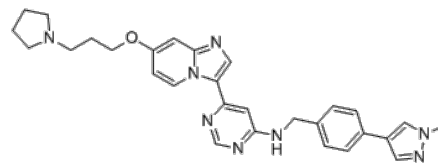


Product Name : M4205
Cat. No. : PC-20159
CAS No. : 2590556-80-0
Molecular Formula : C₂₉H₃₂N₈O
Molecular Weight : 508.63
Target : c-Kit
Solubility : 10 mM in DMSO



Biological Activity

M4205 (IDRX-42) is a potent, highly selective inhibitor of **KIT mutations** with cell IC₅₀ of 52 nM on cKIT autophosphorylation in the GIST430/654 cell line.

M4205 inhibits KIT autophosphorylation at Y703 in the imatinib sensitive GIST430 cell line with an IC₅₀ value of 4 nM.

M4205 inhibits the imatinib-resistant cell line GIST430/654 (exon 11 and exon 13 mutation) and the AML cell line Kasumi-1 (exon 17 mutation N822 K) with IC₅₀ values of 48 and 4 nM, respectively.

M4205 displays high selectivity in a biochemical panel of 398 kinases at 1 μM, only inhibits PDGFRA, PDGFRB, KIT (wt), FLT3, CSF1R, and lymphocyte-specific protein tyrosine kinase (LCK) with >80% inhibition.

M4205 also displays high cellular kinase selectivity at 1 μM using a panel of NanoBRET assays, only binds to KIT (wt) and FLT3 with 65% and 45% occupancy.

M4205 (35 mg/kg) showed strong in vivo tumor growth inhibition and led to regression in mice bearing GIST430/654 tumor, with dose- and exposure-dependent inhibition of KIT autophosphorylation.

References

Andreas Blum, et al. *J Med Chem.* 2023 Feb 23;66(4):2386-2395.

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

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