# Data Sheet (Cat.No.T27499)



#### GSK-A1

### **Chemical Properties**

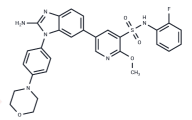
CAS No.: 1416334-69-4

Formula: C29H27FN6O4S

Molecular Weight: 574.63

Appearance: no data available

Storage: Powder: -20°C for 3 years | In solvent: -80°C for 1 year



## **Biological Description**

Description	GSK-A1 is a specific inhibitor of PI4KA, also inhibited calcium-driven PRL secretion without affecting calcium signaling and Prl expression. GSK-A1 has anti-hepatitis C virus (HCV) potential.
Targets(IC50)	HCV Protease,PI4K
In vitro	GSK-A1 is a selective?type III phosphatidylinositol 4-kinase PI4KA (PI4KIIIα)?inhibitor with a?pIC50?of 8.5-9.8[3]. GSK-A1 potently decreases the levels of PtdIns(4)P with a negligible effect on PtdIns(4,5)P2[3].

## **Preparing Stock Solutions**

	1mg	5mg	10mg	
1 mM	1.7403 mL	8.7013 mL	17.4025 mL	
5 mM	0.3481 mL	1.7403 mL	3.4805 mL	
10 mM	0.174 mL	0.8701 mL	1.7403 mL	
50 mM	0.0348 mL	0.174 mL	0.3481 mL	

Please select the appropriate solvent to prepare the stock solution, according to the solubility of the product in different solvents. Please use it as soon as possible.

#### Reference

Beebe SJ, Reimann EM, Schlender KK. Purification and characterization of a cAMP- and Ca2+-calmodulin-independent glycogen synthase kinase from porcine renal cortex. J Biol Chem. 1984 Feb 10;259(3):1415-22.

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