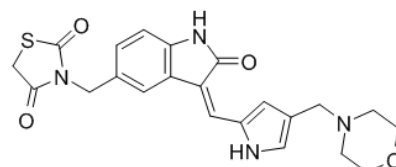


Product Name : S49076
Cat. No. : PC-43299
CAS No. : 1265965-22-7
Molecular Formula : C₂₂H₂₂N₄O₄S
Molecular Weight : 438.4995
Target : TAM Receptor (Tyro3-Axl-Mer)
Solubility : DMSO: ≥ 31 mg/mL



Biological Activity

S49076 is a potent, ATP-competitive tyrosine kinase inhibitor of **MET**, **AXL/MER**, and **FGFR1/2/3** with IC₅₀ of <20 nM, also potently inhibits the kinase activity of mutated isoforms of MET (D1246N, Y1248D, Y1248H) and FGFR1/2. S49076 only inhibits 6% of kinases on a panel of 442 human wild-type and mutated kiTAM/FGFTnases at 100 nM. S49076 inhibits the proliferation of MET- and FGFR2-dependent gastric cancer cells, blocks MET-driven migration of lung carcinoma cells, and inhibits colony formation of hepatocarcinoma cells expressing FGFR1/2 and AXL. S49076 causes tumor growth arrest in bevacizumab-resistant tumors in cancer xenograft models.

References

- Burbridge MF, et al. *Mol Cancer Ther.* 2013 Sep;12(9):1749-62.
Clémenson C, et al. *Mol Cancer Ther.* 2017 Oct;16(10):2107-2119.
Rodon J, et al. *Eur J Cancer.* 2017 Aug;81:142-150.

Caution: Product has not been fully validated for medical applications. Lab Use Only!

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