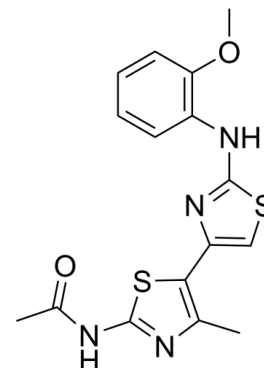


Data Sheet

Product Name:	JNJ0966
Cat. No.:	CS-0027987
CAS No.:	315705-75-0
Molecular Formula:	C ₁₆ H ₁₆ N ₄ O ₂ S ₂
Molecular Weight:	360.45
Target:	MMP
Pathway:	Metabolic Enzyme/Protease
Solubility:	DMSO : 100 mg/mL (ultrasonic)



BIOLOGICAL ACTIVITY:

JNJ0966 is a highly selective **MMP-9 zymogen** inhibitor with an **IC₅₀** of 440 nM. IC₅₀ & Target: IC₅₀: 440 nM (MMP-9 zymogen)^[1] *In Vitro*: JNJ0966 is a highly selective MMP-9 zymogen inhibitor with an IC₅₀ of 440 nM. The activations of proMMP-1, proMMP-2, and proMMP-3 are not significantly different in the presence or absence of 10 μM JNJ0966, whereas proMMP-9 activation by trypsin is significantly attenuated by JNJ0966. The addition of JNJ0966 to the reaction results in a significant reduction in fully processed MMP-9 and an apparent accumulation of the intermediate species^[1]. *In Vivo*: The exposures of JNJ0966 are dose-dependent, with plasma and brain concentrations for the 10-mg/kg dose of 77.5±31.1 ng/mL (215 nM) and 481.6±162.5 ng/g (~1336 nM), respectively, whereas the 30-mg/kg dose achieves 293.6±118.4 ng/mL (815 nM) in plasma and 1394.0±649.1 ng/g (~3867 nM) in brain. JNJ0966 is preferentially partitioned in brain, with brain/plasma ratios of 6.2 for the 10-mg/kg dose and 4.7 for the 30-mg/kg dose^[1].

PROTOCOL (Extracted from published papers and Only for reference)

Cell Assay: ^[1]HT1080 cells are added (**13,000 cells/well**) in 10 μL of serum-free medium to the top chamber with JNJ0966. The bottom feeder tray is filled with serum-free medium supplemented with 6% fetal bovine serum and JNJ0966 and incubated at 5% CO₂, 37°C for 24 h. Calcein AM is added to the feeder layer to label cells, and fluorescence intensity from cells that have migrated to the bottom layer is quantified. Mean percentage inhibition of invasion and IC₅₀ values are calculated. Images of migrated cells are acquired on an inverted microscope with a digital camera^[1]. **Animal Administration:** JNJ0966 is dissolved in 20% hydroxypropyl-β-cyclodextrin.^[1] Encephalomyelitis (EAE) studies are conducted in **female C57Bl/6 mice** age 6 to 8 weeks. Mice are randomly assigned to different groups. Vehicle-treated animals receive 20% hydroxypropyl-β-cyclodextrin via oral gavage. JNJ0966 is dissolved in 20% hydroxypropyl-β-cyclodextrin, such that animals receive **10 or 30 mg of drug per kg of body weight (mg/kg) of JNJ0966** which administered by **oral gavage**. All groups are **dosed twice daily every day until day 17** of the study, at which point animals are sacrificed, and plasma and brain tissue are collected to analyze JNJ0966 levels using mass spectroscopy^[1].

References:

[1]. Scannevin RH, et al. Discovery of a highly selective chemical inhibitor of matrix metalloproteinase-9 (MMP-9) that allosterically inhibits zymogen activation. J Biol Chem. 2017 Oct 27;292(43):17963-17974.

CAIndexNames:

Acetamide, N-[2-[(2-methoxyphenyl)amino]-4'-methyl[4,5'-bithiazol]-2'-yl]-

SMILES:

CC1=C(C2=CSC(NC3=C(OC)C=CC=C3)=N2)SC(NC(C)=O)=N1

Caution: Product has not been fully validated for medical applications. For research use only.

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