Agomelatine hydrochloride

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Cat. No.:	HY-17038A	Q
CAS No.:	1176316-99-6	
Molecular Formula:	C ₁₅ H ₁₈ CINO ₂	
Molecular Weight:	279.76	
Target:	5-HT Receptor; Melatonin Receptor; Endogenous Metabolite	
Pathway:	GPCR/G Protein; Neuronal Signaling; Metabolic Enzyme/Protease	
Storage:	4°C, sealed storage, away from moisture * In solvent : -80°C, 6 months; -20°C, 1 month (sealed storage, away from moisture)	HCI

SOLVENT & SOLUBILITY

In Vitro	DMSO : ≥ 100 mg/mL (357.45 mM) H ₂ O : < 0.1 mg/mL (insoluble) * "≥" means soluble, but saturation unknown.				
	Preparing Stock Solutions	Solvent Mass Concentration	1 mg	5 mg	10 mg
		1 mM	3.5745 mL	17.8725 mL	35.7449 mL
		5 mM	0.7149 mL	3.5745 mL	7.1490 mL
		10 mM	0.3574 mL	1.7872 mL	3.5745 mL
	Please refer to the solubility information to select the appropriate solvent.				
In Vivo	1. Add each solvent one by one: 10% DMSO >> 40% PEG300 >> 5% Tween-80 >> 45% saline Solubility: ≥ 2.5 mg/mL (8.94 mM); Clear solution				
	2. Add each solvent one by one: 10% DMSO >> 90% (20% SBE-β-CD in saline) Solubility: ≥ 2.5 mg/mL (8.94 mM); Clear solution				
	3. Add each solvent one by one: 10% DMSO >> 90% corn oil Solubility: ≥ 2.5 mg/mL (8.94 mM); Clear solution				

Description	Agomelatine hydrochloride (S	-20098 hydrochloride) is a specif	ic agonist of MT1 and MT2 recept	ors with K _i s of 0.1, 0.06, 0.12,	
	and 0.27 nM for CHO-hMT1, HI	EK-hMT1, CHO-hMT2, and HEK-hl	MT2, respectively ^[1] . Agomelatine	hydrochloride is a selective	
	5-HT2C receptor antagonist w	rith pK _i s of 6.4 and 6.2 at native (p	porcine) and cloned, human 5-HT	2C receptors, respectively ^[2] .	
IC ₅₀ & Target	5-HT _{2C} Receptor	5-HT _{2C} Receptor	hMT1	hMT1	
	6.4 (pKi, native porcine)	6.2 (pKi, human)	0.1 (Ki, CHO Cells)	0.06 (Ki, HEK Cells)	

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Proteins

	hMT2 0.12 (Ki, CHO Cells)	hMT2 0.27 (Ki, HEK Cells)
In Vitro	Agomelatine (S 20098) acts as a full agonist of MT1 and MT2 receptors with EC ₅₀ s of 1.6±0.4, 0.10±0.04 nM for CHO hMT1 CHO-hMT2 (hMT1 and hMT2 receptors expressed in CHO or HEK cell membranes) ^[1] . Agomelatine (S20098) also interacts with h5-HT2B receptors (6.6), whereas it shows low affinity at native (rat)/cloned, human 5-HT2A (<5.0/5.3) and 5-HT1A (<5.0/5.2) receptors, and negligible (<5.0) affinity for other 5-HT receptors ^[2] . MCE has not independently confirmed the accuracy of these methods. They are for reference only.	
In Vivo	vo Agomelatine (25, 50, or 75 mg/kg; i.p.) has antioxidant activity in Strychnine (75 mg/kg, i.p.) induced seizure models in mice. Agomelatine dose not have any antioxidant effects on para produced by Pentylenetetrazole (PTZ) or Picrotoxin (PTX) induced seizure models when cor MCE has not independently confirmed the accuracy of these methods. They are for reference	
	Animal Model:	Female Swiss mice (20-30 g) were administered PTZ (85 mg/kg, i.p.), PTX (7 mg/kg, i.p.), strychnine (75 mg/kg, i.p.), Pilocarpine (400 mg/kg, i.p.), respectively ^[3]
	Dosage:	25, 50, or 75 mg/kg
	Administration:	Administered intraperitoneally (i.p.)
	Result:	All dosages showed a significant decrease in thiobarbituric acid reactive substances (TBARS) levels and nitrite content in all brain areas when compared to controls in the Pilocarpine induced seizure model. All dosages decreased TBARS levels in all brain areas, and at low doses (25 or 50 mg/kg) decreased nitrite contents, but only at 25 or 50 mg/kg showed a significant increase in catalase activity in three brain areas when compared to controls in the Strychnine- induced seizure model. Did not have any antioxidant effects on parameters of oxidative stress produced by PTX- or PTZ-induced seizure models when compared to controls.

CUSTOMER VALIDATION

- Protein Cell. 2019 Mar;10(3):178-195.
- Cell Commun Signal. 2023 May 25;21(1):123.
- Pest Manag Sci. 2021 Jul;77(7):3561-3570.

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REFERENCES

[1]. Audinot V, et al. New selective ligands of human cloned melatonin MT1 and MT2 receptors. Naunyn Schmiedebergs Arch Pharmacol. 2003 Jun;367(6):553-61.

[2]. Millan MJ, et al. The novel melatonin agonist agomelatine (S20098) is an antagonist at 5-hydroxytryptamine2C receptors, blockade of which enhances the activity of frontocortical dopaminergic and adrenergic pathways. J Pharmacol Exp Ther. 2003 Sep;306(3):954-64.

[3]. Aguiar CC, et al. Effects of agomelatine on oxidative stress in the brain of mice after chemically induced seizures. Cell Mol Neurobiol. 2013 Aug;33(6):825-35.

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

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