



SAR131675

Catalog No: tcsc0970



Available Sizes

Size: 10mg

Size: 50mg



Specifications

CAS No:

1433953-83-3

Formula:

 $C_{18}H_{22}N_4O_4$

Pathway:

Protein Tyrosine Kinase/RTK

Target:

VEGFR

Purity / Grade:

>98%

Solubility:

DMSO : \geq 28 mg/mL (78.13 mM)

Observed Molecular Weight:

358.39

Product Description

SAR131675 is a potent and selective **VEGFR3** inhibitor with an IC_{50} of 23 nM.

IC50 & Target: IC50: 23 nM (VEGFR3)^[1]

In Vitro: AR131675 is highly selective for VEGFR-3 versus 107 receptors, enzymes, ion channels, and 65 kinases. However, it is moderately active on VEGFR-2 with a VEGFR-3/VEGFR-2 ratio of about 10. SAR131675 inhibits VEGFR-3 tyrosine kinase activity and





VEGFR-3 autophosphorylation in HEK cells with IC $_{50}$ values of 20 and 45 nM, respectively. SAR131675 dose dependently inhibits the proliferation of primary human lymphatic cells, induced by the VEGFR-3 ligands VEGFC and VEGFD, with an IC $_{50}$ of about 20 nM. SSAR131675 has no antiproliferative activity on a panel of 30 tumors and primary cells, further showing its high specificity and indicating that SAR131675 is not a cytotoxic or cytostatic agent^[1].

In Vivo: SAR131675 is very well tolerated in mice and shows a potent antitumoral effect in several orthotopic and syngenic models, including mammary 4T1 carcinoma and RIP1.Tag2 tumors. Interestingly, it significantly reduces lymph node invasion and lung metastasis, showing its antilymphangiogenic activity in vivo. SAR131675 significantly reduces TAM infiltration and aggregation in 4T1 tumors^[1].

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