

产品名称 : JGB1741

同义词 : ILS-JGB-1741

产品货号 : M27505

CAS Number : 1256375-38-8

分子式 : C27H24N2O2S

分子量 : 440.56

化学全名 : ——

产品描述 : JGB1741 is a potent and selective inhibitor of SIRT1 with an IC50 of 15 μ M. JGB1741 modulates Bax/Bcl2 ratio, cytochrome c release and PARP cleavage and increases the acetylated p53 levels leading to p53-mediated apoptosis. JGB1741 can be used in studies about breast cancer. (In Vitro): JGB1741 (1-10000 nM) inhibits the cell proliferation of K562, HepG2 and MDA-MB 231 cell lines. JGB1741 (0.01-1 μ M) induces apoptosis of MDA-MB 231 and shows a cell cycle arrest at G1 phase with more cells entering into sub G0/G1 phase. JGB1741 (0.01-1 μ M) increases the global acetylation of H3K9, acetylated p53K382 levels and p53 expression.

通路 : Chromatin/Epigenetic

靶点 : Sirtuin

受体 : Aurora B

溶解度 : In Vitro: ?DMSO : 5 mg/mL (11.35 mM)

SMILES : C(NCC1=CC=CC=C1)(=O)C=2C3=C(SC2/N=C/C=4C5=C(C=CC4O)C=CC=C5)CCCC3

存储条件 : (-20°C)

稳定性 : ≥ 2 years

参考文献 :

1.1. Naga Rajiv Lakkaniga, et al. Discovery of SP-96, the first non-ATP-competitive Aurora Kinase B inhibitor, for reduced myelosuppression. Eur J Med Chem. 2020 Jul 12; 203:112589.

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