

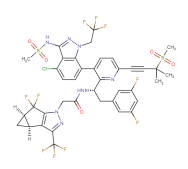
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# Lenacapavir (LEN)

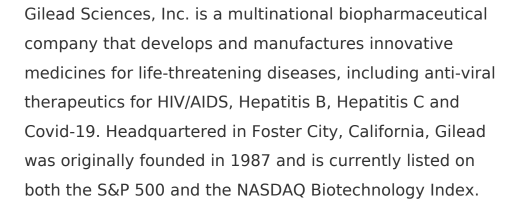
# **Developer(s)**

Gilead Sciences Inc.

Originator

https://www.gilead.com/

**United States** 





# **Drug structure**

Lenacapavir Compound Structure

Sourced From DrugBank

# **Drug information**

# **Associated long-acting platforms**

Aqueous Solution, Oral solid form

# **Administration route**

Oral, Subcutaneous, Intramuscular, To be determined

# Therapeutic area(s)

HIV

# Use case(s)

Pre-Exposure Prophylaxis (PrEP)

Treatment

Prevention

# Use of drug

#### Ease of administration

Administered by a community health worker

Administered by a nurse

Administered by a specialty health worker

Self-administered

To be determined

# **User acceptance**

Not provided

# **Dosage**

## Available dose and strength

LEN oral tablets 300 mg; each injection contains 927 mg of lenacapavir in solution. Dose for investigational Once-Yearly formulation is 5000 mg.

# Frequency of administration

Oral tablets 300 mg taken daily or weekly; Six-monthly injectable; Once-yearly investigational injectable.

#### Maximum dose

5000 mg

## Recommended dosing regimen

For PrEP: Initiation Option 1: Day 1: 927 mg by subcutaneous injection and 600 mg orally (2 x 300-mg tablets). Day 2: 600 mg orally (2 x 300-mg tablets). Initiation Option 2: Day 1: 600 mg orally (2 x 300-mg tablets). Day 2: 600 mg orally (2 x 300-mg tablets). Day 8: 300 mg orally (1 x 300-mg tablet). Day 15: 927 mg by subcutaneous injection. Maintenance: 927 mg by subcutaneous injection every 26 weeks +/- 2 weeks from date of last injection. For the treatment indication, lenacapavir is administered as part of a full treatment regimen with the relevant associated medicines.

#### Additional comments

Not provided

# Dosage link(s)

Not provided

# **Drug information**

# Drug's link(s)

https://go.drugbank.com/drugs/DB15673

#### Generic name

Lenacapavir

#### **Brand name**

Sunlenca

### Compound type

Small molecule

## **Summary**

Lenacapavir (LEN), also known as GS-6207, is a first in-class HIV-1 capsid inhibitor used in combination with other antiretrovirals for the treatment of multi-drug resistant HIV-1 infection, and has potential application as HIV pre-exposure prophylaxis. LEN is utilised combinatorially for HIV-1 treatment, as it displays excellent synergy and no known cross-resistance with any other currently approved class of antiretroviral, in addition to possessing antiviral activity at picomolar levels.

# **Approval status**

for Treatment: Lenacapavir (SUNLENCA) 463.5mg/3ml subcutaneous injection with 300mg oral lead-in tablets are approved for use by 11 regualtory authorities (all for high income countries) for HIV-1 treatment under certain conditions. For PrEP: US FDA has approved lenacapavir (Yeztugo) for HIV-1 prevention. It is also under review in Brazil, EU (+EEA), Australia, Canada and South Africa. Gilead has disclosed that it is preparing additional filings in countries that rely on FDA approval for regulatory submission, including Argentina, Mexico and Peru.

# Regulatory authorities

US FDA granted Breakthrough Therapy Designation for SUNLENCA in combination with other antiretroviral drugs for heavily treatment-experienced patients (HTE) adults with multi-drug resistant (MDR) HIV-1 infection failing their current antiretroviral regimen due to resistance, intolerance, or safety considerations. A European Marketing Authorization was issued for the use of SUNLENCA and it has also been classified as 'Fast-Track Reimbursement' by the Ministry of Health, Labour and Welfare, Japan, and 'Part 1- Schedule 1 & Schedule 3 Poison' by the Department of Health, Hong Kong. WHO guidelines released in July 2025 recommend offering six-monthly injectable lenacapavir as an additional PrEP option.

# **Delivery device(s)**

No delivery device

# Scale-up and manufacturing prospects

### **Scale-up prospects**

Compound is commercially manufactured.

### Tentative equipment list for manufacturing

Equipment for injectable: Stainless steel pharmaceutical reactors, glass-lined reactors, rotary evaporator (rotovap), flash chromatography columns, stainless steel autoclave, cooling bath, silica gel chromatography columns, vacuum distillation apparatus, simulated moving bed chromatography system, Chiralpak columns.

# Manufacturing

Storage of injectable lenacapavir in borosilicate vials is contraindicated due to issues with chemical compatibility. Instead, it is recommended that vials are made from aluminosilicate glass.

# Specific analytical instrument required for characterization of formulation

Proton nuclear magnetic resonance (1H NMR), High-performance liquid chromatography (HPLC), Ultra-Performance Liquid Chromatography (UPLC).

# **Clinical trials**

### **CAPELLA**

### Identifier

NCT04150068

#### Link

https://clinicaltrials.gov/ct2/show/NCT04150068

### Phase

Phase II/III

#### **Status**

Active, not recruiting

# **Sponsor**

Gilead Sciences

#### More details

Not provided

# **Purpose**

Evaluate the antiviral activity of Lenacapavir (formerly GS-6207) administered as an add-on to a failing regimen (functional monotherapy) in people living with HIV with multi-drug resistance.

#### **Interventions**

### Intervention 1

Drug: Oral Lenacapavir

#### **Intervention 2**

Drug: Oral Lenacapavir Placebo

#### **Intervention 3**

Drug: Subcutaneous Lenacapavir

#### **Intervention 4**

Drug: Failing ARV Regimen

#### **Intervention 5**

Drug: Optimized Background Regimen (OBR)

### **Countries**

Canada

France

Germany

Dominican Republic

Italy

Japan

South Africa

Spain

Thailand

United States of America

### Sites / Institutions

Not provided

### **Trials dates**

### **Anticipated Start Date**

Not provided

#### **Actual Start Date**

2019-11-21 00:00:00

### **Anticipated Date of Last Follow-up**

2024-06-26 00:00:00

### **Estimated Primary Completion Date**

Not provided

### **Estimated Completion Date**

2027-01-01 00:00:00

# **Actual Primary Completion Date**

2020-10-05 00:00:00

### **Actual Completion Date**

Not provided

# **Studied populations**

### **Age Cohort**

- Children
- Adolescents
- Adults
- Older Adults

#### **Genders**

All

### Accepts pregnant individuals

Unspecified

# **Accepts lactating individuals**

Unspecified

Accepts healthy individuals

No

Comments about the studied populations

Adult aged  $\geq$  18 years (at all sites) or adolescent aged  $\geq$  12 and weighing  $\geq$  35 kg (at sites in North America and Dominican Republic). Currently receiving a stable failing ARV regimen for > 8 weeks. Have HIV-1 RNA  $\geq$  400 copies/mL at screening. Have multidrug resistance (resistance to  $\geq$ 2 agents from  $\geq$ 3 of the 4 main classes of ARV). Have no more than 2 fully active ARV remaining from the 4 main classes that can be effectively combined to form a viable regimen. Able and willing to receive an Optimized Background Regimen (OBR) together with Lenacapavir.

#### **Health status**

Positive to: HIV

Negative to : HCV

Study type

Interventional (clinical trial)

**Enrollment** 

72

Allocation

Randomized

Intervention model

Sequential assignment

Intervention model description

Not provided

Masking

Quadruple-blind masking

# **Masking description**

Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor)

# Frequency of administration

Once every 6 months

# Studied LA-formulation(s)

Injectable

# Studied route(s) of administration

Oral

Subcutaneous

# Use case

Treatment

# **Key resources**

Туре	Title	Content	Link
Link	Capsid Inhibition w	vith	https://www.nejm.org/doi/10
	Lenacapavir in		
	Multidrug-Resistar	nt	
	HIV-1 Infection		

# **CALIBRATE**

Identifier
NCT04143594
Link
https://clinicaltrials.gov/ct2/show/NCT04143594
Phase
Phase II
Status
Completed
Sponsor
Gilead Sciences
More details
Not provided
Purpose
Evaluate the efficacy of Lenacapavir containing regimens in people living with HIV
Interventions
Intervention 1  Drug: Oral Lenacapavir
Intervention 2
Drug: F/TAF
Intervention 3

Drug: Subcutaneous Lenacapavir

**Intervention 4** 

Drug: TAF

**Intervention 5** 

Drug: BIC

### **Countries**

Dominican Republic

Puerto Rico

United States of America

### Sites / Institutions

Not provided

#### **Trials dates**

### **Anticipated Start Date**

Not provided

**Actual Start Date** 

2019-11-22 00:00:00

**Anticipated Date of Last Follow-up** 

2023-10-03 00:00:00

**Estimated Primary Completion Date** 

Not provided

**Estimated Completion Date** 

Not provided

**Actual Primary Completion Date** 

2021-09-30 00:00:00

**Actual Completion Date** 

### Studied populations

#### **Age Cohort**

- Adults
- Older Adults

#### **Genders**

All

#### Accepts pregnant individuals

Unspecified

### **Accepts lactating individuals**

Unspecified

#### Accepts healthy individuals

No

# Comments about the studied populations

Antiretroviral (ARV) naïve with no use of any ARV within one month of screening. Use of pre-exposure prophylaxis (PrEP) (any duration), post-exposure prophylaxis (PEP) (any duration), or HIV-1 treatment (< 10 days therapy total) > 1 month prior to screening is permitted. HIV-1 RNA  $\ge 200$  copies/mL at screening. CD4+ cell count  $\ge 200$  cells/microliter at screening.

#### **Health status**

Positive to: HIV

Negative to : HCV, HBV

# Study type

Interventional (clinical trial)

183
Allocation
Randomized
Intervention model
Parallel Assignment
Intervention model description
Not provided
Masking
Open label
Masking description
None (Open Label)
Frequency of administration
Once every 6 months
Studied LA-formulation(s)
Injectable
Studied route(s) of administration
Oral Subcutaneous
Use case
Treatment

**Enrollment** 

# **Key resources**

Туре	Title	Content	Link
Link	CROI 2022: Lenacapavir: 54 week results in treatment- naive participants of CALIBRATE study		https://i- base.info/htb/42313
Link	Lenacapavir administered every 26 weeks or daily in combination with oral daily antiretroviral therapy for initial treatment of HIV: a randomised, open- label, active- controlled, phase 2 trial		https://doi.org/10.1016/S235
Link	Interim Resistance Analysis of Long- Acting Lenacapavir in Treatment-Naïve People with HIV at 28 Weeks		https://doi.org/10.1093%2Fd

# GS-US-536-5816

Identifier
NCT04811040
Link
https://clinicaltrials.gov/ct2/show/NCT04811040
Phase
Phase I
Status
Completed
Sponsor
Gilead Sciences
More details
Not provided
Purpose
Evaluate the safety and tolerability of a combination of the broadly neutralizing antibodies (bNAbs) teropavimab (formerly GS-5423) and GS-2872 in combination with the HIV capsid inhibitor lenacapavir
Interventions
Intervention 1

Intervention 2

Drug: Oral Lenacapavir

Drug: Subcutaneous Lenacapavir

**Intervention 3** 

Biological: Teropavimab

**Intervention 4** 

Biological: Zinlirvimab

Countries

United States of America

Sites / Institutions

Not provided

**Trials dates** 

**Anticipated Start Date** 

Not provided

**Actual Start Date** 

2021-04-08 00:00:00

**Anticipated Date of Last Follow-up** 

2023-10-26 00:00:00

**Estimated Primary Completion Date** 

Not provided

**Estimated Completion Date** 

Not provided

**Actual Primary Completion Date** 

2023-04-18 00:00:00

**Actual Completion Date** 

2023-10-17 00:00:00

# Studied populations

### **Age Cohort**

- Adults
- Older Adults

#### **Genders**

All

### **Accepts pregnant individuals**

Unspecified

### **Accepts lactating individuals**

Unspecified

### **Accepts healthy individuals**

No

# Comments about the studied populations

On first-line antiretroviral therapy (ART) for  $\geq 2$  years prior to screening. A change in ART regimen  $\geq 28$  days prior to screening for reasons other than virologic failure (VF) (eg, tolerability, simplification, drug-drug interaction profile) is allowed.

#### **Health status**

Positive to: HIV

Negative to: HCV, HBV

Other health status: No history of opportunistic infection or illness indicative of Stage 3

HIV disease; No comorbid condition(s) requiring ongoing immunosuppression.

### Study type

Interventional (clinical trial)

#### **Enrollment**

32
Allocation
Randomized
Intervention model
Parallel Assignment
Intervention model description
Not provided
Masking
Double-blind masking
Masking description
Double (Participant, Investigator). Clinical pharmacologist and sponsor are not masked to treatment assignment.
Frequency of administration
Other
Studied LA-formulation(s)
Injectable
Studied route(s) of administration
Oral

Subcutaneous

# Use case

Treatment

# **Key resources**

Not provided

# GS-US-200-4072

**Intervention 1** 

**Intervention 2** 

Drug: Lenacapavir Subcutaneous Injection

Identifier
NCT03739866
Link
https://clinicaltrials.gov/ct2/show/NCT03739866
Phase
Phase I
Status
Completed
Sponsor
Gilead Sciences
More details
Not provided
Purpose
Separately evaluate the short-term antiviral activity of both lenacapavir and tenofovir alafenamide with respect to plasma HIV-1 RNA reduction in antiretroviral or capsid inhibitor naïve patients
Interventions

Drug: Placebo

**Intervention 3** 

Drug: B/F/TAF

**Intervention 4** 

Drug: TAF

### Countries

United States of America

### Sites / Institutions

Not provided

#### **Trials dates**

### **Anticipated Start Date**

Not provided

#### **Actual Start Date**

2018-11-26 00:00:00

# **Anticipated Date of Last Follow-up**

2021-03-16 00:00:00

# **Estimated Primary Completion Date**

Not provided

### **Estimated Completion Date**

Not provided

# **Actual Primary Completion Date**

2019-11-14 00:00:00

### **Actual Completion Date**

2020-06-15 00:00:00

# **Studied populations**

### **Age Cohort**

- Adults
- Older Adults

#### **Genders**

All

# Accepts pregnant individuals

No

# **Accepts lactating individuals**

No

# Accepts healthy individuals

No

# Comments about the studied populations

Treatment naïve or experienced but CAI and integrase strand transfer inhibitor (INSTI) naïve, and have not received any antiretroviral therapy (ART) within 12 weeks of screening.

#### **Health status**

Positive to: HIV

# Study type

Interventional (clinical trial)

### **Enrollment**

53

#### Allocation

21.				
Туре	Title	Content	Link	
Key resources	8			
Treatment				
Use case				
Subcutaneous				
Studied route	(s) of administra	ition		
Injectable				
Studied LA-fo	rmulation(s)			
Other : "Single	dose "			
Frequency of	administration			
Double (Particip	oant, Investigator)	)		
Masking desc	-			
Double-blind m	-			
Masking				
Not provided	iodei descriptio	''		
Intervention m	nodel descriptio	n		
Parallel Assignr	ment			
Intervention n	nodel			
Randomized				

Link

Clinical targeting of HIV capsid protein with a long-acting small molecule https://doi.org/10.1038/s415

# GS-US-536-5939

#### Identifier

NCT05729568

#### Link

https://clinicaltrials.gov/study/NCT05729568

### **Phase**

Phase II

### **Status**

Active, not recruiting

## **Sponsor**

Gilead Sciences

#### More details

Not provided

# **Purpose**

Evaluate the Safety and Efficacy of bNAbs GS-5423 and GS-2872 in Combination With Lenacapavir as Long-Acting Treatment Dosed Every 6 Months in Virologically Suppressed Adults With HIV-1 Infection.

#### Interventions

#### Intervention 1

Drug: Teropavimab (Formerly GS-5423)

### **Intervention 2**

Drug: Zinlirvimab (Formerly GS-2872)

**Intervention 3** 

Drug: Lenacapavir Tablet

**Intervention 4** 

Drug: Lenacapavir Injection

**Intervention 5** 

Drug: Antiretroviral Therapy

#### Countries

Australia

Canada

Puerto Rico

United States of America

### Sites / Institutions

Not provided

#### **Trials dates**

#### **Anticipated Start Date**

Not provided

**Actual Start Date** 

2023-05-15 00:00:00

**Anticipated Date of Last Follow-up** 

2024-07-12 00:00:00

**Estimated Primary Completion Date** 

2025-03-01 00:00:00

**Estimated Completion Date** 

2029-12-01 00:00:00

### **Actual Primary Completion Date**

2024-07-02 00:00:00

### **Actual Completion Date**

Not provided

# Studied populations

### **Age Cohort**

- Adults
- Older Adults

#### **Genders**

All

#### Accepts pregnant individuals

Unspecified

### **Accepts lactating individuals**

Unspecified

#### Accepts healthy individuals

No

# Comments about the studied populations

Participants are required to be receiving a stable ART regimen with no clinically significant documented resistance (except isolated NRTI mutations). Plasma HIV-1 RNA < 50 copies/mL at screening visit 2 and documented plasma HIV-1 RNA < 50 copies/mL for  $\geq 12$  months preceding screening visit 2.

#### **Health status**

Positive to: HIV

Negative to : HCV, HBV

# Study type

Enrollment
83
Allocation
Randomized
Intervention model
Parallel Assignment
Intervention model description
Not provided
Masking
Open label
Masking description
None (Open Label)
Frequency of administration
Once every 6 months
Studied LA-formulation(s)
Injectable
Studied route(s) of administration
Subcutaneous
Use case
Treatment

Interventional (clinical trial)

# **Key resources**

Not provided

# **IMEA 070**

France

Identifier
NCT06289361
Link
https://clinicaltrials.gov/study/NCT06289361
Phase
Marketed
Status
Not yet recruiting
Sponsor
Institut de Médecine et d'Epidémiologie Appliquée - Fondation Internationale Léon M'Ba
More details
Immunovirological follow-up and safety of HIV-infected patients receiving lenacapavir under compassionate access in France between 01/01/2021 and 12/31/2023
Purpose
Cohort IMEA 070 -Lenacapavir Compassional
Interventions
Not provided
Countries

### Sites / Institutions

Not provided

#### **Trials dates**

### **Anticipated Start Date**

2024-04-01 00:00:00

### **Actual Start Date**

Not provided

### **Anticipated Date of Last Follow-up**

2024-03-20 00:00:00

# **Estimated Primary Completion Date**

2024-04-15 00:00:00

### **Estimated Completion Date**

2024-07-30 00:00:00

# **Actual Primary Completion Date**

Not provided

# **Actual Completion Date**

Not provided

# Studied populations

### **Age Cohort**

- Adults
- Older Adults

### **Genders**

All

# **Accepts pregnant individuals**

Unspecified
Accepts lactating individuals Unspecified
Accepts healthy individuals No
Comments about the studied populations
Not provided
Health status
Positive to : HIV
Study type
Observational studies (incl. patient registries)
Enrollment
58
Allocation
Not provided
Intervention model
Not provided
Intervention model description
Not provided
Masking
Not provided
Masking description

Not provided

# Frequency of administration

Once every 6 months

# Studied LA-formulation(s)

Injectable

# Studied route(s) of administration

Subcutaneous

#### Use case

Treatment

# **Key resources**

Not provided

# Pharmacokinetics and safety of once-yearly lenacapavir: a phase 1, open-label study

open-label study
Identifier
Not provided
Link
https://doi.org/10.1016/S0140-6736(25)00405-2
Phase
Phase I
Status
Not provided
Sponsor
Gilead Sciences Inc.
More details
Not provided
Purpose
Not provided
Interventions
Intervention 1 Once-yearly intramuscular lenacapavir formulation 1 Dosage: 5000 mg

Intervention 2

Not provided
Sites / Institutions
Not provided
Trials dates
Anticipated Start Date
Not provided
Actual Start Date
Not provided
Anticipated Date of Last Follow-up
Not provided
Estimated Primary Completion Date
Not provided
Estimated Completion Date
Not provided
Actual Primary Completion Date
Not provided
Actual Completion Date
Not provided
Studied populations
Age Cohort
Adults

Once-yearly intramuscular lenacapavir formulation 2

Dosage: 5000 mg

Countries

Accepts healthy individuals Yes
Comments about the studied populations
Not provided
Health status
Not provided
Study type
Interventional (clinical trial)
Enrollment
40
Allocation
Not provided
Intervention model
Parallel Assignment
Intervention model description
Not provided

Genders

Unspecified

Unspecified

Accepts pregnant individuals

Accepts lactating individuals

All

# Masking

Not provided

# **Masking description**

Not provided

# Frequency of administration

Yearly

# Studied LA-formulation(s)

Injectable

# Studied route(s) of administration

Intramuscular

# Use case

PrEP

# **Key resources**

Туре	Title	Content	Link
Link	Pharmacokinetics a safety of once-year lenacapavir: a phas	ly	https://doi.org/10.1016/S014
	1, open-label study		

#### **PURPOSE 1**

#### **Identifier**

NCT04994509

#### Link

https://clinicaltrials.gov/study/NCT04994509

#### Phase

Phase III

#### **Status**

Active, not recruiting

## **Sponsor**

Gilead Sciences

#### More details

The goal of this study is to evaluate the efficacy in preventing HIV infection of the study drugs, lenacapavir (LEN) and emtricitabine/tenofovir alafenamide (F/TAF), in adolescent girls and young women.

# **Purpose**

Pre-Exposure Prophylaxis Study of Lenacapavir and Emtricitabine/Tenofovir Alafenamide in Adolescent Girls and Young Women at Risk of HIV Infection

#### **Interventions**

#### Intervention 1

Oral Lenacapavir (LEN)

#### **Intervention 2**

Subcutaneous (SC) Lenacapavir (LEN)

#### **Intervention 3**

Oral F/TAF

#### Intervention 4

Oral F/TDF

#### **Intervention 5**

Placebo SC LEN

#### **Countries**

South Africa

Uganda

#### Sites / Institutions

Not provided

#### **Trials dates**

#### **Anticipated Start Date**

Not provided

#### **Actual Start Date**

2021-08-30 00:00:00

#### **Anticipated Date of Last Follow-up**

2024-02-26 00:00:00

# **Estimated Primary Completion Date**

2024-09-01 00:00:00

# **Estimated Completion Date**

2027-07-01 00:00:00

# **Actual Primary Completion Date**

Not provided

#### **Actual Completion Date**

Not provided

#### Studied populations

#### **Age Cohort**

- Children
- Adults

#### **Genders**

• Female

### Accepts pregnant individuals

Yes

#### **Accepts lactating individuals**

Yes

#### Accepts healthy individuals

Yes

# Comments about the studied populations

Key Inclusion Criteria: \* Incidence Phase \* HIV-1 status unknown at initial screening and no prior human immunodeficiency virus (HIV)-1 testing within the last 3 months. \* Sexually active (has had  $\gt$  1 vaginal intercourse within the last 3 months) with cisgender male individuals (CGM). \* Randomized Phase \* Negative fourth generation HIV-1 antibody (Ab)/antigen (Ag) test confirmed with central HIV-1 testing. \* Estimated glomerular filtration rate (GFR)  $\succeq$  60 mL/min at screening. \* Body weight  $\succeq$  35 kg. Key Exclusion Criteria: \* Prior receipt of an HIV vaccine. \* Prior use of long-acting systemic HIV pre-exposure prophylaxis (PrEP) or or HIV PEP (postexposure prophylaxis). Note: Other protocol defined Inclusion/Exclusion criteria may apply.

#### Health status

Study type
Interventional (clinical trial)
Enrollment
5368
Allocation
Randomized
Intervention model
Parallel Assignment
Intervention model description
Not provided
Masking
Double-blind masking
Masking description
Double (Participant, Investigator)
Frequency of administration
Once every 6 months
Studied LA-formulation(s)
Injectable
Studied route(s) of administration
Oral

Negative to : HIV

# Subcutaneous

# Use case

PrEP

# **Key resources**

Not provided

## **PURPOSE 2**

#### Identifier

NCT04925752

#### Link

https://clinicaltrials.gov/study/NCT04925752

#### **Phase**

Phase III

#### **Status**

Active, not recruiting

## **Sponsor**

Gilead Sciences

#### More details

The goal of this clinical study is to test how well the study drug, lenacapavir (LEN), works in preventing the risk of HIV.

# **Purpose**

Study of Lenacapavir for HIV Pre-Exposure Prophylaxis in People Who Are at Risk for HIV Infection

#### Interventions

#### **Intervention 1**

Oral Lenacapavir (LEN)

#### Intervention 2

#### Oral F/TDF

#### **Intervention 3**

Subcutaneous (SC) Lenacapavir (LEN)

#### **Intervention 4**

Placebo SC LEN

#### **Intervention 5**

Placebo to match F/TDF

#### Countries

Argentina

Brazil

Peru

Puerto Rico

South Africa

Thailand

United States of America

#### Sites / Institutions

Not provided

#### **Trials dates**

#### **Anticipated Start Date**

Not provided

#### **Actual Start Date**

2021-06-28 00:00:00

# **Anticipated Date of Last Follow-up**

2024-07-11 00:00:00

# **Estimated Primary Completion Date**

2024-12-01 00:00:00

#### **Estimated Completion Date**

2028-05-01 00:00:00

#### **Actual Primary Completion Date**

Not provided

#### **Actual Completion Date**

Not provided

# Studied populations

#### Age Cohort

- Children
- Adults
- Older Adults

#### **Genders**

All

#### **Accepts pregnant individuals**

Unspecified

#### **Accepts lactating individuals**

Unspecified

#### Accepts healthy individuals

Yes

## Comments about the studied populations

Key Inclusion Criteria: Incidence Phase \* CGM, TGW, TGM, and GNB who have condomless receptive anal sex with partners assigned male at birth and are at risk for HIV infection. \* HIV-1 status unknown at screening and no prior HIV-1 testing within the last 3 months. \* Sexually active with  $\geq$  1 partner assigned male at birth (condomless receptive anal sex) in the last 12 months and 1 of the following: \* Condomless receptive anal sex with  $\geq$  2 partners in the last 12 weeks. \* History of syphilis, rectal

gonorrhea, or rectal chlamydia in the last 24 weeks. \* Self-reported use of stimulants with sex in the last 12 weeks. Randomized Phase \* Negative local rapid fourth generation HIV-1/2 Ab/Ag, central fourth generation HIV-1/2 Ab/Ag, and HIV-1 RNA quantitative nucleic acid amplification

#### **Health status**

Negative to : HIV, HCV, HBV Considered high risk to : HIV

## Study type

Interventional (clinical trial)

#### **Enrollment**

3295

#### **Allocation**

Randomized

#### Intervention model

Parallel Assignment

# Intervention model description

Not provided

#### **Masking**

Double-blind masking

# **Masking description**

Double (Participant, Investigator)

# Frequency of administration

Once every 6 months

# Studied LA-formulation(s)

Injectable

# Studied route(s) of administration

Oral

Subcutaneous

# Use case

PrEP

# **Key resources**

Not provided

#### **PURPOSE 3**

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NCT06101329

#### Link

https://clinicaltrials.gov/study/NCT06101329

#### **Phase**

Phase II

#### **Status**

Recruiting

## **Sponsor**

Gilead Sciences

#### More details

Not provided

# **Purpose**

Evaluate the Pharmacokinetics, Safety, and Acceptability of Twice Yearly Long-acting Subcutaneous Lenacapavir for Pre-Exposure Prophylaxis in Cisgender Women in the United States.

#### **Interventions**

#### **Intervention 1**

Drug: Lenacapavir Tablet

#### Intervention 2

Drug: Long-acting Subcutaneous Lenacapavir Injection	
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#### **Intervention 3**

Drug: Emtricitabine/Tenofovir Disoproxil Fumarate (F/TDF)

#### Countries

United States of America

#### Sites / Institutions

Not provided

#### **Trials dates**

#### **Anticipated Start Date**

Not provided

#### **Actual Start Date**

2023-11-17 00:00:00

# **Anticipated Date of Last Follow-up**

2024-08-12 00:00:00

#### **Estimated Primary Completion Date**

2028-01-01 00:00:00

# **Estimated Completion Date**

2028-01-01 00:00:00

# **Actual Primary Completion Date**

Not provided

#### **Actual Completion Date**

Not provided

# Studied populations

#### Age Cohort

#### Adults

Older Adults

#### **Genders**

• Cisgender female

#### **Accepts pregnant individuals**

Unspecified

#### **Accepts lactating individuals**

Unspecified

#### Accepts healthy individuals

Unspecified

# Comments about the studied populations

Cisgender women aged 18 and older who report at least one episode of condomless vaginal or anal sex with a cisgender man in the twelve months prior to enrollment.

#### **Health status**

Considered at low risk of: HIV

Negative to : HIV, HBV

# Study type

Interventional (clinical trial)

#### **Enrollment**

250

#### Allocation

Randomized

#### Intervention model

# Parallel Assignment

# Intervention model description

Not provided

# Masking

Open label

# **Masking description**

None (Open Label)

# Frequency of administration

Once every 6 months

# Studied LA-formulation(s)

Injectable

# Studied route(s) of administration

Oral

Subcutaneous

#### Use case

PrEP

# **Key resources**

Not provided

#### **PURPOSE 4**

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NCT06101342

#### Link

https://clinicaltrials.gov/study/NCT06101342

#### **Phase**

Phase II

#### **Status**

Recruiting

## **Sponsor**

Gilead Sciences

#### More details

PWUD (People Who Use Drugs).

# **Purpose**

Evaluate the Pharmacokinetics and Safety of Twice Yearly Long-Acting Subcutaneous Lenacapavir for Pre-Exposure Prophylaxis in People Who Inject Drugs.

#### **Interventions**

#### **Intervention 1**

Drug: Long-acting Subcutaneous Lenacapavir Injection

#### **Intervention 2**

Drug: Lenacapavir Tablet

#### **Intervention 3**

Drug: Emtricitabine/tenofovir disoproxil fumarate (F/TDF)

#### Countries

United States of America

#### Sites / Institutions

Not provided

#### **Trials dates**

## **Anticipated Start Date**

Not provided

#### **Actual Start Date**

2023-12-13 00:00:00

# **Anticipated Date of Last Follow-up**

2024-08-08 00:00:00

#### **Estimated Primary Completion Date**

2027-07-01 00:00:00

#### **Estimated Completion Date**

2027-07-01 00:00:00

# **Actual Primary Completion Date**

Not provided

#### **Actual Completion Date**

Not provided

# Studied populations

#### Age Cohort

Adults

#### Older Adults

#### **Genders**

All

## Accepts pregnant individuals

Unspecified

#### **Accepts lactating individuals**

Unspecified

#### Accepts healthy individuals

Yes

# Comments about the studied populations

Participant inclusion criteria requires a positive urine drug screen for any drug of misuse including (but not limited to) opioids (eg, fentanyl, heroin), stimulants (eg, cocaine, amphetamines), psychoactive drugs (eg, benzodiazepines), or a combination of these drugs. Participants must also display evidence of recent injection(s) (eg, track marks) and self-report of injection paraphernalia sharing within the last 30 days.

#### **Health status**

Negative to : HIV, HBV, TB

Considered high risk to: HIV

# Study type

Interventional (clinical trial)

#### **Enrollment**

250

#### Allocation

Randomized

#### **PURPOSE 5**

#### Identifier

NCT06513312

#### Link

https://clinicaltrials.gov/study/NCT06513312

#### **Phase**

Phase II

#### **Status**

Not yet recruiting

#### **Sponsor**

Gilead Sciences

#### More details

The goals of this clinical study are to learn more about the study drug lenacapavir (LEN), by comparing the consistent and continuous use of LEN and emtricitabine/tenofovir disoproxil fumarate (coformulated; Truvada®) (F/TDF), then by observing the safety of LEN and F/TDF, evaluating the acceptability of LEN injections and oral F/TDF, and observe how LEN moves throughout the body in people who would benefit from pre-exposure prophylaxis (PrEP). The primary objective of this study is to compare LEN and F/TDF consistent and continuous use among people who would benefit from PrEP.

# **Purpose**

Study of Lenacapavir Taken Twice a Year for HIV Pre-Exposure Prophylaxis (PrEP)

#### **Interventions**

#### Intervention 1

Drug: Lenacapavir Injection

#### **Intervention 2**

Drug: Lenacapavir Tablet

#### **Intervention 3**

Drug: Emtricitabine/tenofovir disoproxil fumarate (F/TDF)

#### Countries

France

United Kingdom

#### Sites / Institutions

Not provided

#### **Trials dates**

# **Anticipated Start Date**

2024-09-01 00:00:00

#### **Actual Start Date**

Not provided

# **Anticipated Date of Last Follow-up**

2024-08-22 00:00:00

#### **Estimated Primary Completion Date**

2027-01-01 00:00:00

#### **Estimated Completion Date**

2029-07-01 00:00:00

#### **Actual Primary Completion Date**

Not provided

#### **Actual Completion Date**

Not provided

## Studied populations

#### Age Cohort

- Adults
- Older Adults

#### **Genders**

- All
- Cisgender female
- Cisgender male
- Transgender female
- Transgender male
- Gender non-binary

#### Accepts pregnant individuals

Unspecified

#### **Accepts lactating individuals**

Unspecified

#### Accepts healthy individuals

Yes

# Comments about the studied populations

Key Inclusion Criteria: - Able to comprehend and provide a signed written informed consent, which must be obtained prior to initiation of study procedures. - Cisgender men who have sex with men, transgender women, transgender men, cisgender women, and nonbinary people - Increased likelihood of HIV acquisition as indicated by at least one of the following: - Condomless sex with  $\geq 2$  partners in the past 6 months - Diagnosis of a bacterial sexually transmitted infection (STI) in the past 12 months - Engagement in sex work or transactional sex in the past 12 months - Use of  $\geq 2$ 

courses of nonoccupational HIV post-exposure prophylaxis (nPEP) in the past 12 months - Condomless sex with a partner living with HIV who has unknown or unsuppressed viral load ( $\geq$  200 copies/mL) in the past 12 months

#### **Health status**

Negative to: HIV

# Study type

Interventional (clinical trial)

#### **Enrollment**

262

## Allocation

Randomized

#### Intervention model

Parallel Assignment

# Intervention model description

Not provided

#### Masking

Open label

# **Masking description**

None (Open Label)

# Frequency of administration

Once every 6 months

# Studied LA-formulation(s)

Injectable

# Studied route(s) of administration

Subcutaneous

Use case

PrEP

# **Key resources**

Not provided

#### 10002211

#### **Identifier**

NCT06819176

#### Link

https://clinicaltrials.gov/study/NCT06819176

#### Phase

Phase I

#### **Status**

Not yet recruiting

## **Sponsor**

National Institute of Allergy and Infectious Diseases (NIAID)

#### More details

Treatment intensification study designed to ascertain the effects of lenacapavir intensification in people living with HIV (PLWH) with viral suppression on daily antiretroviral therapy (ART).

# **Purpose**

Lenacapavir Intensification to Disrupt HIV Reservoirs in Virologically Suppressed People Living With HIV Receiving Antiretroviral Therapy

#### **Interventions**

#### Intervention 1

Lenacapavir

#### Countries

United States of America

#### Sites / Institutions

Not provided

#### **Trials dates**

#### **Anticipated Start Date**

2025-06-05

#### **Actual Start Date**

Not provided

# **Anticipated Date of Last Follow-up**

2025-05-30

#### **Estimated Primary Completion Date**

2028-09-01

## **Estimated Completion Date**

2029-01-24

# **Actual Primary Completion Date**

Not provided

#### **Actual Completion Date**

Not provided

# **Studied populations**

# **Age Cohort**

- Adults
- Older Adults

#### **Genders**

#### Accepts pregnant individuals

Unspecified

#### Accepts lactating individuals

Unspecified

#### Accepts healthy individuals

No

#### Comments about the studied populations

\* INCLUSION CRITERIA: To be eligible to participate in this study, an individual must meet all of the following criteria: 1. Able to provide informed consent. 2. Stated willingness to comply with all study procedures and availability for the duration of the study. 3. Aged 18 years to 75 years. 4. In generally good health with an identified primary health care provider for medical management of HIV infection and willing to maintain a relationship with a primary health care provider while participating in the study. 5. Confirmed HIV-1 infection. 6. Total HIV DNA reservoir size greater than 300 copies/106 CD4+ T cells. 7. CD4+ T cell count \>200 cells/mm\^3 at screening. 8. Documentation of continuous ART treatment \>3 years with suppression of plasma viral level below the limit of quantita

#### **Health status**

Not provided

#### Study type

Interventional (clinical trial)

#### **Enrollment**

50

#### **Allocation**

Randomized
Intervention model
Parallel Assignment
Intervention model description
Not provided
Masking
Single blind masking
Masking description
Not provided
Frequency of administration
Not provided
Studied LA-formulation(s)
Not provided
Studied route(s) of administration
Not provided
Use case
Not provided
Key resources
Not provided

# **Excipients**

# Proprietary excipients used

No proprietary excipient used

Novel excipients or existing excipients at a concentration above Inactive Ingredients Database (IID) for the specified route of administration

No novel excipient or existing excipient used

#### Residual solvents used

No residual solvent used

# Patent info

# **Compound patent families**

#### **Patent informations**

	Representative		Licence with	Patent
Patent description	patent	Categories Patent holder	MPP	source
Lenacapavir use to treat multidrug resistant HIV infection in heavily treatment-experienced Expiry date: 2039-07-15	WO2020018459		Yes	

#### **Patent status**

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted		Australia, United States of America
Filed	China, Albania, Serbia, Türkiye, North Macedonia	Australia, Canada, Liechtenstein, Italy, Norway, Malta, Denmark, Belgium, United Kingdom, Greece, Netherlands, Hungary, Croatia, Switzerland, Spain, San Marino, Slovenia, Austria, Romania, Iceland, Cyprus, Finland, France, Bulgaria, Slovakia, Poland, Latvia, Ireland, Estonia, Germany, Luxembourg, Portugal, Czechia, Lithuania, Monaco, Sweden, Japan, Korea, Republic of, Taiwan, Province of China, United States of America
Not in force	World Intellectual Property Organization (WIPO), Morocco, Tunisia, Bosnia and Herzegovina, Cambodia, Montenegro, Moldova, Republic of	World Intellectual Property Organization (WIPO), Canada, Japan, Korea, Republic of

## **MPP Licence(s)**

Bilateral licence on lenacapavir (LEN)

https://www.gilead.com/-/media/gileadcorpredesign/pdf/Other/LEN-VL.pdf

#### **Patent informations**

			Licence	
	Representative		with	Patent
Patent description	patent	Categories Patent holder	MPP	source
Lenacapavir manufacturing	WO2019161280	Intermediate(s),	Yes	
processess and intermediates		Process		
Expiry date: 2039-02-15				

#### **Patent status**

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	China, Albania, Türkiye, North Macedonia, India	Australia, Liechtenstein, Italy, Norway, Malta, Denmark, Belgium, United Kingdom, Greece, Netherlands, Switzerland, Spain, Slovenia, Austria, Cyprus, Finland, France, Bulgaria, Slovakia, Poland, Latvia, Ireland, Estonia, Germany, Luxembourg, Portugal, Czechia, Sweden, Japan, Korea, Republic of, Taiwan, Province of China, United States of America, Hong Kong
Filed	China, Albania, Serbia, Türkiye, North Macedonia, India	Australia, Canada, Liechtenstein, Italy, Norway, Malta, Denmark, Belgium, United Kingdom, Greece, Netherlands, Hungary, Croatia, Switzerland, Spain, San Marino, Slovenia, Austria, Romania, Iceland, Cyprus, Finland, France, Bulgaria, Slovakia, Poland, Latvia, Ireland, Estonia, Germany, Luxembourg, Portugal, Czechia, Lithuania, Monaco, Sweden
Not in force	World Intellectual Property Organization (WIPO), Argentina, Morocco, Tunisia, Serbia, Bosnia and Herzegovina, Cambodia, Montenegro, Moldova, Republic of	World Intellectual Property Organization (WIPO), Hungary, Croatia, San Marino, Romania, Iceland, Lithuania, Monaco, Japan, Korea, Republic of, Bahamas

#### **MPP Licence(s)**

Bilateral licence on lenacapavir (LEN)

https://www.gilead.com/-/media/gileadcorpredesign/pdf/Other/LEN-VL.pdf

### **Patent informations**

			Licence	
	Representative		with	Patent
Patent description	patent	Categories Patent holder	MPP	source
Crystalline forms of Lenacapavir	WO2019035904	Polymorphs Gilead Sciences, Inc	Yes	
sodium salt				
Expiry date: 2038-08-16				
Lenacapavir solid forms, including				
pharmaceutically acceptable salts				
and cocrystals of the inhibitor, as				
well as crystalline forms of the salts				
and cocrystals, for use in the				
treatment of a Retroviridae viral				
infection including an infection				
caused by the HIV virus. The				
present disclosure also relates to				
pharmaceutical compositions				
containing the novel salts,				
cocrystals, and crystalline forms				
thereof, and methods of treating or				
preventing a Retroviridae viral				
infection.				

### **Patent status**

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	Türkiye, North Macedonia, Albania	Belgium, Germany, France,
		Luxembourg, Netherlands, Switzerland,
		United Kingdom, Sweden, Italy, Austria,
		Liechtenstein, Greece, Spain, Denmark,
		Monaco, Portugal, Ireland, Finland,
		Cyprus, Bulgaria, Czechia, Estonia,
		Slovakia, Hungary, Poland, Malta,
		Norway, San Marino, Romania, Latvia,
		Lithuania, Slovenia, Australia, Canada,
		Japan, Korea, Republic of, Taiwan,
		Province of China, United States of
		America, Hong Kong

Patent status/countries	Low, Low- middle and upper-middle	High income
Filed	Türkiye, North Macedonia, Albania,	Belgium, Germany, France,
	Serbia, China, India	Luxembourg, Netherlands, Switzerland,
		United Kingdom, Sweden, Italy, Austria,
		Liechtenstein, Greece, Spain, Denmark,
		Monaco, Portugal, Ireland, Finland,
		Cyprus, Bulgaria, Czechia, Estonia,
		Slovakia, Hungary, Poland, Iceland,
		Malta, Norway, San Marino, Croatia,
		Romania, Latvia, Lithuania, Slovenia,
		Canada, United States of America, Hong
		Kong
Not in force	World Intellectual Property Organization	World Intellectual Property Organization
	(WIPO), North Macedonia, Albania,	(WIPO), Luxembourg, Denmark, Monaco,
	Bosnia and Herzegovina, Montenegro,	Finland, Cyprus, Bulgaria, Estonia,
	Serbia, Moldova, Republic of, Morocco,	Hungary, Iceland, Malta, San Marino,
	Tunisia, Cambodia, Argentina,	Croatia, Romania, Latvia, Lithuania,
	Bangladesh	Japan, Taiwan, Province of China

### **MPP Licence(s)**

Bilateral licence on lenacapavir (LEN)

https://www.gilead.com/-/media/gileadcorpredesign/pdf/Other/LEN-VL.pdf

### **Patent informations**

			Licence	
	Representative		with	Patent
Patent description	patent	Categories Patent holder	MPP	source
Lenacapavir compound and its use	WO2018035359	Compound Gilead Sciences, In	c Yes	
in HIV (oral and parenteral)				
Expiry date: 2037-08-17				
The present disclosure relates to				
novel compounds for use in the				
treatment of a Retroviridae viral				
infection including an infection				
caused by the HIV virus. The				
present disclosure also relates to				
intermediates for its preparation				
and to pharmaceutical compositions				
containing said novel compound.				

#### **Patent status**

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	Türkiye, North Macedonia, Morocco,	Belgium, Germany, France,
	Brazil, China, Colombia, Dominican	Luxembourg, Netherlands, Switzerland,
	Republic, Turkmenistan, Belarus,	United Kingdom, Sweden, Italy, Austria
	Tajikistan, Kazakhstan, Azerbaijan,	Liechtenstein, Greece, Spain, Denmark
	Kyrgyzstan, Armenia, Mexico, Peru,	Portugal, Ireland, Finland, Cyprus,
	Philippines, Botswana, Gambia (the),	Bulgaria, Czechia, Estonia, Slovakia,
	Ghana, Kenya, Lesotho, Malawi,	Hungary, Poland, Iceland, Malta,
	Mozambique, Namibia, Sierra Leone,	Norway, Croatia, Romania, Latvia,
	Liberia, Sao Tome and Principe, Sudan,	Lithuania, Slovenia, Australia, Canada,
	Eswatini, Tanzania, United Republic of,	Russian Federation, Hong Kong, Israel,
	Zambia, Zimbabwe, Indonesia,	Japan, Korea, Republic of, New Zealand
	Malaysia, Ukraine, South Africa,	Singapore, Taiwan, Province of China,
	Uzbekistan	United States of America, Bahamas,
		Bahrain, Kuwait, Qatar, Saudi Arabia,
		Oman, United Arab Emirates, Macao,
		Panama

Patent status/countries	Low, Low- middle and upper-middle	High income	
Filed	Türkiye, North Macedonia, Albania, Serbia, Morocco, Argentina, Jordan, Philippines, India, Uganda, Egypt, Guatemala, Indonesia, Nigeria, Thailand, Ukraine, Viet Nam	Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Spain, Denmark, Monaco, Portugal, Ireland, Finland, Cyprus, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, San Marino, Croatia, Romania, Latvia, Lithuania, Slovenia, Australia, Costa Rica, Hong Kong, Japan, Singapore, Taiwan, Province of China, United States of America, Saudi Arabia, Panama	
Not in force	World Intellectual Property Organization (WIPO), North Macedonia, Albania, Bosnia and Herzegovina, Montenegro, Serbia, Moldova, Republic of, Argentina, Colombia, Dominican Republic, Ecuador, Peru, Rwanda, Uganda, Bangladesh, Bolivia (Plurinational State of), Cuba, Egypt, Benin, Cameroon, Burkina Faso, Chad, Guinea-Bissau, Comoros, Mali, Senegal, Congo, Guinea, Gabon, Niger, Equatorial Guinea, Mauritania, Togo, Côte d'Ivoire, Central African Republic,	World Intellectual Property Organization (WIPO), Monaco, Malta, San Marino, Chile, Japan, Korea, Republic of, Uruguay, Trinidad and Tobago	

### **MPP Licence(s)**

Bilateral licence on lenacapavir (LEN)

https://www.gilead.com/-/media/gileadcorpredesign/pdf/Other/LEN-VL.pdf

Pakistan, Paraguay, El Salvador, Venezuela (Bolivarian Republic of)

### **Patent informations**

	Danuarantativa		Licence	Datast
Patent description	Representative patent	Categories Patent holder	with MPP	Patent source
Lenacapavir and analogues	WO2014134566	Compound Gilead Sciences, Inc	Yes	
(Markush formula) and their use in HIV				
Expiry date: 2034-02-28				
Compounds of formula (I) or salts				
thereof are disclosed. Also disclosed				
are pharmaceutical compositions				
comprising a compound of formula				
l, processes for preparing				
compounds of formula I,				
intermediates useful for preparing				
compounds of formula I and				
therapeutic methods for treating a				
Retroviridae viral infection including				
an infection caused by the HIV				
virus.				

### **Patent status**

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	Türkiye, North Macedonia, Albania,	Belgium, Germany, France,
	Bosnia and Herzegovina, Montenegro,	Luxembourg, Netherlands, Switzerland,
	Serbia, Brazil, China, Cuba,	United Kingdom, Sweden, Italy, Austria,
	Turkmenistan, Belarus, Tajikistan,	Liechtenstein, Greece, Spain, Denmark,
	Kazakhstan, Azerbaijan, Kyrgyzstan,	Monaco, Portugal, Ireland, Finland,
	Armenia, Mexico, Peru, Philippines,	Cyprus, Bulgaria, Czechia, Estonia,
	Ukraine, Botswana, Gambia (the),	Slovakia, Hungary, Poland, Iceland,
	Ghana, Kenya, Lesotho, Malawi,	Malta, Norway, San Marino, Croatia,
	Mozambique, Namibia, Sierra Leone,	Romania, Latvia, Lithuania, Slovenia,
	Liberia, Rwanda, Sudan, Eswatini,	Australia, Canada, Chile, Costa Rica,
	Tanzania, United Republic of, Zambia,	Russian Federation, Hong Kong, Israel,
	Zimbabwe, Benin, Cameroon, Burkina	Japan, Korea, Republic of, New Zealand
	Faso, Chad, Guinea-Bissau, Comoros,	Singapore, Taiwan, Province of China,
	Mali, Senegal, Congo, Guinea, Gabon,	United States of America, Bahrain,
	Niger, Equatorial Guinea, Mauritania,	Kuwait, Qatar, Saudi Arabia, Oman,
	Togo, Côte d'Ivoire, Central African	United Arab Emirates, Macao, Panama
	Republic, Colombia, Indonesia, Malaysia,	
	Viet Nam, South Africa	

Patent status/countries	Low, Low- middle and upper-middle	High income
Filed	Türkiye, North Macedonia, Albania, Serbia, Argentina, Ukraine, India, Egypt, Thailand	Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Spain, Denmark, Monaco, Portugal, Ireland, Finland, Cyprus, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, San Marino, Croatia, Romania, Latvia, Lithuania, Slovenia, Costa Rica, United States of America
Not in force	World Intellectual Property Organization (WIPO), North Macedonia, Albania, Bosnia and Herzegovina, Montenegro, Serbia, Argentina, Brazil, China, Moldova, Republic of, Peru, Uganda, Bolivia (Plurinational State of), Colombia, Ecuador, Malaysia, Paraguay, Pakistan, El Salvador, Venezuela (Bolivarian Republic of), Viet Nam, South Africa	World Intellectual Property Organization (WIPO), Luxembourg, Denmark, Monaco, Finland, Cyprus, Bulgaria, Estonia, Malta, San Marino, Croatia, Romania, Latvia, Lithuania, Australia, Canada, Hong Kong, Japan, New Zealand, Singapore, United States of America, Uruguay, Bahamas

# MPP Licence(s)

Bilateral licence on lenacapavir (LEN)

https://www.gilead.com/-/media/gileadcorpredesign/pdf/Other/LEN-VL.pdf

# **Supporting material**

# **Publications**

Link JO, Rhee MS, Tse WC, Zheng J, Somoza JR, Rowe W, Begley R, Chiu A, Mulato A, Hansen D, Singer E, Tsai LK, Bam RA, Chou CH, Canales E, Brizgys G, Zhang JR, Li J, Graupe M, Morganelli P, Liu Q, Wu Q, Halcomb RL, Saito RD, Schroeder SD, Lazerwith SE, Bondy S, Jin D, Hung M, Novikov N, Liu X, Villasenor AG, Cannizzaro CE, Hu EY, Anderson RL, Appleby TC, Lu B, Mwangi J, Liclican A, Niedziela-Majka A, Papalia GA, Wong MH, Leavitt SA, Xu Y, Koditek D, Stepan GJ, Yu H, Pagratis N, Clancy S, Ahmadyar S, Cai TZ, Sellers S, Wolckenhauer SA, Ling J, Callebaut C, Margot N, Ram RR, Liu YP, Hyland R, Sinclair GI, Ruane PJ, Crofoot GE, McDonald CK, Brainard DM, Lad L, Swaminathan S, Sundquist WI, Sakowicz R, Chester AE, Lee WE, Daar ES, Yant SR, Cihlar T: Clinical targeting of HIV capsid protein with a long-acting small molecule. Nature. 2020 Aug;584(7822):614-618. doi: https://doi.org/10.1038/s41586-020-2443-1. Epub 2020 Jul 1.

Oral antiretroviral agents provide life-saving treatments for millions of people living with HIV, and can prevent new infections via pre-exposure prophylaxis1-5. However, some people living with HIV who are heavily treatment-experienced have limited or no treatment options, owing to multidrug resistance6. In addition, suboptimal adherence to oral daily regimens can negatively affect the outcome of treatment-which contributes to virologic failure, resistance generation and viral transmission-as well as of pre-exposure prophylaxis, leading to new infections 1, 2, 4, 7-9. Long-acting agents from new antiretroviral classes can provide much-needed treatment options for people living with HIV who are heavily treatment-experienced, and additionally can improve adherence10. Here we describe GS-6207, a small molecule that disrupts the functions of HIV capsid protein and is amenable to long-acting therapy owing to its high potency, low in vivo systemic clearance and slow release kinetics from the subcutaneous injection site. Drawing on X-ray crystallographic information, we designed GS-6207 to bind tightly at a conserved interface between capsid protein monomers, where it interferes with capsid-protein-mediated interactions between proteins that are essential for multiple phases of the viral replication cycle. GS-6207 exhibits antiviral

activity at picomolar concentrations against all subtypes of HIV-1 that we tested, and shows high synergy and no cross-resistance with approved antiretroviral drugs. In phase-1 clinical studies, monotherapy with a single subcutaneous dose of GS-6207 (450 mg) resulted in a mean log10-transformed reduction of plasma viral load of 2.2 after 9 days, and showed sustained plasma exposure at antivirally active concentrations for more than 6 months. These results provide clinical validation for therapies that target the functions of HIV capsid protein, and demonstrate the potential of GS-6207 as a long-acting agent to treat or prevent infection with HIV.

Bester SM, Wei G, Zhao H, Adu-Ampratwum D, Iqbal N, Courouble VV, Francis AC, Annamalai AS, Singh PK, Shkriabai N, Van Blerkom P, Morrison J, Poeschla EM, Engelman AN, Melikyan GB, Griffin PR, Fuchs JR, Asturias FJ, Kvaratskhelia M: Structural and mechanistic bases for a potent HIV-1 capsid inhibitor. Science. 2020 Oct 16;370(6514):360-364. doi: https://doi.org/10.1126/science.abb4808

The potent HIV-1 capsid inhibitor GS-6207 is an investigational principal component of long-acting antiretroviral therapy. We found that GS-6207 inhibits HIV-1 by stabilizing and thereby preventing functional disassembly of the capsid shell in infected cells. X-ray crystallography, cryo-electron microscopy, and hydrogen-deuterium exchange experiments revealed that GS-6207 tightly binds two adjoining capsid subunits and promotes distal intra- and inter-hexamer interactions that stabilize the curved capsid lattice. In addition, GS-6207 interferes with capsid binding to the cellular HIV-1 cofactors Nup153 and CPSF6 that mediate viral nuclear import and direct integration into gene-rich regions of chromatin. These findings elucidate structural insights into the multimodal, potent antiviral activity of GS-6207 and provide a means for rationally developing second-generation therapies.

# **Additional documents**

No documents were uploaded

# **Useful links**

- Gilead Announces First Global Regulatory Approval of Sunlenca® (Lenacapavir)
- PURPOSE PrEP Studies Overview: Prevention With a Purpose
- An Overview of Lenacapavir for PrEP Trials (AVAC)
- Pharmacokinetics and safety of once-yearly lenacapavir: a phase 1, open-label study
- Capsid Inhibition with Lenacapavir in Multidrug-Resistant HIV-1 Infection
- CROI 2022: Lenacapavir: 54 week results in treatment-naive participants of CALIBRATE study
- Lenacapavir administered every 26 weeks or daily in combination with oral daily antiretroviral therapy for initial treatment of HIV: a randomised, open-label, activecontrolled, phase 2 trial
- Interim Resistance Analysis of Long-Acting Lenacapavir in Treatment-Naïve People
   with HIV at 28 Weeks
- Clinical targeting of HIV capsid protein with a long-acting small molecule

# **Access principles**

## Collaborate for development



Consider on a case by case basis, collaborating on developing long acting products with potential significant public health impact, especially for low- and middle-income countries (LMICs), utilising the referred to long-acting technology

Not provided

## **Share technical information for match-making assessment**



Provide necessary technical information to a potential partner, under confidentiality agreement, to enable preliminary assessment of whether specific medicines of public health importance in LMICs might be compatible with the referred to long-acting technology to achieve a public health benefit

Not provided

# Work with MPP to expand access in LMICs



In the event that a product using the referred to long-acting technology is successfully developed, the technology IP holder(s) will work with the Medicines Patent Pool towards putting in place the most appropriate strategy for timely and affordable access in low and middle-income countries, including through licensing

Not provided

# **Comment & Information**

Not provided