# Deschloroclozapine

Cat. No.:	HY-42110			
CAS No.:	1977-07-7			
Molecular Formula:	$C_{18}H_{20}N_4$			
Molecular Weight:	292.38			
Target:	mAChR			
Pathway:	GPCR/G Protein; Neuronal Signaling			
Storage:	Powder	-20°C	3 years	
	In solvent	-80°C	6 months	
		-20°C	1 month	

## SOLVENT & SOLUBILITY

In Vitro	DMSO : 100 mg/mL (342.02 mM; Need ultrasonic)						
	Preparing Stock Solutions	Mass Solvent Concentration	1 mg	5 mg	10 mg		
		1 mM	3.4202 mL	17.1010 mL	34.2021 mL		
		5 mM	0.6840 mL	3.4202 mL	6.8404 mL		
		10 mM	0.3420 mL	1.7101 mL	3.4202 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 (8.55 mM); Clear solution						
	2. Add each solvent one by one: 10% DMSO >> 90% corn oil Solubility: ≥ 2.5 mg/mL (8.55 mM); Clear solution						
	3. Add each solvent one by one: 5% DMSO >> 40% PEG300 >> 5% Tween-80 >> 50% saline Solubility: ≥ 2.5 mg/mL (8.55 mM); Clear solution						
	4. Add each solvent one by one: 5% DMSO >> 95% (20% SBE-β-CD in saline) Solubility: ≥ 2.5 mg/mL (8.55 mM); Clear solution						
	5. Add each solvent one by one: 10% DMSO >> 90% (20% SBE-β-CD in saline) Solubility: ≥ 2.08 mg/mL (7.11 mM); Clear solution						

## **BIOLOGICAL ACTIVITY**

Description

Deschloroclozapine, a metabolite of Clozapine, is a highly potent muscarinic DREADDs agonist. Deschloroclozapine binds to DREADD receptor subtypes hM3Dq and hM4Di with K<sub>i</sub> of 6.3 and 4.2 nM, respectively. [<sup>11</sup>C]-Deschloroclozapine is developed as a promising PET tracer for DREADD imaging<sup>[1][2][3]</sup>.



# Product Data Sheet

IC <sub>50</sub> & Target	mAChR3	mAChR4	
In Vitro	Designer Receptors Exclusively Activated by Designer Drugs (DREADD) are a chemogenetic approach for remote manipulation of neuronal activity in freely-moving animals. DREADDs comprise mutated G protein-coupled receptors (GPCRs) that do not respond to their endogenous neurotransmitter, but do respond to an otherwise "inert" exogenous substance <sup>[2]</sup> . MCE has not independently confirmed the accuracy of these methods. They are for reference only.		
In Vivo	Deschloroclozapine (0.3 mg/kg; intramuscularly) impairs working memory function in male rhesus macaques (aged between 5 and 6 years and weighing 5.5-7.9 kg) <sup>[3]</sup> . Deschloroclozapine (0.1 mg/kg; i.m) is effective at activating DREADD receptors in vivo and reversibly inducing behavioral effects in monkeys <sup>[3]</sup> . MCE has not independently confirmed the accuracy of these methods. They are for reference only.		

### **CUSTOMER VALIDATION**

- Cell. 2023 Nov 22;186(24):5394-5410.e18.
- Nat Neurosci. 2020 Sep;23(9):1157-1167.
- Nat Commun. 2023 Oct 24;14(1):6758.
- Nat Commun. 2023 Jul 24;14(1):4456.
- Nat Commun. 2023 Feb 28;14(1):971.

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#### REFERENCES

[1]. Maggs JL, et al. The metabolic formation of reactive intermediates from clozapine, a drug associated with agranulocytosis in man. J Pharmacol Exp Ther. 1995;275(3):1463-1475.

[2]. Hu F, et al. <sup>18</sup>F-labeled radiotracers for in vivo imaging of DREADD with positron emission tomography. Eur J Med Chem. 2021;213:113047.

[3]. Upright NA, et al. Effect of chemogenetic actuator drugs on prefrontal cortex-dependent working memory in nonhuman primates. Neuropsychopharmacology. 2020;45(11):1793-1798.

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

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