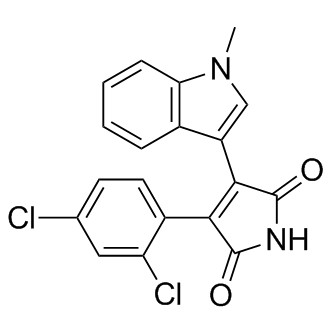
**SB216763 Package : 10mg**

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**Technical literature is available at:** [**www.mesgenbio.com**](http://www.mesgenbio.com)**. E-mail MesGen Technical Services if you have questions on use of this system: tech@mesgenbio.com**

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**Catalog Number :** MG29903

**Appearance :** Off-white to orange(solid)

**Molecular weight :** 371.22

**CAS :** 280744-09-4

**Purity ≥98% by HPLC**

**Molecular formula :** C19H12Cl2N2O2

**Description**

A cell-permeable (arylindole) maleimide compound that acts as a potent, selective, ATP-competitive inhibitor of GSK-3 activity (Ki = 9.1 nM for GSK-3α). At 10 µM concentration, does not significantly inhibit a panel of 24 other protein kinases tested. Reported to stimulate glycogen synthesis in Chang human liver cells (EC50 = 3.6 µM) and induce transcription of β-catenin-LEF/TCF (lymphoid enhancer factor/T cell factor) regulated reporter gene in HEK293 human embryonic kidney cell lines. Offers protection against polyglutamine-induced death in SK-N-SH and COS-7 cells and attenuates hypoxia-induced apoptosis in VSMC and COS-7 cells. Shown to reduce phosphoenolpyruvate carboxykinase (PEPCK) and glucose-6-phosphatase (G6Pase) expression in hepatoma cells. Promotes the survival of CLL lymphocytes via Wnt/β-catenin-mediated transcription activation. Exhibits delayed cardioprotection *in vivo* in a rat model via a KATP (ATP-sensitive potassium channels) and MPTP (mitochondrial permeability transition pore)-dependent mechanism at reperfusion.

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**Solvent & Solubility**

DMSO ≥25mg/mL Water <1.2mg/mL Ethanol <1.2mg/mL

**Storage condition**

-20°C

**For Research Use Only. Not For Use In Diagnostic Procedures.**