Product Data Sheet

GSK-1070916

Cat. No.: HY-70044

CAS No.: 942918-07-2

Molecular Formula: $C_{30}H_{33}N_7O$ Molecular Weight: 507.63

Target: Aurora Kinase; Apoptosis

Pathway: Cell Cycle/DNA Damage; Epigenetics; Apoptosis

Storage: Powder -20°C 3 years

4°C 2 years -80°C 6 months

In solvent -80°C 6 months

-20°C 1 month

SOLVENT & SOLUBILITY

In Vitro

DMSO: 10 mg/mL (19.70 mM; Need ultrasonic)

Preparing Stock Solutions	Solvent Mass Concentration	1 mg	5 mg	10 mg
	1 mM	1.9699 mL	9.8497 mL	19.6994 mL
	5 mM	0.3940 mL	1.9699 mL	3.9399 mL
	10 mM	0.1970 mL	0.9850 mL	1.9699 mL

Please refer to the solubility information to select the appropriate solvent.

In Vivo

- 1. Add each solvent one by one: 10% DMSO >> 40% PEG300 >> 5% Tween-80 >> 45% saline Solubility: ≥ 1 mg/mL (1.97 mM); Clear solution
- 2. Add each solvent one by one: 10% DMSO >> 90% (20% SBE- β -CD in saline) Solubility: \geq 1 mg/mL (1.97 mM); Clear solution
- 3. Add each solvent one by one: 10% DMSO >> 90% corn oil Solubility: ≥ 1 mg/mL (1.97 mM); Clear solution

BIOLOGICAL ACTIVITY

Description GSK-1070916 is a potent and selective ATP-competitive inhibitor of aurora B and aurora C with K_is of 0.38 and 1.5 nM, respectively, and is >250- fold selective over Aurora A.

respectively, and is 250 Tota selective over Autora A.

IC₅₀ & Target Aurora B Aurora C

0.38 nM (Ki) 1.5 nM (Ki)

In Vitro GSK-1070916 potently inhibits Aurora B/INCENP and Aurora C/INCENP kinases with K_is of 0.38±0.29 and 1.45±0.35 nM,

respectively, but is less potent against Aurora A/ TPX2 with a $\rm K_i$ of 492 \pm 61 nM. GSK-1070916 also inhibits FLT1, TIE2, SIK, FLT4, and FGFR1 with IC $_{50}$ values of 42, 59, 70, 74, and 78 nM, respectively. Treatment of A549 human lung cancer cells with GSK-1070916 results in a potent antiproliferative effect (EC $_{50}$ =7 nM)^[1]. GSK-1070916 inhibits a panel of tumor cell lines and is shown o inhibits the phosphorylation of HH3- S10 in all cell lines with average EC $_{50}$ values ranging from 8 to 118 nM^[2]. MCE has not independently confirmed the accuracy of these methods. They are for reference only.

In Vivo

In nude mice implanted with human colon tumor (HCT116) xenografts, a single dose of GSK-1070916 administered i.p. inhibits HH3-S10 phosphorylation in a dose-dependent manner. Repeated i.p. administration of GSK-1070916 produces complete or partial antitumor activity in 4 of 8 tumor types [lung, A549; colon, HCT116; acute myelogenous leukemia (AML), HL60; and chronic myelogenous leukemia, K562], stable disease in 3 of 8 (colon, Colo205; lung, H460; and breast, MCF-7), and tumor growth delay in 1 of 8 tumor types (colon, SW620). Daily administration of GSK-1070916 is generally well-tolerated [2].

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PROTOCOL

Cell Assay [2]

A panel of tumor cell lines are plated in 96-well plates in the recommended growth media and incubated at 37° C in 5% CO₂ overnight. The following day, the cells are treated with serial dilutions of GSK-1070916. At this time, one set of cells is treated with CellTiter-Glo for a time equal to 0 (T=0) measurement. Following a 6- to 7-d incubation with compound, cell proliferation is measured using the CellTiter-Glo reagent^[2].

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Animal Administration [2]

Mice: Tumors are initiated by injection of tumor cell suspensions (A549, SW620, HCT116, H460, MCF-7, HL60, K562) or tumor fragments (Colo205) s.c. into nude (A549, SW620, HCT116, H460, MCF-7, HL60, and Colo205) or severe combined immunodeficient (SCID; K562) mice. When the tumors reach a volume of 80 to 200 mm³, the mice are randomized into groups of 5 to 10 mice per group. GSK-1070916 is administered at 25, 50, or 100 mg/kg once daily for 5 consecutive days-on, 2d-off, schedule for two (Colo205 and HL60) or three (A549, SW620, HCT116, H460, MCF-7, K562) cycles. Tumors are measured twice weekly^[2].

 $\label{eq:mce} \mbox{MCE has not independently confirmed the accuracy of these methods. They are for reference only.}$

CUSTOMER VALIDATION

- J Biomol Screen. 2013 Oct;18(9):1062-71.
- Harvard Medical School LINCS LIBRARY

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REFERENCES

 $[1]. Adams \, ND, \, et \, al. \, Discovery \, of \, GSK-1070916, \, a \, potent \, and \, selective \, inhibitor \, of \, Aurora \, B/C \, kinase. \, J \, Med \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, GSK-1070916, \, a \, potent \, and \, selective \, inhibitor \, of \, Aurora \, B/C \, kinase. \, J \, Med \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, GSK-1070916, \, a \, potent \, and \, selective \, inhibitor \, of \, Aurora \, B/C \, kinase. \, J \, Med \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, GSK-1070916, \, a \, potent \, and \, selective \, inhibitor \, of \, Aurora \, B/C \, kinase. \, J \, Med \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, GSK-1070916, \, a \, potent \, and \, below \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, GSK-1070916, \, a \, potent \, and \, below \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, GSK-1070916, \, a \, potent \, and \, below \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \, 27; \\ 53(10):3973-4001. \, Discovery \, of \, Chem. \, 2010 \, May \,$

[2]. Hardwicke MA, et al. GSK-1070916, a potent Aurora B/C kinase inhibitor with broad antitumor activity in tissue culture cells and human tumor xenograft models. Mol Cancer Ther. 2009 Jul;8(7):1808-17.

 $\label{lem:caution:Product} \textbf{Caution: Product has not been fully validated for medical applications. For research use only.}$

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