



SX-3228

Catalog No: tcsc0018443

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Available Sizes

Size: 1mg

Size: 5mg

Size: 10mg



Specifications

CAS No:

156364-04-4

Formula:

 $C_{18}H_{18}N_4O_3$

Pathway:

Neuronal Signaling; Membrane Transporter/Ion Channel

Target:

GABA Receptor; GABA Receptor

Purity / Grade:

>98%

Solubility:

10 mM in DMSO

Observed Molecular Weight:

338.36

Product Description

SX-3228 is a selective **benzodiazepine1** (**BZ1**) receptor agonist with an IC_{50} of 17 nM.

IC50 & Target: IC50: 17 nM (BZ1 receptor)^[1]

In Vitro:





SX-3228 is a selective ligand for the BZ1 receptor. Among the BZ-receptor subtypes, SX-3228 preferentially binds to the BZ1 receptor (IC $_{50}$ =17 nM). It has very weak affinity for the BZ2 receptor (spinal cord: IC $_{50}$ =127 nM), and virtually no affinity for the peripheral type BZ receptor (kidney: IC $_{50}$ >10000 nM). A compound with similar selectivity, SX-3228 has been shown to bind to BZ receptors, but not to dopamine (D $_1$, D $_2$), serotonin (5-HT $_1$, 5-HT $_2$ and 5-HT $_3$ subtypes), noradrenaline (α_1 , α_2 , β), GABA or acetylcholine (muscarinic) subtypes^[1].

In Vivo: Administration of 0.5-2.5 mg/kg SX-3228 to rats during the light phase induces a significant reduction of rapid-eye-movement sleep (REMS) (P[1].

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