Data Sheet (Cat.No.T13786)



MZP-55

Chemical Properties

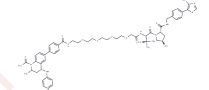
CAS No.: 2010159-48-3

Formula: C57H70ClN7O10S

Molecular Weight: 1080.73

Appearance: no data available

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



Biological Description

Description	MZP-55 is a selective BRD3/4 degrader based on PROTAC technology(Brd4BD2 with Kd of 8 nM)
Targets(IC50)	Epigenetic Reader Domain,PROTACs
In vitro	MZP-55 binds to VHL-EloC-EloB protein (VCB) with a Kd of 105 \pm 24 nM. MZP-55 shows an inhibitory activity against MV4;11 and HL60 cells, with pEC50s of 7.31 \pm 0.03 and 6.57 \pm 0.02, respectively[1].

Solubility Information

Solubility

DMSO: 50 mg/mL (46.27 mM), Sonication is recommended.

(< 1 mg/ml refers to the product slightly soluble or insoluble)

Preparing Stock Solutions

	1mg	5mg	10mg	
1 mM	0.9253 mL	4.6265 mL	9.253 mL	
5 mM	0.1851 mL	0.9253 mL	1.8506 mL	
10 mM	0.0925 mL	0.4627 mL	0.9253 mL	
50 mM	0.0185 mL	0.0925 mL	0.1851 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

Chan KH, et al. Impact of Target Warhead and Linkage Vector on Inducing Protein Degradation: Comparison of Bromodomain and Extra-Terminal (BET) Degraders Derived from Triazolodiazepine (JQ1) and Tetrahydroquinoline

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Tel:781-999-4286 E_mail:info@targetmol.com Address:36 Washington Street,Wellesley Hills,MA 02481

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