Bictegravir + Lenacapavir: Baseline and Week 48 Resistance Analyses in ARTISTRY-1 Phase 2

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Conclusions

- The high frequency of baseline RAMs among VS participants in ARTISTRY-1 Phase 2 was consistent with the study requirement that participants be on a complex regimen
- The presence of baseline RAMs did not affect the efficacy of BIC + LEN
- 93% treatment success overall, regardless of baseline RAMs, including baseline INSTI RAMs
- On-treatment emergence of capsid polymorphism (N74T) was observed in one participant who resuppressed on a LEN-containing regimen
- No phenotypic loss of susceptibility to LEN
- These data underscore the potential of the use of once-daily BIC + LEN to optimize therapy in people receiving a complex ART regimen

Plain Language Summary

- At least two or more drugs (called antiretroviral drugs) are needed to treat people who have human immunodeficiency virus (HIV). Some antiretroviral drugs may not work for some people with HIV, due to side effects or because the virus has become resistant to the drugs (in other words, those specific drugs no longer kill the virus). As a result, these people sometimes need to take multiple pills each day to treat their HIV; this can be hard to do.
- Lenacapavir and bictegravir are two drugs that are approved to treat HIV infection and are each taken together with other drugs. Combining these two drugs into one pill taken once a day could help lower the number of pills that people need to take to treat their HIV infection, and make their treatment easier to take.
- People in this study took an average of three pills per day to prevent the growth of their HIV and had no HIV in their blood at the start of the study. After 48 weeks of treatment, people who switched their treatment plan to the combination of lenacapavir plus bictegravir experienced a similar result as when they were still taking many pills to treat their HIV. This showed that combining lenacapavir and bictegravir was an effective treatment.
- Even though many participants with HIV had mutations (changes) in their HIV showing resistance (some drugs may no longer kill the virus) when this study started, giving lenacapavir and bictegravir together stopped the virus from making copies of itself.

Introduction

- Once-daily single tablet regimens (STRs) are standard of care for HIV treatment
- However, many people with HIV (PWH) take complex antiretroviral therapy (ART) regimens for reasons including drug resistance, intolerance, toxicity, drug-drug interactions, and contraindications to existing STRs¹⁻⁴
- A bictegravir + lenacapavir (BIC + LEN) STR could optimize the treatment of virologically suppressed (VS) PWH who are unable to take currently available STRs
- BIC is a global, guideline-recommended integrase strand transfer inhibitor (INSTI) with a high barrier to resistance^{1,5-7}
- LEN is a first-in-class capsid inhibitor with an orthogonal resistance profile to existing/major drug classes⁸
- There is an absence of capsid inhibitor resistance in the HIV community⁹
- ARTISTRY-1 (NCT05502341) is a Phase 2/3 study evaluating the safety and efficacy of a complete regimen combining BIC and LEN - At Baseline, 43%, 19%, 11%, and 27% of participants were taking 2, 3, 4, or \geq 5 pills/day, respectively
- In Phase 2 of ARTISTRY-1, the efficacy of BIC + LEN was compared with remaining on a complex multi-tablet ART regimen in VS participants (n = 128)
- Arm 1: BIC 75 mg + LEN 25 mg, n = 51; Arm 2: BIC 75 mg + LEN 50 mg, n = 52; Arm 3: Complex ART regimen, n = 25
- High rates of virologic suppression were observed across treatment groups at Week 48 (90.4% to 100%; no statistical differences between groups)¹⁰

References: 1. Department of Health and Human Services. https://clinicalinfo.hiv.gov/sites/default/files/guidelines/documents/ adult-adolescent-arv/guidelines-adult-adolescent-arv.pdf (accessed April 23, 2024). 2. Chang H-M, et al. BMC Infect Dis. 2022;22:2. 3. Rolle C-P, et al. J Virus Erad. 2020;6:100021. 4. Colloty J, et al. Br J Hosp Med (Lond). 2023;84:1-9. 5. Acosta RK, et al. Antimicrob Agents Chemother. 2019;63:e02533-18. 6. European AIDS Clinical Society. https://www.eacsociety.org/media/guidelines-12.0.pdf (accessed July 10, 2024). 7. Gandhi RT, et al. JAMA. 2023;329:63-84. 8. Margot N, et al. Antimicrob Agents Chemother. 2021;65:e02057-20. 9. Marcelin A-G, et al. J Antimicrob Chemother. 2020;75:1588-90. 10. Mounzer K, et al. Clin Infect Dis. November 2024 [ePub]. doi:10.1093/cid/ciae522. 11. Demirdjian S, et al. Poster 681 presented at: Conference on Retroviruses and Opportunistic Infections (CROI); March 3-6, 2024; Denver, CO.



Objectives

Characterize the baseline resistance profile of participants

Analyze the potential emergence of resistance to study drugs after 48 weeks of treatment

Methods

• ARTISTRY-1 is a Phase 2/3, randomized, open-label, multicenter study

Study Design of Phase 2 of ARTISTRY-1 (NCT05502341)

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Results

Baseline Pre-Existing Resistance Mutations Per Treatment Group



	Participants With Virologic Suppression, % (n/N) ^a					
	BIC-LEN 25 mg	BIC-LEN 50 mg	Complex ART	All		
All participants	92.2 (47/51)	90.4 (47/52)	100 (25/25)	93.0 (119/128)		
With NRTI-RAMs	92.5 (37/40)	91.9 (34/37)	100 (18/18)	93.7 (89/95)		
Without NRTI-RAMs	88.9 (8/9)	86.7 (13/15)	100 (5/5)	89.7 (26/29)		
<i>P</i> value ^b	0.57	0.62	1.0	0.44		
With NNRTI-RAMs	92.6 (25/27)	93.5 (29/31)	100 (16/16)	94.6 (70/74)		
Without NNRTI-RAMs	90.9 (20/22)	85.7 (18/21)	100 (7/7)	90.0 (45/50)		
<i>P</i> value ^b	1.0	0.38	1.0	0.48		
With PI-RAMs	100 (22/22)	88.2 (15/17)	100 (12/12)	96.1 (49/51)		
Without PI-RAMs	85.2 (23/27)	91.4 (32/35)	100 (11/11)	90.4 (66/73)		
<i>P</i> value ^b	0.12	1.0	1.0	0.31		
With INSTI-RAMs ^c	88.9 (8/9)	0	100 (1/1)	90.0 (9/10)		
Without INSTI-RAMs ^c	88.5 (23/26)	89.7 (35/39)	100 (19/19)	91.7 (77/84)		
<i>P</i> value ^b	1.0	NA	1.0	1.0		
US EDA-defined Snapshot algorithm: treatment success if HIV-1 RNA < 50 c/mL b Eisher exact test $^{\circ}$ Proviral DNA analysis only (n with data = 94). For all three other mutation categorie						

ID	INSTI RAMs	Treatment Group	Treatment Outcome at Week 48
1	T66T/A E92E/Q S147S/G Q148Q/R N155N/H	BIC-LEN 25 mg	Virologic suppression
2	R263R/K	BIC-LEN 25 mg	Virologic suppression
3	T66T/A	BIC-LEN 25 mg	Virologic suppression
4	G140G/S Q148Q/H	BIC-LEN 25 mg	Virologic suppression
5	Q148Q/H	BIC-LEN 25 mg	Virologic suppression
6	Q148Q/R	Complex ART	Virologic suppression
7	N155N/H	BIC-LEN 25 mg	Participant's decision to discontinue (virologic suppression at last visit)
8	N155N/H	BIC-LEN 25 mg	Virologic suppression
9	N155N/H	BIC-LEN 25 mg	Virologic suppression
10	G140G/R	BIC-LEN 25 mg	Virologic suppression

Disclosures: All authors are employees and stock holders of Gilead Sciences, Inc.

None of the participants with baseline INSTI RAMs experienced treatment failure with BIC + LEN

Participant With HIV-1 RNA ≥ 50 c/mL at Week 48 – 1 of 3



BIC, bictegravir; c, copies; LEN, lenacapavir.

- FDA Snapshot HIV-1 RNA ≥ 50 c/mL at Week 48: HIV-1 RNA = 50 c/mL

Participant With HIV-1 RNA ≥ 50 c/mL at Week 48 – 2 of 3



BIC, bictegravir; c, copies; LEN, lenacapavir.

- FDA Snapshot HIV-1 RNA ≥ 50 c/mL at Week 48: HIV-1 RNA = 98 c/mL

Participant With HIV-1 RNA \geq 50 c/mL at Week 48 – 3 of 3



BIC, bictegravir; c, copies; CA, capsid; DTG, dolutegravir; LEN, lenacapavir; MVC, maraviroc; SC, subcutaneous.

- FDA Snapshot HIV-1 RNA ≥ 50 c/mL at Week 48: HIV-1 RNA = 305 c/mL
- Resistance analysis showed emergence of N74T in capsid with PhenoSense assay failure;

Phenotypic Characterization of N74T Capsid Mutation

- Capsid mutants containing N74T were generated by site-directed mutagenesis

Mutant	n ^a	Mean LEN EC₅₀ Fold Change from WT				
wutant		MT-2	RevLun	Monogram		
N74wt	3	1.0	1.0	1.0		
N74T	3	1.4	1.2	1.1		
N74D	3	24.0	12.5	14.1		

EC₅₀, effective concentration to inhibit 50% of viral replication; LEN, lenacapavir; WT, wild type.

- N74T does not confer resistance to LEN In contrast to N74D control mutants
- N74T is a novel capsid polymorphism
- No change in susceptibility to BIC

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BIC 75 mg + LEN 25 mg

• Participant did not meet threshold for resistance analysis (viral load $[VL] \ge 200 \text{ c/mL}$) and resuppressed at next visit

BIC 75 mg + LEN 25 mg



• Participant did not meet threshold for resistance analysis (VL \ge 200 c/mL) and resuppressed at next visit

BIC 75 mg + LEN 50 mg

participant resuppressed subsequently with new regimen of dolutegravir + maraviroc + subcutaneous LEN

— Laboratory strain (HIV-1XXLAI) and three clinical samples with diverse capsid sequence

• Mutants were tested in single-cycle (Monogram Biosciences) and multicycle (MT-2 and RevLun) assays

^aEach mutant was tested in three to four to different genetic constructs; repeated at least three times per construct.