




**Australian Government**  
**Department of Health and Aged Care**  
Therapeutic Goods Administration

## Public Summary

<b>Summary for ARTG Entry:</b>	377442	APRETUDE cabotegravir (as sodium) 30 mg film coated tablet bottle
<b>ARTG entry for</b>	Medicine Registered	
<b>Sponsor</b>	ViiV Healthcare Pty Ltd	
<b>Postal Address</b>	PO Box 18095, MELBOURNE CITY MC, VIC, 8001 Australia	
<b>ARTG Start Date</b>	11/08/2022	
<b>Product Category</b>	Medicine	
<b>Status</b>	Active	
<b>Approval Area</b>	Drug Safety Evaluation Branch	

  
**Medicine under  
additional monitoring**

### Conditions

Conditions applicable to the relevant category and class of therapeutic goods as specified in the document "Conditions Applying to Therapeutic Goods Accepted for Registration or Listing as Goods Currently Supplied at the Commencement of the Therapeutic Goods Act 1989"

Conditions applicable to all therapeutic goods as specified in the document "Standard Conditions Applying to Registered or Listed Therapeutic Goods Under Section 28 of the Therapeutic Goods Act 1989" effective 1 February 1992, other than condition No. 8 and condition No. 9.

Conditions applicable to the relevant category and class of therapeutic goods as specified in the document "Standard Conditions Applying to Registered or Listed Therapeutic Goods Under Section 28 of the Therapeutic Goods Act 1989" effective 1 February 1992, other than condition No. 11.

Except where the sponsor has been contracted by an overseas party to manufacturer the goods and that party will be responsible for placing the goods on the market in countries other than Australia, the sponsor must have and shall retain, while the goods remain listed, evidence necessary to substantiate and support the accuracy of the indications in relation to the listed goods, and upon the request of the Head, Office of Prescription Medicines Authorisation Branch, Therapeutic Goods Administration, shall produce such evidence to this officer.

The conditions applying to these goods when they are exported from Australia are given below:

Conditions applicable to all therapeutic goods as specified in the document "Standard Conditions Applying to Registered or Listed Therapeutic Goods Under Section 28 of the Therapeutic Goods Act 1989" effective 1 July 1995, other than condition No. 8 and condition No. 9.

Conditions applicable to the relevant category and class of therapeutic goods as specified in the document "Standard Conditions Applying to Registered or Listed Therapeutic Goods Under Section 28 of the Therapeutic Goods Act 1989" effective 1 July 1995, other than condition No.11

Conditions applicable to all therapeutic goods as specified in the document "Standard Conditions Applying to Registered or Listed Therapeutic Goods Under Section 28 of the Therapeutic Goods Act 1989" effective 1 July 1995.

Conditions applicable to the relevant category and class of therapeutic goods as specified in the document "Standard Conditions Applying to Registered or Listed Therapeutic Goods Under Section 28 of the Therapeutic Goods Act 1989" effective 1 July 1995.

The therapeutic goods grouped in this registration/listing as export only goods under the following names are included in the Australian Register of Therapeutic Goods and may not be supplied in Australia. Export Name(s) to be added:

### Products

#### 1 . APRETUDE cabotegravir (as sodium) 30 mg film coated tablet bottle

<b>Product Type</b>	Single Medicine Product	<b>Effective Date</b>	18/09/2023
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#### Permitted Indications

No Permitted Indications included on Record

#### Indication Requirements

No Indication Requirements included on Record

#### Standard Indications

No Standard Indications included on Record

#### Specific Indications

APRETUDE is indicated in at-risk adults and adolescents (at least 12 years of age) and weighing at least 35 kg for pre-exposure prophylaxis (PrEP) to reduce the risk of sexually acquired HIV-1 infection..APRETUDE tablets may be used as an oral lead-in to assess tolerability of cabotegravir prior to administration of cabotegravir injections or as short-term oral PrEP in individuals who will miss planned dosing with cabotegravir injections..Individuals must have a documented negative HIV-1 test prior to initiating APRETUDE for HIV-1 PrEP.

#### Warnings

See Product Information and Consumer Medicine Information for this product

Public Summary



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**Additional Product information**

*This product is included in the Black Triangle Scheme*

**Container information**

Type	Material	Life Time	Temperature	Closure	Conditions
Bottle	HDPE	36 Months	Store below 30 degrees Celsius	Child resistant closure	Not recorded

**Pack Size/Poison information**

Pack Size	Poison Schedule
30	(S4) Prescription Only Medicine

**Components**

**1 . APRETUDE cabotegravir 30 mg film-coated tablet bottle**

<b>Dosage Form</b>	Tablet, film coated
<b>Route of Administration</b>	Oral
<b>Visual Identification</b>	White, oval, film-coated, tablets, debossed with SV CTV on one side

**Active Ingredients**

<b>cabotegravir sodium</b>	<b>31.62 mg</b>
Equivalent: cabotegravir	30 mg

**Other Ingredients (Excipients)**

hypromellose  
lactose monohydrate  
macrogol 3350  
magnesium stearate  
microcrystalline cellulose  
sodium starch glycollate type A  
titanium dioxide

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Public Summary