

Product Data Sheet

Dexmedetomidine

Cat. No.: HY-12719

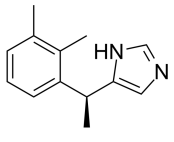
CAS No.: 113775-47-6Molecular Formula: $C_{13}H_{16}N_2$ Molecular Weight: 200.28

Target: Adrenergic Receptor

Pathway: GPCR/G Protein; Neuronal Signaling

Storage: 4°C, protect from light

* In solvent : -80°C, 6 months; -20°C, 1 month (protect from light)



SOLVENT & SOLUBILITY

In Vitro

DMSO: 125 mg/mL (624.13 mM; Need ultrasonic)

Preparing Stock Solutions	Solvent Mass Concentration	1 mg	5 mg	10 mg
	1 mM	4.9930 mL	24.9650 mL	49.9301 mL
	5 mM	0.9986 mL	4.9930 mL	9.9860 mL
	10 mM	0.4993 mL	2.4965 mL	4.9930 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: \geq 2.08 mg/mL (10.39 mM); Clear solution
- 2. Add each solvent one by one: 10% DMSO >> 90% (20% SBE-β-CD in saline) Solubility: ≥ 2.08 mg/mL (10.39 mM); Clear solution
- 3. Add each solvent one by one: 10% DMSO >> 90% corn oil Solubility: ≥ 2.08 mg/mL (10.39 mM); Clear solution

BIOLOGICAL ACTIVITY

Description	Dexmedetomidine ((+)-Medetomidine) is a potent, selective and orally active agonist of $\alpha 2$ -adrenoceptor, with a K_i of 1.08 nM. Dexmedetomidine shows 1620-fold selectivity against $\alpha 1$ -adrenoceptor. Dexmedetomidine exhibits anxiolysis, sedation, and modest analgesia effects [1][2][3].	
IC ₅₀ & Target	α2-adrenergic receptor 1.08 nM (Ki)	
In Vitro	Medetomidine has high selectivity for $\alpha 2$ adrenoceptors (K_i =1.08 nM) over $\alpha 1$ adrenoceptors (K_i =1750 nM) in rat brain membranes ^[1] .	

	Medetomidine (0.1-100 nM) inhibits the twitch response in field-stimulated mouse vas deferens, with a pD_2 of $9.0^{[1]}$. MCE has not independently confirmed the accuracy of these methods. They are for reference only.		
In Vivo	Medetomidine (10-100 μ g/kg; i.v. at 5-min intervals) produces a dose-dependent pupillary dilatation in pentobarbitone-anaesthetized rats ^[1] . MCE has not independently confirmed the accuracy of these methods. They are for reference only.		
	Animal Model:	Female Sprague-Dawley rats (270-350 g) ^[1]	
	Dosage:	1, 5, 10, 50, 100 mg/kg	
	Administration:	I.v. at 5-min intervals	
	Result:	Produced the pupil dilatation of 2.5 mm (approximately half of the maximum effect) at the cumulative dose of 4 $\mu g/kg$.	

CUSTOMER VALIDATION

- Nat Commun. 2023 Jul 7;14(1):4011.
- Biomed Pharmacother. 2023 Nov 23:169:115915.
- Cardiovasc Drugs Ther. 2023 Jul 1.
- Eur J Neurosci. 2021 Nov 4.
- J Neuropathol Exp Neurol. 2022 Jul 11;nlac055.

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REFERENCES

- [1]. Virtanen R, et, al. Characterization of the selectivity, specificity and potency of medetomidine as an alpha 2-adrenoceptor agonist. Eur J Pharmacol. 1988 May 20;150(1-2):9-14.
- $[2]. \ Gertler \ R, et, al. \ Dex medetomidine: a novel sedative-analgesic agent. \ Proc \ (Bayl \ Univ \ Med \ Cent). \ 2001 \ Jan; 14(1): 13-21.$
- [3]. Sajid B, et, al. A comparison of oral dexmedetomidine and oral midazolam as premedicants in children. J Anaesthesiol Clin Pharmacol. Jan-Mar 2019;35(1):36-40.

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

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