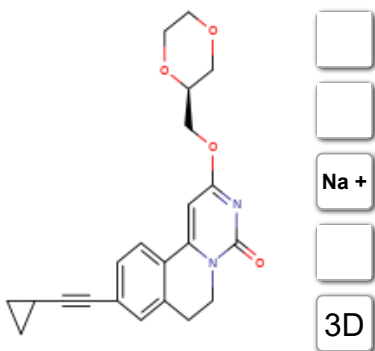


TOXNET > [ChemIDplus](#) > Substance

# ChemIDplus

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 equals  
**Substance Name: GLPG-1205**

RN: 1445847-37-9

UNII: [K9WR6LRA5D](#)

InChIKey: IRBAWVGZNIROV-SFHVURJKSA-N

**Molecular Formula**C<sub>22</sub>H<sub>22</sub>N<sub>2</sub>O<sub>4</sub>**Molecular Weight**

378.4258

\* denotes mobile formatted website

**Links to Resources**[DrugPortal](#)Search for this InChIKey on the [Web](#)**Names and Synonyms****Name of Substance**

GLPG-1205

**Synonyms**

4H-Pyrimido(6,1-a)isoquinolin-4-one, 9-(2-cyclopropylethynyl)-2-((2S)-1,4-dioxan-2-ylmethoxy)-6,7-dihydro-  
 GLPG-1205  
 GLPG1205  
 UNII-K9WR6LRA5D

**Registry Numbers****CAS Registry Number**

1445847-37-9

**FDA UNII**

K9WR6LRA5D

**System Generated Number**

1445847379

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**Structure Descriptors**

**InChI**

1S/C22H22N2O4/c25-22-23-21(28-14-18-13-26-9-10-27-18)12-20-19-6-5-16(4-3-15-1-2-15)11-17(19)7-8-24(20)22/h5-6,11-12,15,18H,1-2,7-10,13-14H2/t18-/m0/s1

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**InChIKey**

IRBAWVGZNIROV-SFHVURJKSA-N

[Search the web for this InChIKey](#)

**Smiles**

O=C1N=C(OC[C@@H]2COCCO2)C=C3N1CCc4cc(ccc34)C#CC5CC5

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