# Data Sheet (Cat.No.T13018)



#### SU3327

## **Chemical Properties**

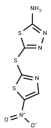
CAS No.: 40045-50-9

Formula: C5H3N5O2S3

Molecular Weight: 261.3

Appearance: no data available

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



# **Biological Description**

Description SU3327 (halicin) is a potent, selective and substrate-competitive inhibitor o 0.7 $\mu$ M).		
Targets(IC50)	JNK	
In vitro	TNF-α stimulated phosphorylation of c-Jun in HeLa cells inhibited by SU3327 with EC50 of 6.23 μM[1]. SU3327 (25 nM) pretreatment of human-derived cerebral microvascular endothelial cells (hCMEC/D3) effectively reduces LPS-induced polymorphonuclear leukocytes (PMN) rolling/adhesion to hCMEC/D3, prevents activation of AP-1, and significantly reduces expression of VCAM-1[3].	
In vivo	In male BKS.Cg-+Leprdb/+Leprdb/OlaHsd db/db mice, SU3327 (25 mg/kg; intraperitoneal injection; ) treatment possesses the ability to restore insulin sensitivity in mice models of diabetes[1]. SU3327 has favorable microsomal and plasma stability (T1/2 = 27 min)[1].	

# **Solubility Information**

DMSO: 62.5 mg/mL (239.19 mM), Sonication is recommended.	
(< 1 mg/ml refers to the product slightly soluble or insoluble)	

#### **Preparing Stock Solutions**

	1mg	5mg	10mg
1 mM	3.827 mL	19.1351 mL	38.2702 mL
5 mM	0.7654 mL	3.827 mL	7.654 mL
10 mM	0.3827 mL	1.9135 mL	3.827 mL
50 mM	0.0765 mL	0.3827 mL	0.7654 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.

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## Reference

De SK, et al. Design, synthesis, and structure-activity relationship of substrate competitive, selective, and in vivo active triazole and thiadiazole inhibitors of the c-Jun N-terminal kinase. J Med Chem. 2009 Apr 9;52(7):1943-52.



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