## CCT 137690

Cat. No.:	HY-10804		
CAS No.:	1095382-05	-0	
Molecular Formula:	C <sub>26</sub> H <sub>31</sub> BrN <sub>8</sub> O		
Molecular Weight:	551.48		
Target:	Aurora Kinase; Apoptosis		
Pathway:	Cell Cycle/DNA Damage; Epigenetics; Apoptosis		
Storage:	Powder	-20°C	3 years
		4°C	2 years
	In solvent	-80°C	2 years
		-20°C	1 year

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## SOLVENT & SOLUBILITY

		Solvent Mass Concentration	1 mg	5 mg	10 mg		
	Preparing Stock Solutions	1 mM	1.8133 mL	9.0665 mL	18.1330 mL		
		5 mM	0.3627 mL	1.8133 mL	3.6266 mL		
	10 mM	0.1813 mL	0.9067 mL	1.8133 mL			
	Please refer to the so	Please refer to the solubility information to select the appropriate solvent.					
ı Vivo		one by one: 10% DMSO >> 40% PEG ng/mL (3.03 mM); Clear solution	G300 >> 5% Tween-80	) >> 45% saline			
		2. Add each solvent one by one: 10% DMSO >> 90% (20% SBE-β-CD in saline) Solubility: ≥ 1.67 mg/mL (3.03 mM); Clear solution					
	3. Add each solvent one by one: 10% DMSO >> 90% corn oil Solubility: ≥ 1.67 mg/mL (3.03 mM); Clear solution						

BIOLOGICAL ACTIVITY			
Description	CCT 137690 is a potent and c respectively.	orally available aurora kina	se inhibitor with $\rm IC_{50}s$ of 15, 25, and 19 nM for aurora A, B and C,
IC₅₀ & Target	Aurora A 15 nM (IC <sub>50</sub> )	Aurora B 25 nM (IC <sub>50</sub> )	Aurora C 19 nM (IC <sub>50</sub> )
In Vitro	CCT 137690 displays antipro	liferative activity in a range	e of human tumor cell lines, including SW620 colon carcinoma (GI <sub>50</sub>

	=0.30 μM) and A2780 ovarian cancer cell line (GI <sub>50</sub> =0.14 μM). CCT 137690 inhibits in vitro phosphorylation of histone H3. CCT 137690 is a moderate inhibitor of the hERG ion-channel (IC <sub>50</sub> =3.0 μM) <sup>[1]</sup> . CCT137690 efficiently inhibits histone H3 and TACC3 phosphorylation (Aurora B and Aurora A substrates, respectively) in HCT116 and HeLa cells. Continuous exposure of tumour cells to the inhibitor causes multipolar spindle formation, chromosome misalignment, polyploidy and apoptosis <sup>[2]</sup> . MCE has not independently confirmed the accuracy of these methods. They are for reference only.
In Vivo	CCT 137690 slows the growth of the SW620 xenografts with no observed toxicity <sup>[1]</sup> . CCT 137690 significantly inhibits tumour growth in a transgenic mouse model of neuroblastoma (TH-MYCN) that overexpresses MYCN protein and is predisposed to spontaneous neuroblastoma formation <sup>[2]</sup> . MCE has not independently confirmed the accuracy of these methods. They are for reference only.

PROTOCOL	
Cell Assay <sup>[2]</sup>	Cells are plated in 96-well plates at 3,000 cells per well and are treated with a range of 0 to 25 mol/L of CCT137690 for 72 h. Cell proliferation assays are performed by colorimetric 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) <sup>[2]</sup> . MCE has not independently confirmed the accuracy of these methods. They are for reference only.
Animal Administration <sup>[2]</sup>	Mice: Animals are randomized into two groups, group 1: treatment with 100 mg/kg CCT137690 n=4 or group 2: vehicle control n=4. Treatment is administered via oral gavage twice daily. Tumour volumes are measured at day 0, 3 (48 hours after treatment started), 7 and 10 using <sup>1</sup> H MRI <sup>[2]</sup> . MCE has not independently confirmed the accuracy of these methods. They are for reference only.

## **CUSTOMER VALIDATION**

• SSRN. 2022 Nov 21.

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## REFERENCES

[1]. Bavetsias V, et al. Imidazo[4,5-b] pyridine derivatives as inhibitors of Aurora kinases: lead optimization studies toward the identification of an orally bioavailable preclinical development candidate. J Med Chem. 2010 Jul 22;53(14):5213-28.

[2]. Faisal A, et al. The aurora kinase inhibitor CCT137690 downregulates MYCN and sensitizes MYCN-amplified neuroblastoma in vivo. Mol Cancer Ther. 2011 Nov;10(11):2115-23.

Caution: Product has not been fully validated for medical applications. For research use only.

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