Data Sheet (Cat.No.T6534)



HCI

Histamine dihydrochloride

Chemical Properties

CAS No.: 56-92-8

Formula: C5H9N3·2HCl

Molecular Weight: 184.07

Appearance: no data available

Storage: Pure form: -20°C for 3 years | In solvent: -80°C for 1

year year

NH₂

Biological Description

Description

	an organic nitrogen compound which is a powerful stimulant of gastric secretion, a constrictor of bronchial smooth muscle, a vasodilator, and also a centrally acting neurotransmitter.
Targets(IC50)	Endogenous Metabolite, Histamine Receptor
In vitro	Histamine suppresses the generation of ROS through the Histaminetype-2 receptor (H2 receptor).[1] Histamine inhibits the generation and release of reactive oxygen species (ROS) by monocytes/macrophages (MO) during respiratory burst. Histamine and interleukin-2 (IL-2) act synergistically to activate NK cell cytotoxicity (NKCC). Histamine combined with IL-2 might improve response rates and disease-free survival by protecting the cells of the immune system from oxidative stress and inducing natural endogenous immune cytotoxicity. [2]
In vivo	Histamine treatment (0.5 mg/kg or 5.0 mg/kg, twice daily) protects against liver injury as evident by normal serum transaminase levels and significantly reduced liver pathology scores in a rat model with early alcohol-induced liver injury. The protective effect of histamine is blocked by Ranitidine (10 mg/kg), an H2 receptor antagonist, indicating that the histamine effect is predominantly mediated through the H2 receptor. [1] Histamine (30 pg/rat, icv) increases both 3,4-dihydroxyphenylalanine accumulation and 3,4-dihydroxyphenylalanine acid concentrations in the nucleus accumbens in male rats, and this effect is not affect by H2 antagonist zolantidine, indicating that histamine stimulates mesolimbic DA neurons through an action at the H1 receptor. [3] Histamine (0.5 mg/kg s.c.) reduces the liver tumour weight by 46% and subcutaneous tumour weight by 41% versus rats receiving subcutaneous saline injections. The anti-tumour effect observed by subcutaneous histamine injections is inhibited by Ranitidine (50 mg/kg s.c.) in rats sarcoma. [4] Histamine (1000 mg/kg s.c.) displays acute tissue damage after 24 hours and indications of pathological inflammation at the injection sites at 5 days and 28 days in Sprague-Dawley rats. Histamine (1000 mg/kg s.c.) results in Cmax of 167 mM, tmax of 0.5 hour, t1/2 of 0.95 and AUC of 186 mmol-h/L in male Sprague-Dawley rats. [5]

Histamine dihydrochloride (Ceplene) is dihydrochloride form of Histamine. Histamine is

Solubility Information

A DRUG SCREENING EXPERT

Solubility	H2O: 18.4 mg/mL (99.96 mM), Sonication is recommended.	
	DMSO: 70 mg/mL (380.29 mM), Sonication is recommended.	
	(< 1 mg/ml refers to the product slightly soluble or insoluble)	

Preparing Stock Solutions

Tel:781-999-4286

	1mg	5mg	10mg	
1 mM	5.4327 mL	27.1636 mL	54.3272 mL	
5 mM	1.0865 mL	5.4327 mL	10.8654 mL	
10 mM	0.5433 mL	2.7164 mL	5.4327 mL	
50 mM	0.1087 mL	0.5433 mL	1.0865 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.

Reference

Hornyak SC, et al. Inflammation, 2003, 27(5), 317-327.

Agarwala SS, et al. Expert Opin Biol Ther, 2001, 1(5), 869-879.

Fleckenstein AE, et al. Naunyn Schmiedebergs Arch Pharmacol, 1993, 347(1), 50-54.

Rizell M, et al. Anticancer Res, 2002, 22(4), 1943-1948.

Karavodin L, et al. Drug Chem Toxicol, 2003, 26(1), 35-49.

 $\textbf{Inhibitor} \cdot \textbf{Natural Compounds} \cdot \textbf{Compound Libraries} \cdot \textbf{Recombinant Proteins}$

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