

HIV-1 RNA BLIPS AND LOW-LEVEL REPLICATION DURING PHASE II/IIB CABOTEGRAVIR + RILPIVIRINE LONG-ACTING STUDIES ARE SIMILAR TO ORAL **3-DRUG THERAPY AND NOT ASSOCIATED WITH WEEK 48 VIROLOGIC OUTCOME**

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Introduction

- Findings from phase III/IIIb studies demonstrated that long-acting (LA) cabotegravir (CAB) + rilpivirine (RPV) dosed every 4 weeks (Q4W) was noninferior to current antiretroviral (ART) regimen (CAR; FLAIR¹ and ATLAS²), and dosing every 8 weeks (Q8W) was noninferior to Q4W (ATLAS-2M³) through Week 48
- Presence of transient, elevated HIV-1 RNA levels as a predictor for durability of ART is of clinical interest,⁴ but the clinical relevance of such HIV-1 RNA blips is unknown⁵
- This analysis evaluates the robustness of antiviral potency of CAB LA + RPV LA using these measures (Table 1):
- Number of participants with HIV-1 RNA blips overall and at individual study visits
- Qualitative detection of HIV-1 RNA
- HIV-1 RNA low-copy analysis
- Presence or absence of HIV-1 RNA blips in participants with HIV-1 RNA <50 c/mL at Week 48 (Snapshot analysis)

Methods

Study Design

- FLAIR (ClinicalTrials.gov identifier: NCT02938520), ATLAS (ClinicalTrials.gov identifier: NCT02951052), and ATLAS-2M (ClinicalTrials.gov identifier: NCT03299049) are phase III/IIIb, randomized (1:1), open-label studies designed to assess antiviral activity and safety of intramuscular (IM) CAB + RPV LA (Q4W or Q8W)
- Eligible participants were adults (aged ≥18 years) with HIV-1 infection who were treatment naive (FLAIR) or had no history of virologic failure to ART (ATLAS and ATLAS-2M) and HIV-1 RNA <50 c/mL before randomization
- All participants who had not previously taken CAB + RPV LA received a 4-week oral lead-in of CAB 30 mg + RPV 25 mg once daily to assess individual tolerability prior to receiving LA injections
- Participants were given IM injections of CAB LA 400 mg + RPV LA 600 mg Q4W (FLAIR/ATLAS/ATLAS-2M) Q4W group) or CAB LA 600 mg + RPV LA 900 mg Q8W (ATLAS-2M Q8W group) as maintenance doses or remained on CAR

Endpoints and Assessments

• Plasma samples were collected at study visits to quantitatively and qualitatively analyze HIV-1 RNA (Table 1)

Table 1. Outcome Definitions

Outcome	Definition	Assay	Assessment
Blip	HIV-1 RNA 50 to <200 c/mL with adjacent values <50 c/mL ⁴	Abbott RealTime HIV-1 Assay	BL to Week 48
TD/TND	Qualitative outcome for HIV-1 RNA <40 c/mL	Abbott RealTime HIV-1 Assay	BL to Week 48
Low copy	HIV-1 RNA <2 c/mL limit of detection ⁶	bioMONTR Labs HIV-1 SuperLow Assay	BL and Week 48

BL, Baseline; TD, target detected; TND, target not detected.

Results

Participant Disposition and Baseline Characteristics

 Demographic characteristics of the FLAIR, ATLAS, and ATLAS-2M study participants were generally similar between treatment groups and have been previously presented¹⁻³

Blip Outcomes

- Overall, proportions of participants with HIV-1 RNA blips were similar in the Q4W CAB + RPV LA and CAR groups in FLAIR and ATLAS and in the Q4W and Q8W CAB + RPV LA groups in ATLAS-2M (Table 2)
- Blip rate was higher in treatment-naive participants from FLAIR vs treatment-experienced participants from either treatment group in ATLAS and ATLAS-2M

Table 2. Participants With Increased Viral Load: Intention-to-Treat–Exposed Populations

	FLAIR		ATLAS		ATLAS-2M	
Parameter, n (%)	Q4W IM (n=283)	ABC/DTG/3TC (n=283)	Q4W IM (n=308)	CAR (n=308)	Q8W IM (n=522)	Q4W IM (n=523)
Blip	36 (13)	37 (13)	18 (6)	22 (7)	18 (3)	29 (6)
≥2 consecutive viral loads ≥50 to <200 c/mL	6 (2)	4 (1)	0 (0)	6 (2)	2 (<1)	4 (<1)



90 80 70 60 50 40 20 10 blips

n=

Qualitative Outcomes for Target Not Detected

HIV-1 Low-Copy Assay Outcomes

• Across treatment groups and studies, the majority of study participants virologically suppressed (HIV-1 RNA <50 c/mL) had plasma HIV-1 RNA <2 c/mL at Baseline and Week 48 (Figure 4)

 Proportions of participants with HIV-1 RNA blips were similar for each study week and consistently occurred in <5% of participants with available HIV-1 RNA data (Figure 1)

Figure 1. Proportion of Participants With Blips by Visit: Intention-to-Treat-Exposed Populations of (A) FLAIR, (B) ATLAS, and (C) ATLAS-2M



^aWeek 4 data not reported for FLAIR and ATLAS because only participants who received CAB + RPV LA had viral load measured at Week 4 per study protocol

• Few participants with HIV-1 RNA blips in these trials had HIV-1 RNA ≥50 c/mL at Week 48, and participants with or without blips and HIV-1 RNA <50 c/mL were comparable using the Snapshot algorithm (Figure 2) • Of 17 participants with confirmed virologic failure (CVF) in CAB + RPV LA groups,¹⁻³ only 1 had a blip before reaching CVF criteria (2 consecutive HIV-1 RNA measurements ≥200 c/mL)

Figure 2. Snapshot Analysis Outcomes by Presence of Blips in Intention-to-Treat–Exposed Populations of (A) FLAIR, (B) ATLAS, and (C) ATLAS-2M



aNo virologic data reported due to participant discontinuation for adverse events, deaths, or other events or participant on study but missing data at Week 48. bThe participant with blips in ATLAS-2M Q8W had CVF.

• Among participants with plasma HIV-1 RNA <40 c/mL at any timepoint, the majority (>75%) had qualitative outcomes for TND (Figure 3)

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Conclusions

- CAR aroups through Week 48
- analysis) at Week 48
- phase III studies
- These data further reinforce the potency and robustness of CAB + RPV LA for treatment of HIV-1 infection

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 Proportions of participants in phase III/IIIb studies with HIV-1 RNA blips, TND viral load results. and HIV-1 RNA <2 c/mL were similar in the Q4W and Q8W CAB + RPV LA and the oral 3-drug

 Majority (89%-100%) of participants with HIV-1 RNA blips was virologically suppressed (Snapshot) Occurrence of blips through Week 48 did not correlate with Week 48 Snapshot outcomes in any group from