



SCH 563705

Catalog No: tcsc0833

Available Sizes		
Size: 2mg		
Size: 5mg		
Size: 10mg		
Specifications		
CAS No: 473728-58-4		
Formula: C ₂₃ H ₂₇ N ₃ O ₅		
Pathway: GPCR/G Protein;Immunology/Inflammation		
Target: CXCR;CXCR		
Purity / Grade: >98%		
Solubility: DMSO : ≥ 30 mg/mL (70.51 mM)		
Observed Molecular Weight:		

Product Description

425.48

SCH 563705 is a potent and orally available **CXCR2** and **CXCR1** antagonist, with IC_{50} s of 1.3 nM, 7.3 nM and K_i s of 1 and 3 nM, respectively.



IC50 & Target: IC50: 1.3 nM (CXCR2), 7.3 nM (CXCR1)^[1], 5.2 nM (Mouse CXCR2)^[2]

Ki: 1 nM (CXCR2), 3 nM (CXCR1)^[1]

In Vitro: SCH 563705 (Compound 16) is a potent and orally available CXCR2 and CXCR1 antagonist, with IC $_{50}$ s of 1.3 nM, 7.3 nM and K $_{i}$ s of 1 and 3 nM, respectively. SCH 563705 shows potent inhibition against both Gro-a and IL-8 induced human neutrophil migration (chemotaxis IC $_{50}$ = 0.5 nM, against 30 nM of Gro-a; chemotaxis IC $_{50}$ = 37 nM, against 3 nM of IL-8)^[1]. SCH 563705 potently inhibits mouse CXCR2 (IC $_{50}$ = 5.2 nM)^[2].

In Vivo: SCH 563705 has good oral pharmacokinetic profiles in rats, mice, monkeys and dogs^[1]. SCH 563705 (50 mg/kg p.o) reduces blood Ly6G⁺ Ly6C⁺ neutrophil frequency and unchanged levels of Ly6GLy6Chi monocytes. SCH563705 (3-30 mg/kg p.o) treatment causes a dosedependent elevation in plasma levels of CXCL1^[2].

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