



GSK2239633A

Catalog No: tcsc0018178

Available Sizes	
Size: 1mg	
Size: 5mg	
Size: 10mg	
Specifications	
CAS No: 1240516-71-5	
Formula: $C_{24}H_{25}CIN_4O_5S_2$	
Pathway: Immunology/Inflammation;GPCR/G Protein	
Target: CCR;CCR	
Purity / Grade: >98%	
Solubility: 10 mM in DMSO	
Observed Molecular Weight: 549.06	

Product Description

GSK2239633A is a CC-chemokine receptor 4 (**CCR4**) antagonist, which inhibits the binding of [^{125}I]-TARC to human CCR4 with a **pIC** 50 of $^{7.96 \pm 0.11}$.





IC50 & Target: CCR4^[1]

In Vitro: The GSK2239633A is an allosteric antagonist of human CCR4. GSK2239633A inhibits the binding of [125 I]-TARC to human CCR4 with a pIC $_{50}$ of 7.96±0.11 and also inhibits thymus- and activation-regulated chemokine-induced (TARC)-induced increases in the F-actin content of isolated human CD4⁺ CCR4⁺ T-cells with a pA $_2$ of 7.11±0.29^[1]. The effect of GSK2239633A (Compound 3) on CCL17-induced increases in the F-actin content of human CD4⁺ CCR4⁺ T cells is measured. The pEC $_{50}$ value is 8.79±0.22^[2].

In Vivo: Following intravenous dosing, plasma GSK2239633A displays rapid, bi-phasic distribution and slow terminal elimination ($t_{1/2}$: 13.5 hours), suggesting that GSK2239633A is a low to moderate clearance drug. Following oral dosing, blood levels of GSK2239633A reach C_{max} rapidly (median t_{max} : 1.0-1.5 hours). Estimated GSK2239633A bioavailability is low with a maximum value determined of only $16\%^{[1]}$. GSK2239633A (Compound 9) demonstrates good pharmacokinetic data in preclinical animal studies (bioavailability in rats and beagle dogs 85% and 97% respectively)^[3].

All products are for RESEARCH USE ONLY. Not for diagnostic & therapeutic purposes!