



Y-27632 (dihydrochloride)

Catalog No: tcsc0878

Available Sizes	
Size: 5mg	
Size: 10mg	
Size: 50mg	
Size: 100mg	
Size: 200mg	
Size: 500mg	
Size: 1g	
Size: 2g	
Specifications	
CAS No: 129830-38-2	
Formula: C ₁₄ H ₂₃ Cl ₂ N ₃ O	
Pathway: Apoptosis; Cell Cycle/DNA Damage; Cytoskeleton; Stem Cell/Wnt; TGF-beta/Smad	
Target: Apoptosis; ROCK	
Form: White to off-white (Solid)	
Purity / Grade:	



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98.01%

Solubility:

DMSO: 33.33 mg/mL (104.07 mM; Need ultrasonic) H2O: 100 mg/mL (312.25 mM; Need ultrasonic)

Storage Instruction:

2-8°C, sealed storage, away from moisture

Alternative Names:

Cyclohexanecarboxamide, 4-[(1R)-1-aminoethyl]-N-4-pyridinyl-, hydrochloride (1:2), trans

Observed Molecular Weight:

320.26

References

[1]. Ishizaki T, et al. Pharmacological properties of Y-27632, a specific inhibitor of rho-associated kinases. Mol Pharmacol. 2000 May;57(5):976-83. [2]. Xue ZW, et al. Rho-associated coiled kinase inhibitor Y-27632 promotes neuronal-like differentiation of adult human adipose tissue-derived stem cells. Chin Med J (Engl). 2012 Sep;125(18):3332-5. [3]. Inan S, et al. Antiepileptic effects of two Rho-kinase inhibitors, Y-27632 and fasudil, in mice. Br J Pharmacol. 2008 Sep;155(1):44-51. [4]. Tada S, et al. A selective ROCK inhibitor, Y27632, prevents dimethylnitrosamine-induced hepatic fibrosis in rats. J Hepatol. 2001 Apr;34(4):529-36

Product Description

Y-27632 dihydrochloride is an ATP-competitive inhibitor of **ROCK-I** and **ROCK-II**, with **K**_i of 220 nM and 300 nM for **ROCK-I** and **ROCK-II**, respectively.

IC50 & Target: Ki: 220/300 nM (ROCK-I/II)^[1]

In Vitro: Y-27632 inhibits the ROCK family of kinases 100 times more potently than other kinases including protein kinase C, cAMP-dependent kinase and myosin light chain kinase. Y-27632 prolongs the lag time and delays the appearance of BrdU-labeled cells in a concentration-dependent manner, delays of about 1 and 4 h are noticed in the Swiss 3T3 cells treated with 10 and 100 μ M Y-27632, respectively^[1]. Y-27632 promotes neuronal differentiation of adipose tissue-derived stem cells (ADSCs). Compared to 1.0 and 2.5 μ M Y-27632 induced groups, percentages of neuroal-like cells achieved a peak in the 5.0 μ M Y-27632 induced group^[2].

In Vivo: Y-27632 (5 and 10 mg/kg) significantly prolongs the onset time of myoclonic jerks when compare with saline group. Y-27632 (5 and 10 mg/kg) significantly prolongs the onset time of clonic convulsions when compare with saline group^[3]. Treatment with Dimethylnitrosamine (DMN) causes a significant decrease in rat body and liver weight (DMN-S group) compared with control animals (S-S group). Oral Y27632 (30 mg/kg) essentially prevents this DMN-induced rat body and liver weight loss (DMN-Y group)^[4].





All products are for RESEARCH USE ONLY. Not for diagnostic & therapeutic purposes!