



A 438079 (hydrochloride)

Catalog No: tcsc1292



Available Sizes

Size: 10mg

Size: 50mg



Specifications

CAS No:

899431-18-6

Formula:

 $C_{13}H_{10}CI_3N_5$

Pathway:

Membrane Transporter/Ion Channel

Target:

P2X Receptor

Purity / Grade:

>98%

Solubility:

 $H2O : \ge 350 \text{ mg/mL} (1021.57 \text{ mM})$

Observed Molecular Weight:

342.61

Product Description

A 438079 (hydrochloride) is a potent, and selective $P2X_7$ receptor antagonist with pIC₅₀ of 6.9.

IC50 & Target: pIC50: 6.9

In Vitro: In 1321N1 cells stably expressing rat P2X₇ receptors, A 438079 blocks BzATP-(10 μ M) evoked changes in intracellular calcium concentrations with an IC₅₀ of 321 nM. A 438079 is also selective for the P2X₇ receptor, at concentrations up to 100 μ M^[1].





In Vivo: A 438079 (80 μmol/kg, i.v.) reduces noxious and innocuous evoked activity of different classes of spinal neurons in neuropathic rats. A 438079 (100 and 300 μmol/kg, i.p.) significantly raises withdrawal thresh-olds in both the SNL and CCI models^[1]. Intraperitoneal injection of A 438079 (5 and 15 mg/kg) 60 min after triggering seizures reduces seizure severity and neuronal death within the hippocampus. A 438079 has superior neuroprotective effects compared with an equally dose of phenobarbital (25 mg/kg) ^[2]. A 438079 partially but significantly prevents the 6-OHDA-induced depletion of striatal DA stores^[3]. Pretreatment with A 438079 reduces nociceptive behaviour scores in the HC model^[4].

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