



PI3kδ inhibitor 1

Catalog No: tcsc0675

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Z	4		

Available Sizes

Size: 5mg



Specifications

CAS No:

1332075-63-4

Formula:

 $C_{28}H_{33}FN_{6}O_{2}$

Pathway:

PI3K/Akt/mTOR

Target:

PI3K

Purity / Grade:

>98%

Solubility:

10 mM in DMSO

Observed Molecular Weight:

504.6

Product Description

PI3k δ inhibitor 1 is a potent and selective **PI3K\delta** inhibitor with an **IC**₅₀ of 3.8 nM.

IC50 & Target: IC50: 3.8 nM (PI3Kδ)^[1]

In Vitro: PI3k δ inhibitor 1 (Compound 3) is a potent inhibitor of PI3K δ that is 200-400 fold selective for all three remaining Class I PI3K isoforms and extremely selective relative to 239 kinases tested in SelectScreen service (0/239 kinases showing >50% inhibition when tested at 1 μ M; mTOR, DNA-PK, VPS34, PI4K α and PI4K β are inhibited at 10% or less when tested at 1 μ M; PIKC2A and PIKC2B are inhibited at 11% and 42%, respectively, at this same concentration and show less than 10% inhibition when tested at 0.1 μ M; the





PIKK family kinases ATM and ATR are not assessed)^[1].

In Vivo: The pharmacokinetic properties of PI3k δ inhibitor 1 (Compound 3) are evaluated in mice and rats when dosed IV and orally. Good plasma exposures and reasonable half-lives are observed upon oral dosing, a reflection of high oral bioavailability (80% and 90% at a low dose for mouse and rat, respectively), moderate volume of distribution, and moderate clearance. PI3k δ inhibitor 1 has moderate terminal elimination half-life ($t_{1/2}$ =2.6 h, 2.9 h, 5 h, 2.6, 3.8 and 4.8 h for mouse (5 mg/kg, po), mouse (20 mg/kg, po), mouse (40 mg/kg, po), rat (5 mg/kg, po), rat (10 mg/kg, po), rat (30 mg/kg, po)). Plasma exposures and Cmax levels increase with dose in both mice and rats, important in that inflammatory disease models utilize these two species. Plasma protein binding for PI3k δ inhibitor 1 ranges from 80-88% in rodents and is consistent with values obtained in human plasma (86%)^[1].

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