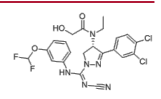


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Data Sheet

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Product Name	:	BAY-598
Cat. No.	:	PC-60420
CAS No.	:	906919-67-2
Molecular Formula	:	C ₂₂ H ₂₀ Cl ₂ F ₂ N ₆ O ₃
Molecular Weight	:	525.3
Target	:	Histone Methyltransferase (HMTase)
Solubility	:	10 mM in DMSO



Biological Activity

BAY-598 (BAY598) is a potent, selective, and cell-active, substrate-competitive inhibitor of **SMYD2** with IC50 of 27±7 nM in the biochemical SPA assay.

BAY-598 displays >100-fold selectivity over other HMTases (SMYD3, SUV420H1, SUV420H2, etc.) and other non-epigenetic targets.

BAY-598 inhibits the methylation of p53K370 in cells with IC50 <1 uM, has properties that are compatible with in vivo experiments.

References

Eggert E, et al. *J Med Chem.* 2016 May 26;59(10):4578-600.

Caution: Product has not been fully validated for medical applications. Lab Use Only! E-mail: tech@probechem.com