Product Data Sheet

SJPYT-195

Cat. No.:HY-150616CAS No.:2973762-16-0Molecular Formula: $C_{35}H_{34}N_6O_8$ Molecular Weight:666.68

Target: Molecular Glues

Pathway: PROTAC

Storage: Powder -20°C 3 years

4°C 2 years

In solvent -80°C 6 months

-20°C 1 month

SOLVENT & SOLUBILITY

In Vitro

DMSO: 50 mg/mL (75.00 mM; ultrasonic and warming and heat to 60°C)

Preparing Stock Solutions	Solvent Mass Concentration	1 mg	5 mg	10 mg
	1 mM	1.5000 mL	7.4999 mL	14.9997 mL
	5 mM	0.3000 mL	1.5000 mL	2.9999 mL
	10 mM	0.1500 mL	0.7500 mL	1.5000 mL

Please refer to the solubility information to select the appropriate solvent.

In Vivo

- 1. Add each solvent one by one: 10% DMSO >> 90% (20% SBE- β -CD in saline) Solubility: 2.08 mg/mL (3.12 mM); Suspended solution; Need ultrasonic
- 2. Add each solvent one by one: 10% DMSO >> 40% PEG300 >> 5% Tween-80 >> 45% saline Solubility: ≥ 1.67 mg/mL (2.50 mM); Clear solution
- Add each solvent one by one: 10% DMSO >> 90% corn oil Solubility: ≥ 1.67 mg/mL (2.50 mM); Clear solution

BIOLOGICAL ACTIVITY

Description

SJPYT-195 is a cytotoxic GSPT1 degrader and can be used for PROTAC synthesis^[1].

In Vitro

SJPYT-195 (24 h) potently and efficaciously reduces endogenous PXR (pregnane X receptor) protein in the colorectal SNU-C4 cell line (SNU-C4 3xFLAG-PXR KI cells), with a half maximal degradation concentration (DC₅₀) of 310 \pm 130 nM and maximum degradation efficacy (D_{Max}) of 85 \pm 1%^[1].

SJPYT-195 reduces PXR protein through the degradation of GSPT1^[1].

 ${\tt MCE}\ has\ not\ independently\ confirmed\ the\ accuracy\ of\ these\ methods.\ They\ are\ for\ reference\ only.$

REFERENCES					
	: A Designed Nuclear Recep	tor Degrader That Functions as	a Molecular Glue Degrader of G	SPT1. ACS Medicinal Chemistry l	etters, 2022.
		not been fully validated for r			
	Tel: 609-228-6898 Address:	Fax: 609-228-5909 1 Deer Park Dr, Suite Q, Monr	E-mail: tech@MedCh		
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