either during the OOLI phase, if applicable, or directly before their first CAB+RPV LA injection for those on the DTI pathway
- Clarified the study design and population for retrospective and prospective participants
- Anticipated country scope revised to include Canada, Japan, South Korea, and Taiwan.

Changes in data sources:

- Added that the baseline survey will be administered prior to the first CAB+RPV LA injection

Changes in study objectives:

1. Primary objective:
 - a. Updated adherence outcome definition to include both early as well as delayed injections.
 - b. Added clarifying text to outcome definitions including persistence and effectiveness
2. Secondary objective:
 - a. Added additional subgroups for BMI (<30 , ≥ 30 -40, ≥ 40 kg/m²) and sex/race category
 - b. Added treatment satisfaction and treatment preference to be collected as part of the participant questionnaire
 - c. Added clarification that for data collected beyond the two-year follow-up period, outcomes will be analyzed at the extended time points

Changes in Outcome definitions:

- Updated adherence outcome definition to include both early as well as delayed injections.
- Added clarifying text to outcome definitions including persistence, discontinuation and effectiveness
- Added references to the HIVTSQs, HIVTSQc and treatment preference to be collected as part of the participant questionnaire
- Added assessment of quality of life and treatment related worries and stigma at discontinuation in addition to baseline and month 12

Changes in Confounder and effect modifiers:

- Added additional variables to be collected if available (i.e., CD4/CD8 ratio, CD4 nadir prior to initiation of CAB+RPV LA, Type of viral load assay for testing, VACS Mortality Index, ASCVD risk score, Framingham Risk Score, additional comorbidities for mental health disorders

Changes in study size:

- Sample size increased from 1000 to up to 2,000

Changes in data management:

- Clarified that baseline survey will be completed at the site and follow-up survey either on participant's own device or tablet provided to sites by RTI-HS
- Added a sentence to allow comparison of study data to data from other protocols or studies

Changes in data analysis:

- Added clarification in the main analytical approach and secondary analysis

Changes in Ethics approval and subject consent:

- Removed text outlining country specific ethics and regulatory requirements; this information will be captured in a separate document

Changes in the milestones table:

- Simplified the milestones table

Updated ‘individuals’ to ‘participants’ throughout the protocol as appropriate

Amendment 2:

Changes to background and rationale:

- Added language to describe the CABFIDENCE platform program with the planned shared data lake and use of a master protocol of which CABFIDENCE CAB+RPV LA would be a sub-protocol

Changes to study design/ study population and setting:

- Clarified that the date of first CAB+RPV injection is considered the index date for participants
- Anticipated country scope revised to remove Canada, Germany, South Korea, and Taiwan.
- Added exclusion criteria for excluding individuals with prior resistance if data is available and individuals who are unable to complete the consent process

Changes in data sources:

- Added 6 month data collection of participant surveys and participant medical information via site-completed eCRF

Changes in Outcome definitions:

- Updated patient-reported outcomes to reflect which items will be included in the added 6 month participant survey

Changes in data management:

- Added language to account for the addition of the 6 month data collection timepoint
- Added platform program language

Changes in Ethical approval and subject consent:

- Added platform program language

Changes in Management and reporting of adverse events:

- Added some clarifications to this section

Changes in the milestones table:

- Updated the milestones table

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LIST OF ABBREVIATIONS

ADR	adverse drug reaction
AE	adverse event
ART	antiretroviral therapy
ARV	antiretroviral
BMI	body mass index
CAB	cabotegravir
CAB+RPV LA	cabotegravir + rilpivirine long-acting
CD4	cluster of differentiation 4
CVF	confirmed virologic failure
DTI	Direct to injection
eCRF	electronic case report form
EDC	electronic data capture
EU	European Union
GPP	Good Pharmacoepidemiology Practices
HCRW	Health and Care Research Wales
HIV	human immunodeficiency virus
HIVTSQc	HIV Treatment Satisfaction Questionnaire change version
HIVTSQs	HIV Treatment Satisfaction Questionnaire status version
ICMJE	International Committee of Medical Journal Editors
ICSR	individual case safety reports
ID	identifier
INI	integrase inhibitor
INSTI	integrase strand transfer inhibitor

ISPE	International Society for Pharmacoepidemiology
LA	long-acting
NNRTI	nonnucleoside reverse transcriptase inhibitor
OOLI	Optional oral lead-in
RPV	rilpivirine
RTI-HS	RTI Health Solutions
SAE	serious adverse event
UK	United Kingdom
US	United States
VL	viral load
WHOQOL-HIV BREF	World Health Organization Quality of Life-Human Immunodeficiency Virus BREF

TRADEMARK INFORMATION

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VOCABRIA™	

1. RESPONSIBLE PARTIES

MARKETING AUTHORISATION HOLDER

ViiV Healthcare Company

Sponsor Legal Registered Address:

ViiV Healthcare Company

410 Blackwell St.

Durham, NC 27701

1.1. Sponsor Signatory

Title: CABFIDENCE CAB+RPV LA Study: Global Real-World Evidence Cohort Study to evaluate Utilization, Clinical and Patient-Reported Outcomes among CAB+RPV LA users

Compound Number: GSK4071318

PPD

Gayathri Sridhar
Primary Author/NI Scientific Lead

29 April 2026

Date (DD Month YYYY)

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Vani Vannappagari
Therapy Area Leader/+1 Manager

30 April 2026

Date (DD Month YYYY)

PPD

Nassrin Payvandi
VP & Head, Safety & Pharmacovigilance

30-Apr-2026

Date (DD Month YYYY)

PPD

Jens Ulrich Stegmann
Qualified Person for Pharmacovigilance /Delegate

30-Apr-2026

Date (DD Month YYYY)

PPD

Bryn Jones
GSK APBI Signatory

20th April 2026

Date of Certification
(Date DD Month YYYY)

**I certify that this protocol is in accordance with the
ABPI Code of Practice.**

Note: Not applicable if an eSignature process is used to get the sponsor approval.

1.2 Investigator Protocol Agreement Page

- I confirm agreement to conduct the study in compliance with the protocol.
- I acknowledge that I am responsible for overall study conduct. I agree to personally conduct or supervise the described study.
- I agree to ensure that all associates, colleagues, and employees assisting in the conduct of the study are informed about their obligations. Mechanisms are in place to ensure that site staff receive the appropriate information throughout the study.

Investigator Name:

David Richardson

PPD



Investigator Signature

Date (DD Month YYYY)

2. SYNOPSIS

Title CABFIDENCE CAB+RPV LA Study: Global Real-World Evidence Cohort Study to evaluate Utilization, Clinical and Patient-Reported Outcomes among CAB+RPV LA users

Rationale and Background

Cabotegravir + rilpivirine long-acting (CAB+RPV LA) injectable is the first complete long-acting antiretroviral (ARV) regimen approved for treatment of human immunodeficiency virus type 1 (HIV-1). It is indicated for people with HIV (PWH) who are virologically suppressed (HIV-1 RNA viral load [VL] < 50 copies per mL) on a stable ARV regimen, with no history of treatment failure and no known or suspected resistance to CAB or RPV (in the US)/agents of the nonnucleoside reverse transcriptase inhibitor (NNRTI) and integrase inhibitors (INI) class (in Europe). While clinical studies have established the efficacy, safety, and tolerability of CAB+RPV LA, this study, as part of the CABFIDENCE platform program, will assess the global utilization patterns, adherence, persistence, discontinuation, virologic effectiveness, and patient-reported outcomes of this regimen in a real-world clinical setting.

Research Question and Objective(s)

The overall objective of this global study is to describe characteristics, assess clinical outcomes, and patient-reported outcomes among people with HIV who receive CAB+RPV LA regimen.

Primary Objectives:

1. To describe baseline demographics, clinical characteristics and patterns of use among participants receiving CAB+RPV LA
2. To assess the persistence, discontinuation, and adherence among participants receiving CAB+RPV LA
3. To assess virologic effectiveness among participants receiving CAB+RPV LA
4. To assess factors associated with confirmed virologic failure (CVF) among participants receiving CAB+RPV LA regimen

Secondary Objectives:

1. Subgroup analyses by body mass index (BMI) (< 30, ≥ 30-<40 and ≥40 kg/m²) at initiation, age at initiation (18-49, 50-64, ≥ 65 years), sex assigned at birth (female, male), race/ethnicity, sex/race, country (if sample sizes allow), to assess characteristics, adherence, persistence, discontinuation and virologic effectiveness
2. To estimate the frequency of documented injection site reactions and hypersensitivity reactions among participants receiving CAB+RPV LA injections

3. To describe HIV resistance and proviral DNA testing at the time of discontinuation/CVF, history of resistance and proviral DNA testing prior to discontinuation/CVF, and the ART regimen and virologic outcomes after discontinuation/CVF
4. To describe reasons for initiating CAB+RPV LA, health-related quality of life, treatment satisfaction, treatment preference, treatment-related stigma, and benefits, and reasons for missed injections and discontinuation among prospective participants (new initiators) of CAB+RPV LA
5. For data collected beyond the two-year follow-up period, outcomes will be analyzed at the extended time points

Study Design

This is a non-interventional prospective observational cohort study using data from individual medical records from participating clinical sites and participant-reported data from electronic questionnaires. The study population will include treatment-experienced virologically suppressed (VL < 50 copies/mL) individuals who are 18 years or older and received at least one CAB+RPV LA injection with every-2 month dosing schedule, .

Retrospective participants will include individuals who initiated CAB+RPV LA from January 2023 and will be followed for up to two years. Prospective participants will include individuals enrolled from November 2025 until study size is reached and will be followed for a minimum of two years, unless they withdraw from the study, are lost to follow-up, or pass away. Participants who discontinue and switch to a subsequent regimen or who experience CVF will be followed-up for a minimum of 6 months and may be extended until viral load assessment is completed, with a maximum duration of 12 months. Study end is the date when the last participant completes the required follow-up assessment (including any required post-discontinuation/CVF follow-up).

Population

The study population will consist of people living with HIV who are aged 18 years and older on CAB+RPV LA from participating clinical sites. For the retrospective participants, those who have discontinued the treatment for any reason will be included and individuals who are deceased or lost to follow up will not be included. Anticipated countries include United Kingdom, Italy, Portugal, France, Spain, Japan, Australia, and the United States.

Inclusion Criteria:

- Diagnosed with HIV-1
- Aged ≥ 18 years at time of CAB+RPV LA injection
- Treatment experienced
- VL < 50 copies/mL at initiation

- Initiated every 2 month dosing CAB+RPV LA
 - For prospective participants, study enrollment should occur prior to the first CAB+RPV LA injection, either during the optional oral lead-in (OOLI) phase, if applicable, or directly before their first CAB+RPV LA injection for those on the direct-to-injection (DTI) pathway
 - For retrospective participants, individuals who initiated CAB+RPV LA from January 2023 will be enrolled

Exclusion Criteria:

- Received CAB+RPV LA regimen as part of a randomized, controlled clinical trial
- Treatment naïve
- VL $\geq$ 50 copies/mL at initiation
- Individuals with documented prior virologic failure to agents of the NNRTI or INSTI class (if data is available)
- Individuals with documented present or past resistance to agents of the NNRTI or INSTI class (if data is available)
- Individuals who are unable to complete the consent process

Variables

Exposure:

Exposure is defined as receiving CAB+RPV LA injectable regimen

Outcomes:

- Persistence
- Discontinuation
- Adherence
- Virologic effectiveness
- Resistance
- ART regimen after discontinuation /CVF
- Health-related quality of life
- Treatment satisfaction
- Treatment preference
- Treatment-related stigma, and benefits
- Reasons for initiating CAB+RPV LA
- Reasons for discontinuing CAB+RPV LA
- Reasons for missed injection(s)

Potential Confounders to be Evaluated:

Baseline: age, sex, gender, race, ethnicity, region, marital status, risk of HIV transmission,

payer type, cluster of differentiation 4 (CD4) cell count, VL, prior regimen, years since HIV diagnosis and antiretroviral therapy (ART) initiation, weight, BMI, co-infections, comorbidities, and concomitant medications

Data Sources

Data for this study will be obtained via the following sources:

- **Participant medical information via site-completed electronic case report form (eCRF):** The investigator or authorized medical staff will record clinical and treatment data from participants' medical records into an eCRF at enrollment, at month 6, and a minimum of two annual follow-ups (unless participant discontinued CAB+RPV LA) and at study end.
- **Participant surveys:** Prospective participants of CAB+RPV LA will be asked to complete an online survey at enrollment, at month 6, at month 12, and if they discontinue CAB+RPV LA. The baseline survey will be administered prior to the first CAB+RPV LA injection. Participants will be asked to complete the World Health Organization Quality of Life-HIV BREF (WHOQOL-HIV BREF) HIVTSQs, and the HIVTSQc (O'Connell, 2012; Woodcock and Bradley, 2001; Woodcock and Bradley, 2006) and several bespoke questionnaires developed for this study. Appropriate agreements with the copyright owners will be in place for use in this study. It is estimated that the participant surveys will take approximately 20 minutes to complete.

Study Size

The study aims to enroll up to approximately 2,000 participants. Among the participants, it is estimated that 250 participants are already receiving CAB+RPV LA (retrospective participants), and 1,750 participants will be newly initiating CAB+RPV LA (prospective participants). A participant will be considered as a prospective participant of CAB+RPV LA if study enrollment occurred prior to the first CAB+RPV LA injection, either during the OOLI phase, if applicable, or directly before their first CAB+RPV LA injection for those on the DTI pathway.

Data Analysis

Baseline characteristics and outcomes will be described using counts and relative frequencies for categorical variables. Continuous variables will be summarized using summary statistics (means, standard deviations, medians, minimums, maximums, and quartiles 1 and 3). For event outcomes assessed at any point during follow-up, incidence rates will be estimated using unadjusted Poisson regression, accounting for person-time since index (i.e., first CAB+RPV LA injection) where appropriate. Additional analytic approaches may also be considered (e.g., Kaplan-Meier estimator/curve) and multivariable models such as logistic regression as applicable. Analyses will be stratified by BMI ($< 30 \text{ kg/m}^2$ vs. $\geq 30 \text{ kg/m}^2$), age (18-49, 50-64, ≥ 65 years), sex assigned at birth (female, male) and race and ethnicity, and country, if sample sizes allow, at initiation of CAB+RPV LA regimen. Factors associated with CVF will be assessed using a logistic regression model. The number and proportion of participants with history of resistance to INSTI or NNRTI classes and INSTI and/or NNRTI RAMs at discontinuation/CVF with

results will be described. Subsequent regimen after discontinuation/CVF and virologic outcomes will be assessed. Participant surveys will be summarized descriptively for item and summary score (where applicable). The WHOQOL-HIV BREF, HIVTSQs, and HIVTSQc will be scored according to instrument scoring guidelines and derived scores will be summarized descriptively. All analyses will be performed on the observed data only and no missing values will be imputed.

Milestones

The study will start after the protocol is approved, with data collection until follow-up is completed. The final study report will be completed 12 months after end of data collection.

3. AMENDMENTS AND UPDATES

Amendment or update no	Date	Section of study protocol	Amendment or update	Reason
1	19 Oct 2025	Study objectives, study design, study population and setting, outcome definitions, confounders and effect modifiers, data sources, study size, data management, data analysis, ethics approval and subject consent and milestones	Study design/study population and setting: - Updated inclusion criteria for prospective participants (new initiators) that study enrollment should occur prior to the first CAB+RPV LA injection, either during the OOLI phase, if applicable, or directly before their first CAB+RPV LA injection for those on the DTI pathway - Clarified the study design and population for retrospective and prospective participants - Anticipated country scope revised to include Canada, Japan, South Korea, and Taiwan. Data sources: - Added that the baseline survey will be administered prior to the first CAB+RPV LA injection Primary objectives: - Updated adherence outcome definition to include both early as well as delayed injections. - Added clarifying text to outcome definitions	Protocol amended to reflect change in inclusion criteria for prospective participants to ensure baseline survey data is administered prior to first CAB+RPV LA injection, clarify study design and population for retrospective and prospective participants, to expand data collected to support primary and secondary objectives, to ensure clarity of outcome definitions, and to account for additional countries and increased estimated sample size

			including persistence and effectiveness Secondary objectives: - Added additional subgroups for BMI (<30, ≥ 30-40, ≥ 40 kg/m²) and sex/race category - Added treatment satisfaction and treatment preference to be collected as part of the participant questionnaire - Added clarification that for data collected beyond the two-year follow-up period, outcomes will be analyzed at the extended time points Outcome definitions: - Updated adherence outcome definition to include both early as well as delayed injections. - Added clarifying text to outcome definitions including persistence, discontinuation and effectiveness - Added references to the HIVTSQs, HIVTSQc and treatment preference to be collected as part of the participant questionnaire - Added assessment of quality of life and treatment related worries and stigma at discontinuation in addition to baseline and month 12	
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			Confounder and effect modifiers: - Added additional variables to be collected if available (i.e., CD4/CD8 ratio, CD4 nadir prior to initiation of CAB+RPV LA, Type of viral load assay for testing, VACS Mortality Index, ASCVD risk score, Framingham Risk Score, additional comorbidities for mental health disorders Study size: - Sample size increased from 1000 to up to 2,000 Data management: - Clarified that baseline survey will be completed at the site and follow-up survey either on participant's own device or tablet provided to sites by RTI-HS - Added a sentence to allow comparison of study data to data from other protocols or studies Data analysis: - Added clarification in the main analytical approach and secondary analysis Ethics approval and subject consent: - Removed text outlining country specific ethics and regulatory requirements; this information will be captured in a separate document	
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			Milestones: Simplified the milestones table Updated ‘individuals’ to ‘participants’ throughout the protocol as appropriate	
2	30 Apr 2026	Rationale and background, study design/study population and setting, outcome definitions, data sources, data management	Rationale and background - Added description of the CABFIDENCE platform program with the shared data lake and use of a master protocol of which CABFIDENCE would be a sub-protocol Study design/study population and setting: - Clarified that the date of first CAB+RPV injection is considered index date - Anticipated country scope revised to remove Canada, Germany, South Korea, and Taiwan. - Added exclusion criteria for excluding individuals with prior resistance if data is available and individuals who are unable to complete the consent process Outcome definitions: - Added 6 month data collection of select patient-reported outcomes Data sources: - Added 6 month data collection of participant surveys and participant medical information via site-completed eCRF	Protocol amended to reflect the addition of a 6 month data collection timepoint for both participant surveys and participant medical information via site-completed eCRF, add platform program language, to clearly define the index date as the date of first CAB+RPV injection, to reflect updates for planned country inclusion, update exclusion criteria, adverse events reporting and milestones

			Data management: • Language updated to be inclusive of newly added 6 month data collection timepoint • Added platform program language Ethical approval and subject consent: • Added platform program language Management and reporting of adverse events • Added some clarifications to this section Milestones: • Updated the milestones table	
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4. MILESTONES

Milestone	Planned/Estimated date
Draft Protocol	7 March 2025
Registered with HMA-EMA catalogue	30 April 2025
Final Protocol	01 May 2025
Protocol Amendment 1	19 October 2025
Protocol Amendment 2	30 April 2026
Start of data collection	November 2025
First Interim Tables (Data through October 2026)	December 2026
Second Interim Tables (Data through October 2027)	December 2027
Third Interim Tables (Data through October 2028)	December 2028
Final report (Data through when follow-up is completed on all participants)	12 months after end of data collection

5. RATIONALE AND BACKGROUND

CABFIDENCE platform program. The CABFIDENCE platform program is designed to generate a comprehensive, contemporary understanding of HIV treatment experiences and outcomes across antiretroviral therapy (ART) modalities within routine clinical practice. A platform approach enables coordinated evidence generation across related interventions, reduces redundancy, and improves speed and consistency of decision-making, while maintaining scientific rigor and regulatory acceptability.

A central element of the CABFIDENCE platform program is the use of a shared data lake, or data warehouse, to support studies on currently licensed ART regimens while future-proofing the program for regimens in clinical development. By leveraging aligned informed consent processes, common data models, and coordinated operational teams, the data lake enables efficient integration of data across studies while preserving study-specific attributes.

Another central element of the CABFIDENCE platform program is the use of a master protocol which harmonizes design elements across multiple ART specific observational study protocols. The master protocol defines all shared scientific, operational, and analytic elements of the platform, while individual study protocols independently describe treatment specific objectives, populations, and analyses within this common framework.

The platform design with the shared data lake and use of a master protocol efficiently accommodate sub-protocols for multiple drug products or indications with a shared comparator. The focus of this document is the first of the CABFIDENCE platform non-interventional, observational subprotocols to enroll people with HIV (PWH) who are receiving CAB+RPV LA.

Cabotegravir (CAB), an HIV-1 integrase strand transfer inhibitor (INSTI), in combination with rilpivirine (RPV), an HIV-1 NNRTI, is indicated as a complete regimen for the treatment of HIV-1 infection in adults who are virologically suppressed (HIV-1 RNA < 50 copies /mL) on a stable ART regimen without present or past evidence of viral resistance to, and no prior virological failure with CAB or RPV (in the US)/agents of the NNRTI and INI class (in Europe).

The novel long-acting ART delivered through intramuscular injection—CAB+RPV LA—has been shown to be non-inferior to daily oral therapy. Before starting the regimen, the healthcare provider should carefully select people with HIV who agree to the required injection schedule and counsel individuals about the importance of adherence to scheduled dosing visits to help maintain viral suppression and reduce the risk of viral rebound and potential development of resistance associated with missed doses. After discontinuation of CAB+RPV LA injection, it is essential to adopt an alternative, fully suppressive ARV regimen.

CAB in combination with RPV has a prolonged pharmacokinetic tail, which presents individuals with the opportunity to take their HIV treatment once monthly or once every 2 months instead of daily, potentially allowing for much improved adherence. While clinical studies have established the efficacy, safety, and tolerability of CAB+RPV LA, this study, as part of the CABFIDENCE platform program, will assess global utilization

patterns, adherence, persistence, discontinuation, effectiveness, and resistance among virologic failures with available resistance testing data, as well as patient-reported outcomes of this regimen in a real-world clinical setting.

6. RESEARCH QUESTION AND OBJECTIVE(S)

6.1. Primary Objective

The overall objective of this global study is to describe participant characteristics and assess clinical outcomes and patient-reported outcomes among people with HIV who receive CAB+RPV LA regimen.

1. To describe baseline demographics, clinical characteristics and patterns of use among participants receiving CAB+RPV LA
 - Descriptive analysis of study population by baseline demographic and clinical characteristics
 - Monitor for use of OOLI
2. To assess the persistence, discontinuation, and adherence among participants receiving CAB+RPV LA
 - Persistence: Proportion of participants still on CAB+RPV LA at time of analysis, after 6, 12, and 24 months of use and at study end
 - Proportion of participants discontinuing the regimens of interest will be assessed
 - a. Reasons for discontinuation will be assessed
 - Non-adherence to the dosing schedule will be assessed by:
 - a. Estimating the number of participants who missed 1 or more consecutive injections without taking daily oral bridging therapy or any other oral ARV regimen while not on CAB+RPV LA and estimating the mean and median number of injections missed during a 12-month period
 - b. Estimating the number of participants who received the injections more than 7 days earlier or later than their scheduled injection visit and median time from target injection to actual injection date of early delayed injections
 - c. Estimating the number of participants who missed 1 or more consecutive injections without taking daily oral bridging therapy or those who received the injections more than 7 days earlier or later than their scheduled injection visit (combined group A or B) and describe individual characteristics for the participants who are non-adherent.
3. To assess virologic effectiveness (i.e., proportion of participants experiencing virologic suppression and CVF) among participants receiving CAB+RPV LA regimen
 - Estimate overall number and proportion of participants with CVF and time to CVF (months)

- Estimate the proportion of participants with CVF during the first 6 months after initiation of CAB+RPV LA
 - Estimate the proportion of participants with CVF 6, 12, and 24 months after initiation of CAB+RPV LA
 - Assessment of virologic suppression among participants on CAB+RPV LA regimen during first 6 months and at 6, 12, and 24 months after initiation of the regimen
 - Assessment of number and proportion of participants who remain suppressed (VL <50 c/mL) throughout all follow-up and at last follow-up
4. To assess factors associated with CVF among participants receiving CAB+RPV LA regimen

6.2. Secondary Objectives

1. Subgroup analyses by BMI (< 30, ≥ 30 -<40 and ≥ 40 kg/m²) at initiation, age at initiation (18-49, 50-64, ≥ 65 years), sex assigned at birth (female, male), race and ethnicity, sex/race, and country, if sample sizes allow, to assess characteristics, adherence, persistence, discontinuation, and virologic effectiveness
2. To estimate the frequency of documented injection site reactions and hypersensitivity reactions among participants receiving CAB+RPV LA injections
3. To describe resistance and ART regimen among participants after discontinuation/CVF of CAB+RPV LA, where VL data are available and resistance testing has been done as part of routine clinical practice
 - Describe resistance (where this has been done as part of routine clinical practice) at the time of discontinuation/CVF and history of resistance and proviral DNA testing before discontinuation/CVF
 - Describe the ART regimen and virologic outcomes after discontinuation/CVF of CAB+RPV LA
 - To describe the demographic and clinical characteristics of participants who continue to stay on CAB+RPV LA as compared to those who switched regimens after discontinuation or CVF
 - To describe virologic response (suppression, non-response, CVF) to ART regimen after discontinuation/CVF of CAB+RPV LA
4. To describe reasons for initiating CAB+RPV LA, health-related quality of life, treatment satisfaction, treatment preference, treatment-related stigma, benefits, and reasons for missed injections and discontinuation among prospective participants (new initiators) of CAB+RPV LA
5. For data collected beyond the two-year follow-up period, outcomes will be analyzed at the extended time points

7. RESEARCH METHODS

7.1. Study Design

This is a non-interventional prospective observational cohort study using data from individual medical records, assessments from participating clinical sites, and participant-reported data from electronic questionnaires. Prospective participants will be followed from the date of their first CAB+RPV injection (index date) for a minimum of two years, unless they withdraw from the study, are lost to follow-up, or pass away. Participants that discontinue CAB+RPV LA and switch to a different ART regimen or to monthly CAB+RPV LA dosing schedule or experience CVF, will continue to be followed to describe virologic response and treatment switching. Retrospective participants will be included in the study regardless of whether they are still on CAB+RPV at the time of the study start and will have all applicable data collected. Participants who are deceased or lost to follow up will not be included.

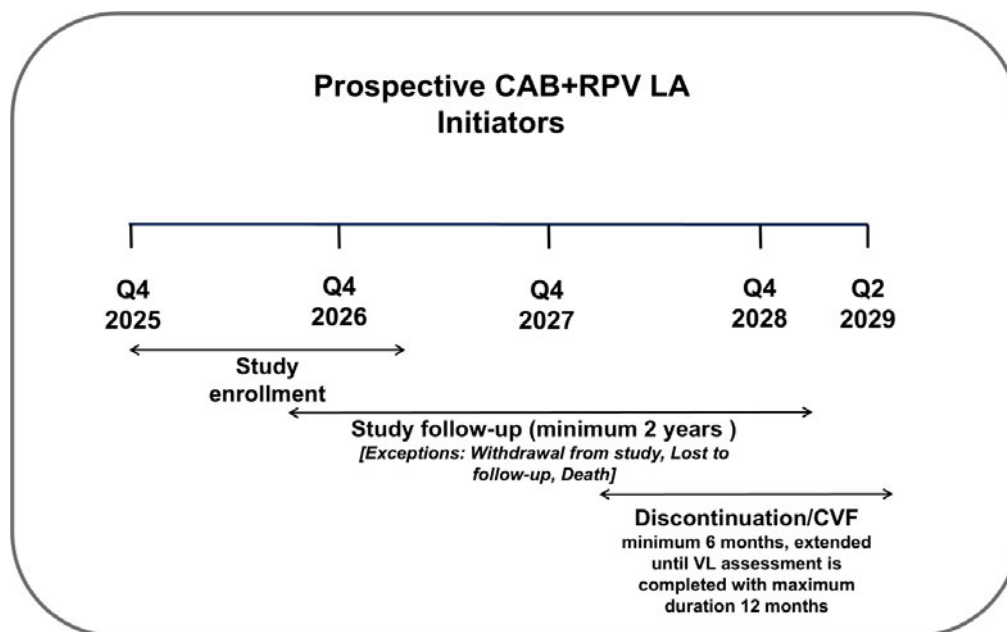
For this non-interventional study, treatment and laboratory testing decisions will be made by the treating physician according to standard practice, taking into account the treatment history, individual characteristics, the approved prescribing information or summary of product characteristics for CAB+RPV LA formulation, contemporary regimen and local guideline or recommendations. Testing for resistance and subtype at discontinuation/virologic failure will be part of the standard of care at selected sites, and the study will include only the clinical sites that conduct resistance and subtype testing as standard of care. The study protocol will be implemented by RTI Health Solutions (RTI-HS).

7.2. Study Population and Setting

The study population includes treatment-experienced, virologically suppressed ($VL < 50$ copies/mL) individuals who are 18 years or older and received at least one CAB+RPV LA injection with every-2 month dosing schedule, from participating clinical sites across several countries. Anticipated countries include the UK, Italy, Portugal, France, Spain, Japan, Australia, and the US. Enrollment will take place as part of a participant's standard of care visit at a participating site.

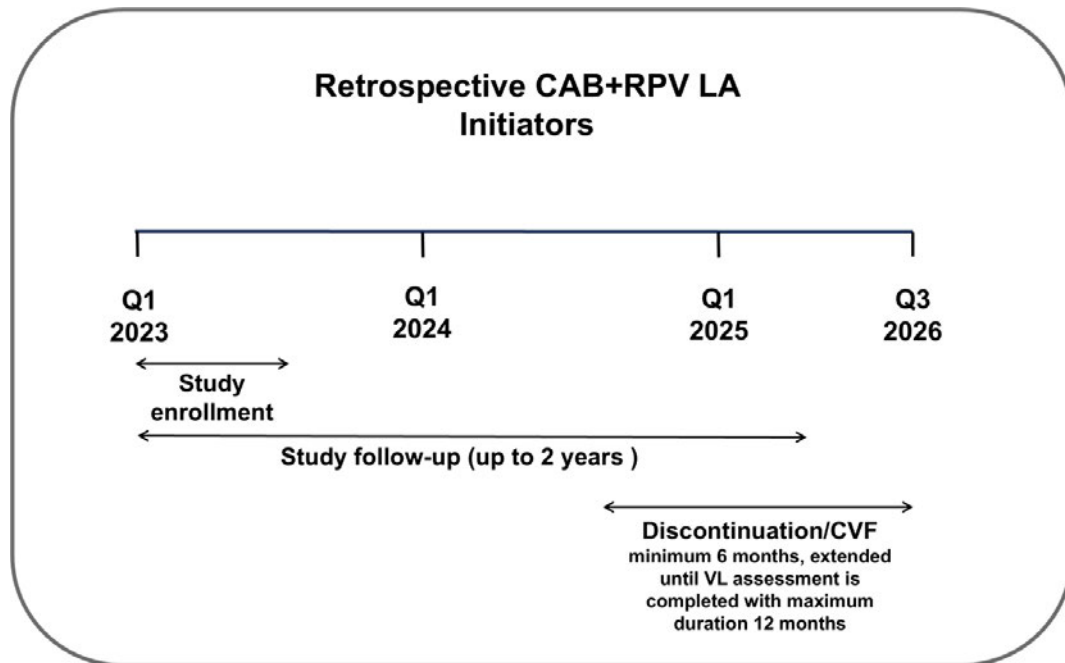
Prospective Participants:

Prospective participants will include individuals enrolled from November 2025 until study size is reached. Participants will be enrolled prior to their first CAB+RPV LA injection, either during the optional oral lead-in (OOLI) phase, if applicable, or directly before their first CAB+RPV LA injection for those on the direct-to-injection (DTI) pathway. The baseline survey will be administered before the first CAB+RPV LA injection. The participants will be followed for a minimum of two years, unless they withdraw from the study, are lost to follow-up, or pass away. Participants who discontinue and switch to a subsequent regimen or who experience CVF will be followed-up for a minimum of 6 months and may be extended until viral load assessment is completed, with a maximum duration of 12 months. Study end is the date when the last participant completes the required follow-up assessment (including any required post-discontinuation/CVF follow-up).

**Retrospective Participants:**

Retrospective participants will include individuals who initiated CAB+RPV LA from January 2023 and will be followed for up to two years. Participants who have discontinued CAB+RPV LA for any reason will be included and participants who are deceased or lost to follow up will not be included. Participants who discontinue and switch to a subsequent regimen or who experience CVF will be followed-up for a minimum of 6 months and may be extended until viral load assessment is completed, with a maximum duration of 12 months. Study end is the date when the last participant completes the required follow-up assessment (including any required post-discontinuation/CVF follow-up).

The sampling methodology will be designed with the goal of maximising the probability that each individual treated with CAB+RPV LA will have an equal opportunity to be selected. Participants will be enrolled as they come in for a scheduled visit, however if the volume of participants at a site is large, a customized, site-specific sampling approach may be devised to achieve an efficient method of obtaining a representative sample of participants. In some sites, it is likely that all treated participants will be selected.



The study will aim to select a diverse group of medical practices to represent diversity in clinical practice settings and to support recruitment of a diverse sample of participants (e.g., prioritizing non-cisgender males, individuals aged 50 years or older, and individuals with BMI ≥ 30 kg/m²).

Individuals who are viremic (VL ≥ 50 copies/mL) at initiation will not be included in this study. Individuals with VL results available around the time of initiation at VL < 50 copies/mL will be included in the study regardless of VL levels following enrollment.

7.2.1 Inclusion criteria:

- Diagnosis of HIV-1
- Aged ≥ 18 years at time of CAB+RPV LA injection
- Treatment experienced
- VL < 50 copies/mL at initiation
- Initiated every 2 month dosing CAB+RPV LA
 - For prospective participants, study enrollment should occur prior to the first CAB+RPV LA injection, either during the OOLI phase, if applicable, or directly before their first CAB+RPV LA injection for those on the DTI pathway
 - For retrospective participants, individuals who initiated CAB+RPV LA from January 2023 will be enrolled

7.2.2 Exclusion Criteria

- Received CAB+RPV LA regimen as part of a randomized controlled clinical trial
- Treatment naïve

- VL $\geq$ 50 copies/mL at initiation
- Individuals with documented prior virologic failure to agents of NNRTI or INSTI class (if data is available)
- Individuals with documented present or past resistance to agents of the NNRTI or INSTI class (if data is available)
- Individuals who are unable to complete the consent process

7.3. Variables

7.3.1. Exposure Definitions

All users of CAB+RPV LA regimen treated per the approved prescribing information or summary of product characteristics for CAB+RPV LA will be included in the study.

7.3.2. Outcome Definitions

Endpoint	Definition
Primary	
Persistence	▪ Proportion of participants still on CAB+RPV LA at time of analysis, after 6, 12, and 24 months of use and at study end
Discontinuation	▪ Proportion discontinuing after last injection: ○ Date of discontinuation will be the date of the last injection prior to: ▪ no injections and no documentation of oral bridging for more than 127 days after any injection. ▪ Physician confirmed treatment switch from CAB+RPV LA to another ARV regimen ▪ Physician confirmed treatment stop of CAB+RPV LA ▪ Median time to discontinuation ▪ Proportion of participants who discontinued and reinitiated the regimen within 6 months from discontinuation

Endpoint	Definition																					
Adherence	- Adherence to the dosing schedule will be assessed as the proportion of participants who received their injections within the +/- 7-day dosing window relative to their target treatment date as detailed below (see table: Days After Last Injection) - Non-adherence to the dosing schedule will be assessed as follows: - Describe the proportion of participants who received CAB+RPV LA more than 7 days earlier or later than their target treatment date - Mean/median (range) number and time from target injection date to actual injection date of early or delayed injections and number of participants with 1, 2, 3 and ≥4 early or delayed injections - Describe the proportion of participants who missed one consecutive injection with and without taking daily oral therapy or any other oral ARV regimen while not on CAB+RPV LA - Mean and median number of injections missed, time from target injection date to actual injection date, and number of participants with 1, 2, 3 and ≥4 missed injections																					
	<table><tr><td></td><th colspan="2">Days after last injection</th></tr><tr><td></td><th>Q2M 2nd initiation</th><th>Q2M continuation injection</th></tr><tr><td>On-time</td><td>23-37</td><td>53-67</td></tr><tr><td>Late</td><td>38-67</td><td>68-127</td></tr><tr><td>Delayed (Short delay)</td><td>38-67</td><td>68-97</td></tr><tr><td>Long delay (requiring re-initiation)</td><td>NA</td><td>98-127</td></tr><tr><td>Requiring re-initiation/ Discontinuation</td><td>>67</td><td>>127</td></tr></table>		Days after last injection			Q2M 2 nd initiation	Q2M continuation injection	On-time	23-37	53-67	Late	38-67	68-127	Delayed (Short delay)	38-67	68-97	Long delay (requiring re-initiation)	NA	98-127	Requiring re-initiation/ Discontinuation	>67	>127
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Long delay (requiring re-initiation)	NA	98-127																				
Requiring re-initiation/ Discontinuation	>67	>127																				

Endpoint	Definition
CVF	Proportion of participants with: ▪ 2 consecutive HIV RNA VL $\geq$ 200 copies/mL prior to/by assessment timepoint - OR ▪ 1 HIV RNA VL level $\geq$ 200 copies/mL prior to/by assessment timepoint followed by core agent/regimen discontinuation within 4 months of HIV RNA VL level $\geq$ 200 copies/mL
Virologic suppression	▪ Proportion of participants with VL measurement $<$ 50 copies/mL during follow-up
Non-response	▪ Not achieving re-suppression after CVF, consisting of VL $\geq$ 50 copies/mL following CVF with no VL of $<$ 50 copies/mL following CVF
Secondary	
Injection site reactions	▪ Proportion of participants with documented diagnostic code/diagnosis of injection site reaction or nodules attributed to administration of CAB+RPV LA ▪ Proportion of participants with Division of AIDS grading 1-4 to classify severity
Hypersensitivity reactions	▪ Proportion of participants with documented diagnosis of Hypersensitivity, anaphylactic reaction, allergic reaction or drug allergy to CAB+RPV LA ▪ 2 or more events are reported from 2 or more of the following groups of signs/symptoms: ○ Rash, fever, gastrointestinal symptoms (nausea, vomiting, diarrhea, abdominal pain), constitutional symptoms (lethargy, fatigue, malaise, myalgia, arthralgia, general ill feeling) respiratory symptoms (dyspnea, sore throat, cough, chest x-ray changes, predominantly infiltrates, which can be localized) ▪ Cases where there is a negative rechallenge with CAB+RPV LA ▪ Cases where symptoms resolved (or did not worsen/result in withdrawal from the study)
Emergent resistance	▪ Proportion of participants with resistance and proviral DNA testing at discontinuation/CVF.

Endpoint	Definition
	▪ Proportion of participants with history of resistance and proviral DNA testing prior to discontinuation/CVF. ▪ The proportion of participants tested with any resistance mutation will be reported at discontinuation/CVF and history of prior to discontinuation/CVF, as will mutations associated with reduced susceptibility at the class-level and drug-level ○ All mutations will be reported; specific resistance mutations of interest to CAB+RPV LA can be highlighted and identified using the International Antiviral Society mutations list and Stanford HIV drug resistance database.
Patient-reported outcomes	
Reasons for initiating CAB+RPV LA	▪ Descriptive summary of bespoke questionnaire (2 items) completed at baseline
Baseline adherence	▪ Descriptive summary of bespoke questionnaire (7 items) completed at baseline
HIV health-related quality of life	▪ Descriptive summary of WHOQOL-HIV BREF (31 items) completed at baseline, month 6, month 12 and discontinuation, if applicable
HIV treatment-related worries and stigma	▪ Descriptive summary of HIV Treatment-Related Worries (HRW; 4 items) completed at baseline, month 6, month 12 and discontinuation, if applicable
Treatment-related benefits	▪ Descriptive summary of bespoke questionnaire (7 items) completed at month 6, month 12, if applicable
Treatment satisfaction	▪ Descriptive summary of HIVTSQs (12 items) completed at baseline, month 6, and month 12, if applicable
Change in treatment satisfaction	▪ Descriptive summary of HIVTSQc (12 items) completed at month 6, month 12, if applicable
Treatment preference	▪ Descriptive summary of bespoke questionnaire (3 items) completed at month 6 and month 12, if applicable
Reasons for missed injection(s)	▪ Descriptive summary of bespoke questionnaire (4 items) completed at month 6 and month 12, if applicable
Reasons for discontinuation	▪ Descriptive summary of bespoke questionnaire (3 items) completed at discontinuation
Social determinants of health (SDoH)	▪ Descriptive summary of bespoke questionnaire (7 items) completed at baseline, month 6, and month 12, if applicable

ARV = antiretroviral; CAB = cabotegravir; CVF = confirmed virologic failure; HIV = human immunodeficiency virus; HIVTSQs = HIV Treatment Satisfaction Questionnaire status version; HIVTSQc = HIV Treatment Satisfaction Questionnaire change version; LA = long-acting; RPV = rilpivirine; VL = viral load; WHOQOL-HIV BREF = World Health Organization Quality of Life-Human Immunodeficiency Virus BREF.

7.3.3. Confounders and Effect Modifiers

Standard variables to be evaluated and included at the index date:

Demographic variables at index

- Age (years, also categorized as 18-25, 26-49, 50-64, ≥ 65 years)
- Sex assigned at birth (female, male)
- Gender identity
- Race
- Ethnicity
- Marital status (single, married/domestic partner, widowed, separated/divorced, unknown)
- Geographic region
- Route of HIV infection (men who have sex with men, people who inject drugs, heterosexual, vertical, Other)
- Payer type (Medicaid, Medicare, commercial insurance, Veteran's Affairs, AIDS Drug Assistance Programs/Ryan White, cash with self-pay, other) (US only)

Virologic variables at regimen initiation

- HIV VL at initiation of CAB+RPV LA regimen
 - Continuous (copies/mL)
 - Categorical:
 - < 50 copies/mL (or $<$ limit of detection in clinics where limit of detection > 50 copies/mL)
 - ≥ 50 to < 200 copies/mL
 - ≥ 200 to $< 1,000$ copies/mL
 - $\geq 1,000$ to $< 10,000$ copies/mL
 - $\geq 10,000$ to $< 100,000$ copies/mL
 - $\geq 100,000$ copies/mL
 - Missing
- Type of VL assay
 - Abbott RealTime
 - Aptima Quant Dx
 - Roche 6800

- Roche Cobas

Immunologic variables at regimen initiation

- CD4 cell count at initiation of CAB+RPV LA regimen
 - Continuous (cells/ μ L)
 - Categorical:
 - > 500 cells/ μ L
 - > 350 to ≤ 500 cells/ μ L
 - > 200 to ≤ 350 cells/ μ L
 - > 50 to ≤ 200 cells/ μ L)
 - ≤ 50 cells/ μ L
 - Missing
- CD4/CD8 ratio
- CD4 nadir prior to initiation of CAB+RPV LA

Clinical variables at regimen initiation

- Number of years since HIV diagnosis
- Number of years since first ART initiation
- HIV subtype where available
- History of previous ARV exposure before CAB+RPV LA
 - Number of core agents experienced
 - Number of ARV classes experienced
 - Duration of last ARV regimen
 - Prior core agent class
- History of previous virologic failure
- History of previous HIV drug resistance and any historical HIV-1 subtype information, where available
- BMI, height, weight
- History of AIDS-defining events
- Veterans Aging Cohort Study (VACS) Mortality Index score
 - Characteristics for mortality index calculation include age, CD4 count, HIV-1 RNA level, Hemoglobin, FIB-4 index, eGFR, HCV status, albumin, BMI, white blood cell count
- ASCVD risk score
 - Characteristics for CVD risk calculation include age, sex, race, total, LDL and HDL cholesterol, systolic and diastolic blood pressure, on

antihypertensive therapy, on a statin, on aspirin therapy, history of diabetes, and smoking status.

- Framingham risk score
 - Characteristics for CVD risk calculation include age, sex, total and high-density lipoprotein cholesterol, systolic blood pressure, antihypertensive therapy, and smoking status.
- Co-infections
 - Hepatitis B co-infection including serology status
 - Hepatitis C co-infection
 - Syphilis infection (ever)
- Comorbidities
 - Autoimmune disease
 - Cardiovascular disease
 - Invasive cancer
 - Endocrine disorder
 - Mental health disorder
 - Depression
 - Anxiety
 - Sleeplessness
 - Suicidality
 - Liver disease
 - Bone disorder
 - Peripheral neuropathy
 - Renal disease
 - Hypertension
 - Substance abuse
 - Non-AIDS–defining malignancies (excluding hepatocellular carcinoma)
 - Any of the above
- Concomitant medications
 - Anticonvulsants
 - Carbamazepine
 - Oxcarbazepine
 - Phenobarbital
 - Phenytoin

 - Antimycobacterials

- Rifampin
- Rifapentine
- Rifabutin
- Glucocorticoids
 - Dexamethasone
- Macrolide or ketolide antibiotics
 - Azithromycin
 - Clarithromycin
 - Erythromycin
- Narcotic analgesic
 - Methadone

7.4. Data Sources

- **Participant surveys:** Prospective participants of CAB+RPV LA will be asked to complete an online survey at enrollment, at month 6 (if applicable), at month 12 (if applicable), and if they discontinue CAB+RPV LA. The baseline survey will be administered prior to the first CAB+RPV LA injection. The online survey will include the WHOQOL-HIV BREF, the HIVTSQs, and the HIVTSQc (O’Connell, 2012; Woodcock and Bradley, 2001; Woodcock and Bradley, 2006) and several bespoke questionnaires developed for this study. Appropriate agreements with the copyright owners will be in place for use in this study. It is estimated that the participant surveys will take approximately 20 minutes to complete.
- **Participant medical information via site-completed eCRF:** The investigator or authorized medical staff will record clinical and treatment data from participants’ medical records into an eCRF at enrollment, month 6, and a minimum of two annual follow-ups (unless participant discontinued CAB+RPV LA) and at study end.

Participant surveys and medical record information will be linked by a common participant identifier (ID) code that will be retained by the site.

7.5. Study Size

The study will aim to enroll up to approximately 2,000 participants. Among them, it is estimated that 250 participants are already receiving CAB+RPV LA at the time of study start, and 1,750 participants will be newly initiating CAB+RPV LA.

There is no formal statistical hypothesis to be tested in these analyses. The study will be descriptive in nature and will focus on absolute effectiveness estimation. A precision calculation was performed using power analysis and sample size (PASS) software version 14 using the exact Clopper-Pearson method based on the cumulative probabilities of the binomial distribution for 1 proportion.

In order to demonstrate the estimated precision for the total sample size and also among the groups of participants already receiving and newly initiating CAB+RPV LA, Table 1 presents the precision calculations for the width of 95% confidence intervals for a range of proportions and a combination of sample sizes. It is anticipated that data from 2,000

participants are thought adequate to provide reasonable precision around a point estimate (here defined as a proportion) to meet the study objectives. With 2,000 participants, the 2-sided 95% confidence interval precision width for the estimation of virologic failure rate will range from 1.28% and 2.13% for a failure rate ranging from 2% to 6%.

Table 1. Exact Confidence Intervals for Various Combinations of Sample Size and Proportions

Confidence level	Sample size (N)	Proportion (P)	95% Lower limit	95% Upper limit	Actual width
0.95	250	0.0200	0.0065	0.0461	0.0395
	250	0.0300	0.0126	0.0595	0.0469
	250	0.0400	0.0193	0.0723	0.0530
	250	0.0500	0.0265	0.0848	0.0583
	250	0.0600	0.0340	0.0970	0.0631
	500	0.0200	0.0096	0.0365	0.0268
	500	0.0300	0.0169	0.0490	0.0321
	500	0.0400	0.0246	0.0611	0.0365
	500	0.0500	0.0326	0.0729	0.0403
	500	0.0600	0.0408	0.0846	0.0437
	750	0.0200	0.0112	0.0328	0.0215
	750	0.0300	0.0190	0.0449	0.0259
	750	0.0400	0.0271	0.0566	0.0295
	750	0.0500	0.0355	0.0681	0.0326
	750	0.0600	0.0441	0.0795	0.0354
	1,000	0.0200	0.0123	0.0307	0.0185
	1,000	0.0300	0.0203	0.0426	0.0222
	1,000	0.0400	0.0287	0.0541	0.0253
	1,000	0.0500	0.0373	0.0654	0.0281
	1,000	0.0600	0.0461	0.0766	0.0305
	1,250	0.0200	0.0130	0.0294	0.0164
	1,250	0.0300	0.0213	0.0410	0.0198
	1,250	0.0400	0.0298	0.0524	0.0226
	1,250	0.0500	0.0386	0.0636	0.0250
	1,250	0.0600	0.0475	0.0746	0.0272
	1,500	0.0200	0.0135	0.0284	0.0149
	1,500	0.0300	0.0220	0.0399	0.0180
	1,500	0.0400	0.0307	0.0512	0.0205
	1,500	0.0500	0.0395	0.0623	0.0228
	1,500	0.0600	0.0485	0.0732	0.0247
	1,750	0.0200	0.0140	0.0277	0.0137
	1,750	0.0300	0.0225	0.0391	0.0166
	1,750	0.0400	0.0313	0.0503	0.0190
	1,750	0.0500	0.0403	0.0613	0.0210
	1,750	0.0600	0.0493	0.0722	0.0228
	2,000	0.0200	0.0143	0.0271	0.0128

Confidence level	Sample size (N)	Proportion (P)	95%	95% Upper limit	Actual width
			Lower limit		
	2,000	0.0300	0.0230	0.0385	0.0155
	2,000	0.0400	0.0318	0.0495	0.0177
	2,000	0.0500	0.0409	0.0605	0.0196
	2,000	0.0600	0.0500	0.0713	0.0213

7.6. Data Management

A central element of the CABFIDENCE platform program is the use of a shared data lake, or data warehouse. This centralized data environment serves as a single, secure repository for study data and is designed to ensure consistent data structure, standardized variable definitions, and harmonized data capture processes across cohorts developed in sub-protocols. The shared data lake is housed within RTI-HS secure infrastructure and resides behind the RTI firewall. Access controls, technical safeguards, and governance processes are implemented in alignment with all applicable data privacy, security, and regulatory requirements, as well as the data protection commitments described in the informed consent documents. All data use within the repository is governed by the scope of participant consent and overseen through established RTI-HS data governance procedures to ensure confidentiality, integrity, and appropriate use of study data.

The Qualtrics electronic data capture (EDC) system will be used to capture data from eCRFs and participant surveys. The system will be programmed and tested by RTI-HS data management staff. Participants who consent and are formally enrolled in the study will be asked to complete the baseline survey on a tablet provided to sites by RTI-HS while at the clinical site and the months 6 and 12 follow-up surveys either on their own device via an electronic link provided by RTI-HS or on a tablet provided to sites by RTI-HS. If participants skip a question, they will be prompted to complete it. If they still wish to skip it, they will be able to continue to the next question. Investigators or authorized staff will enter clinical data from participants' medical records directly into the EDC system at the protocol specified timepoints. There is no plan to reconcile any discrepancies in data between individual responses to the survey and their medical record data.

Data will be linked by a common participant identification (ID) code (i.e., participant data are de-identified). Limited personal IDs (e.g., age but not names or email addresses) will be stored with the data.

All data management activities will be fully documented in a detailed data management plan. The plan will describe the EDC system used in the study, user acceptance testing of the system, data sources, data collection/entry methods, querying of missing or unclear data, data cleaning plans, data storage procedures during the study, data set creation, database lock, and data archival. Data from this study may be utilized to compare to outcomes among individuals on other ARV regimens collected as part of other protocols or studies.

7.7. Data Analysis

7.7.1. Primary Analysis

7.7.1.1. Main analytical approach

Baseline characteristics and outcomes will be described using counts and relative frequencies for categorical variables and means, standard deviations, medians, minimums, maximums, and quartiles 1 and 3 for continuous variables. For event outcomes, incidence rates will be estimated using unadjusted Poisson regression, accounting for person-time since index (i.e., first CAB+RPV LA injection) where appropriate. Additional analytic approaches may also be considered for select time-to-event outcomes (e.g., time to discontinuation) such as estimators that take into account right-censoring (e.g., Kaplan-Meier estimator) and multivariable models such as logistic regression as applicable.

Analyses will be stratified by BMI (< 30 , ≥ 30 -40 and ≥ 40 kg/m²), age (18-49, 50-64, ≥ 65 years), sex (male, female), race/ethnicity, sex/race, and country at first CAB+RPV LA injections, as sample size allows, to assess characteristics, adherence, persistence, discontinuation, and virologic effectiveness. Other demographic and clinical variables may also be considered for stratification, if relevant.

Additionally, factors such as, age, sex assigned at birth, race, geographic region, BMI, injection drug use (IDU), CD4 count, history of AIDS-defining events, core class in prior regimen and comorbidities associated with CVF will be explored using a logistic regression model.

The number and proportion of participants with history of resistance to INSTI or NNRTI and individual ARTs and the results of any historical resistance testing will be assessed. Similarly number and proportion of participants with discontinuation/CVF will be assessed for resistance testing and specific INSTI and/or NNRTI RAMs will be described. Subsequent regimen after discontinuation/CVF and virologic outcomes will be described. Participant surveys will be summarized descriptively for item and summary score (where applicable). The WHOQOL-HIV BREF, HIVTSQs, and HIVTSQc will be scored according to instrument scoring guidelines and derived scores will be summarized descriptively. All analyses will be performed on the observed data only and no missing values will be imputed.

7.7.1.2. Data handling conventions

Data handling conventions applicable to this study design as described in the Guidelines for Good Pharmacoepidemiology Practices (GPP) issued by the International Society for Pharmacoepidemiology (ISPE) will be followed to ensure ethical and scientific recording and reporting of non-interventional trials that involve data from human subjects. The following procedures will be undertaken:

- Automatic checks and prompts are built into the EDC program (e.g., checks for out-of-range numeric values). Each individual eCRF will be further evaluated for logical consistency between study variables, potential outliers, and missing information.

- Inconsistencies noted in the eCRF data will be queried with the site using data-clarification procedures. Sites will be queried for clarification via the EDC system if any such issues are identified. Database lock will be issued when all queries have been resolved.
- Participant medical chart data will be abstracted into the eCRF by qualified site staff, identified only by the unique subject identification number. Linkage between the identification number and the participant records will be recorded at a site-level only and will not be shared externally with RTI-HS or ViiV Healthcare. Data will be abstracted by site staff from participant medical records and entered into an eCRF hosted on the secure EDC system. The link between the subject identification code and the individual participant must be destroyed by the site at the end of the study.

7.7.1.3. Sensitivity analyses

No sensitivity analyses are planned.

7.7.2. Secondary Analysis/Exploratory Analysis

Individual surveys will be summarized descriptively for item and summary score (where applicable). The WHOQOL-HIV BREF, HIVTSQs, and the HIVTSQc will be scored according to instrument scoring guidelines. All analyses will be performed on the observed data only, and no missing values will be imputed.

7.8. Quality Control and Quality Assurance

This project will be conducted in accordance with all applicable regulatory requirements. The Office of Quality Assurance, an independent unit that reports to the vice president of RTI-HS, will oversee quality assurance for this study.

Standard operating procedures will be used to guide the conduct of the study. These procedures include rules for secure and confidential data storage, methods to maintain and archive project documents, quality control procedures for programming, standards for writing analysis plans, and requirements for senior scientific review.

All programming written by one study analyst will be independently reviewed by a different analyst, with oversight by a senior statistician.

All key study documents, such as the analysis plan, data collection forms, and study reports, will undergo quality control review, senior scientific review, and editorial review.

7.8.1. Site Training

Investigators and study coordinators will be trained via an initial web-based training session on the protocol, including adverse event (AE) reporting, study flow, EDC system, documentation, site responsibilities and expectations, and any applicable study processes. Any new information relevant to the performance of this study will be forwarded to the medical staff during the study.

No on-site monitoring visits will be performed for this study unless there is a specific need to meet with the site team. Remote data monitoring will be conducted during the life of the study to ensure linkage and integrity between the eCRFs and the participant surveys and to identify missing or unclear data in the eCRFs and issue queries.

RTI-HS will closely monitor the participant recruitment and data collection on a regular basis. Specifically, RTI-HS will provide oversight of activities, including screening participants, recruiting and obtaining consent from participants, and completion of electronic surveys and eCRFs. RTI-HS will maintain regular communication with all sites and will assess progress and site performance and address any issues as they arise.

7.9. Limitations of the Research Methods

Limitations of this study are common to non-randomized non-interventional study. This is a study of routine clinical care and reflects treatment practice across multiple countries and diverse clinical sites. This will be a single treatment group analysis with no formal statistical comparisons. While the study aims to include approximately 2,000 users of the CAB+RPV LA regimen, market uptake of the regimen and participants willingness to participate will dictate timelines for enrollment and data collection. The results from this study should be interpreted cautiously, with careful consideration given to the limitations of the observational study design.

7.9.1. Study Closure/Uninterpretability of Results

Not applicable; these are descriptive analyses of observational, real-world data.

7.10. Other Aspects

None

8. PROTECTION OF HUMAN SUBJECTS

8.1. Ethical Approval and Subject Consent

The study will be conducted in accordance with legal and regulatory requirements; with scientific purpose, value, and rigor; and will follow generally accepted research practices described in GPP issued by Public Policy Committee of the ISPE.

Ethics review requirements and processes for prospective studies vary among countries and are governed by local standards. This study will be conducted in accordance with such standards in each country.

RTI-HS, with assistance from local clinical research associates, will work in collaboration with the site Principal Investigators to submit the necessary ethics review applications in each country at both the site and the national or regional levels, as well as notify/register the local Health Authority, as and where applicable.

Informed consent will be obtained from participants (or their legally acceptable representative) by site study staff before data collection. The person obtaining consent

will be responsible for ensuring that each participant fully understands the nature, purpose, risks, and benefits associated with participation. Each participants will be provided with a copy of his or her signed informed consent form. The investigator, or a person designated by the investigator, will retain the original of each participant's signed consent form.

Data from this study may be utilized to compare to outcomes among individuals on other ARV regimens collected as part of other sub-protocols developed under the master protocol. Participants enrolled under each sub-protocol provide informed consent for analysis of their data across the CABFIDENCE study program. This consent framework and the master protocol allow data collected under different sub-protocols to be combined and compared within the centralized data lake, supporting integrated analyses across treatment modalities. In addition, participant consent permits the use of their data in conjunction with existing and future comparable HIV sub-protocols within the CABFIDENCE platform program, where scientifically appropriate and consistent with ethical, regulatory, and privacy requirements. This forward-looking consent approach is intended to maximize the long-term scientific value of the data while maintaining transparency and respect for participant autonomy.

Participants (or the participant's legally acceptable representative) may choose to withdraw consent at any time. At time of withdrawal, all data that have already been processed and analyzed will be kept to guarantee the integrity of the study. Once data has been processed and analyzed (i.e., at the point of data analysis), the information may continue to be used.

8.2. Subject Confidentiality

ViiV and all investigators will ensure adherence to applicable data privacy protection regulation. The research team at RTI-HS and any approved vendors will have access to pseudonymized individual data. Sites will maintain a subject identification log with the names/initials/identifiers of participants who consented to participate in the study and study subject ID. Sites will assign subject ID at time of participant enrollment. Subject identification logs will be maintained at the sites and will never be shared with the research team at RTI-HS, third parties, or ViiV. Additionally, the collected data (eCRF and individual questionnaires) will not contain any participant -identifying information to further protect confidentiality (for instance, age or year of birth will be collected in lieu of a full birth date). To comply with applicable data protection regulations, European Union (EU) and UK personal data will be collected and stored in the EU or UK and only staff dedicated to this project and located in EU will have access to these. Pseudonymized data sets will be transferred to the US. Finally, only aggregated data in the form of analysis tables will be made available to ViiV. Thus, any reports generated will not contain any participant identifiers.

9. LEGAL BASIS FOR PROCESSING INDIVIDUAL HUMAN DATA

The authors confirm that study data is Individual Human Data not owned by ViiV, but that the proposed use of the Individual Human Data aligns with the ‘purpose of use’ outlined in the source contract and/or the terms and conditions of use of the data source and it will comply with any specified prohibitions of use.

10. MANAGEMENT AND REPORTING OF ADVERSE EVENTS/ADVERSE REACTIONS

This study has safety objectives. Safety collection and reporting will correspond to the following study types based on the specific data sources:

Participant surveys

The participant surveys represent study type 1: Primary data collection studies with or without sites/investigators with safety objectives where there is consent and/or data collection direct from individuals. This part of the study is able to identify solicited events only.

Collection of Adverse Events/Reactions (Solicited Events)

Solicited Events are defined as adverse events related to the GSK/ViiV product under evaluation and identified for collection in the study database as per study objectives.

The purpose of the study is to collect real-world data on global utilization patterns, adherence, persistence, discontinuation, virologic effectiveness, and patient-reported outcomes for initiators of CAB+RPV LA.

Solicited events related to CAB+RPV LA must be collected in the study database and reported to GSK for entry into the GSK Safety database. All AEs/SAEs, pregnancies and incidents, systematically collected in the Participant Surveys, which will be based on closed ended questions¹ only, should be reported to the GSK Safety department (timing and contact information provided in the safety management plan). For primary data collection studies, where there is direct collection of data from individuals, these will be classified as solicited individual case safety reports (ICSRs). ICSR will also be summarised in interim, if applicable, and final study reports.

¹ A closed ended question is where a respondent can only choose a pre-set response.

Reporting of Adverse Events/Reactions (Spontaneous Events)

Adverse events not related to CAB+RPV LA cannot be reported in the participant surveys due to the interview/survey structure being based on closed ended questions only.

Site completed e-CRF

The site completed e-CRF represent study type 5: Secondary data collection studies including unstructured data with human review with safety objectives. These studies can identify solicited events in aggregate at study end but cannot identify spontaneous events.

Collection of adverse events/reactions (Solicited Events)

Solicited Events are defined as pre-specified adverse events related to the GSK/ViiV product under evaluation and identified for collection as per study objectives.

The purpose of the study is to collect real-world data on global utilization patterns, adherence, persistence, discontinuation, virologic effectiveness, and patient-reported outcomes for initiators of CAB+RPV LA.

For CAB+RPV LA regimen, pre-defined safety events of interest persistence, adherence, discontinuation and virologic effectiveness, as documented in participants' medical records will be systematically recorded in aggregate. These will be summarised in final study reports. This part of the study is based on secondary use of existing health data and as such Individual Case Safety Reporting (ICSRs) to regulatory agencies is not required. These events are not entered into the GSK Safety database.

Reporting of adverse events/reactions (Spontaneous Events)

This study is based on data previously collected for other purposes e.g., routine healthcare encounters. As such, there is no requirement for the collection and reporting of Individual Case Safety Reports (ICSRs). Although the study is based on human review of unstructured data, the nature of the secondary data protocol driven data collection and analysis does not allow for reporting of serious and non-serious AEs, pregnancy exposures, or incidents related to any GSK/ViiV product during the conduct of this research. In addition, the minimum criteria of identifiable patient, reporter, exposure and event, needed to report individual case safety reports may not be present in the information reviewed within the context of the study. The data also may lack an identifiable patient and reporter and may be insufficient to establish attribution between a potential safety event and an individual patient using a GSK/ViiV product.

11. PLANS FOR DISSEMINATING AND COMMUNICATING STUDY RESULTS

The target audience for these data includes healthcare providers, health plan population-based decision-makers, and regulatory and health authorities.

Abstracts will be prepared for both the interim and final analysis, with a manuscript developed from the final data set. All publications will comply with the *International Committee of Medical Journal Editors* (ICMJE) guidelines. A final full study report will be submitted to the Sponsor. Study results will be submitted to scientific conferences and for peer-reviewed journal publication.

12. REFERENCES

O'Connell, KA, Skevington, SM. An international quality of life instrument to assess wellbeing in adults who are HIV-positive: a short form of the WHOQOL-HIV (31 items). *AIDS Behav.* 2012;16:452-60. doi:10.1007/s10461-010-9863-0.

Woodcock, A., Bradley, C. Validation of the HIV Treatment Satisfaction Questionnaire (HIVTSQ). *Qual Life Res.* 2001;10:517-31

Woodcock A, Bradley C. Validation of the revised 10-item HIV Treatment Satisfaction Questionnaire status version and new change version. *Value Health.* 2006;9(5):320-33.