# Data Sheet (Cat.No.T4453)



## JD-5037

### **Chemical Properties**

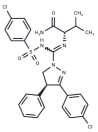
CAS No.: 1392116-14-1

Formula: C27H27Cl2N5O3S

Molecular Weight: 572.51

Appearance: no data available

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



## **Biological Description**

Description	JD-5037 is a novel, peripherally restricted CB1R antagonist with an IC50 of 1.5 nM.			
Targets(IC50)	Cannabinoid Receptor			
In vivo	JD5037, when administered at a dosage of 3 mg/kg/day intraperitoneally (i.p.), effectively induces uniform reductions in body weight and mitigates high-fat diet (HFD)-induced hyperglycemia, hepatic injury, and steatosis in obese Magel2-null mice. Likewise, oral administration of JD5037 (3 mg/kg/day, p.o.) significantly diminishes tumor size and eliminates tumors in DEN-treated mice. Moreover, JD5037 reduces anandamide (AEA) levels in hepatocellular carcinoma (HCC) samples from mice.			
Animal Research	Mice: JD-5037 is formulated in vehicle (V; 1% Tween80, 4% DMSO, 95% Saline). Obese mice are treated chronically (28 d) with vehicle (V; 1% Tween80, 4% DMSO, 95% Saline), JD5037, or SLV319 at a dose of 3 mg/kg, i.p. Body weight and food intake are monitored daily. Mice are euthanized by cervical dislocation under anesthesia; the brain, hypothalamus, liver, and combined fat pads are removed, weighed, and snap-frozen, and trunk blood is collected for determining the endocrine and biochemical parameters			

### **Solubility Information**

Solubility	DMSO: 50 mg/mL (87.33 mM),
	(< 1 mg/ml refers to the product slightly soluble or insoluble)

Page 1 of 2 www.targetmol.com

#### **Preparing Stock Solutions**

	1mg	5mg	10mg
1 mM	1.7467 mL	8.7335 mL	17.4669 mL
5 mM	0.3493 mL	1.7467 mL	3.4934 mL
10 mM	0.1747 mL	0.8733 mL	1.7467 mL
50 mM	0.0349 mL	0.1747 mL	0.3493 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

Tan S, Liu H, Ke B, et al. The peripheral CB1 receptor antagonist JD5037 attenuates liver fibrosis via a CB1 receptor/β-arrestin1/ Akt pathway. The peripheral CB1 receptor antagonist JD5037 attenuates liver fibrosis via a

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Page 2 of 2 www.targetmol.com