Spebrutinib

Cat. No.:	HY-18012		
CAS No.:	1202757-89	-8	
Molecular Formula:	C ₂₂ H ₂₂ FN ₅ O	3	
Molecular Weight:	423.44		
Target:	Btk		
Pathway:	Protein Tyr	osine Kin	ase/RTK
Storage:	Powder	-20°C	3 years
		4°C	2 years
	In solvent	-80°C	6 months
		-20°C	1 month

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SOLVENT & SOLUBILITY

In Vitro	DMSO : ≥ 45 mg/mL (106.27 mM) * "≥" means soluble, but saturation unknown.					
		Solvent Mass Concentration	1 mg	5 mg	10 mg	
	Preparing Stock Solutions	1 mM	2.3616 mL	11.8080 mL	23.6161 mL	
		5 mM	0.4723 mL	2.3616 mL	4.7232 mL	
		10 mM	0.2362 mL	1.1808 mL	2.3616 mL	
	Please refer to the so	lubility information to select the app	propriate solvent.			
In Vivo	1. Add each solvent o Solubility: ≥ 2.5 m	one by one: 10% DMSO >> 40% PEC g/mL (5.90 mM); Clear solution	6300 >> 5% Tween-80) >> 45% saline		
	2. Add each solvent of Solubility: ≥ 2.5 m	one by one: 10% DMSO >> 90% cor g/mL (5.90 mM); Clear solution	n oil			

BIOLOGICAL ACTIVITY				
Description	Spebrutinib (AVL-292; CC-292) is a covalent, orally active, and highly selective with an IC ₅₀ of 0.5 nM.			
IC ₅₀ & Target	IC50: <0.5 nM (Btk) ^[1]			
In Vitro	Spebrutinib (CC-292) is a covalent, highly selective, orally active inhibitor of Btk with IC ₅₀ value of 0.5 nM. Spebrutinib also less potently inhibits Yes, c-Src, Brk, Lyn, and Fyn with IC ₅₀ s of 723 nM, 1.729 μM, 2.43 μM, 4.4 μM, and 7.15 μM, rspectively. Extensive analysis has revealed that the EC ₅₀ of Btk occupancy from a Spebrutinib dose-response in Ramos cells (EC ₅₀ =6 nM) correlated directly with the cellular EC ₅₀ of Btk kinase inhibition with Spebrutinib (EC ₅₀ =8 nM). Furthermore, the concentration at which Spebrutinib inhibits 90% of Btk activity in Ramos cells is 35 nM while the concentration of			

Product Data Sheet



Spebrutinib required for 90% occupancy of Btk is $39 \text{ nM}^{[1]}$.

MCE has not independently confirmed the accuracy of these methods. They are for reference only.

PROTOCOL	
Cell Assav ^[1]	Cells are incubated in serum-free RPMI media for 1-15 hours. Isolated human B cells are incubated with Spebrutinib at a
	final concentration of 0.001, 0.01, 0.1 and 1 μ M. Ramos cells are incubated with 0.1 nM-3 μ M Spebrutinib. Cells are then
	incubated in the presence of compound for 1 hour at 37°C. Following incubation, cells are centrifuged and resuspended
	100 μL of serum-free RPMI and BCR is stimulated with addition of 5 μg/mL α-human IgM. Samples are centrifuged, washe
	phosphate-buffered saline (PBS), and lysed in 100 μL of Cell Extraction Buffer plus 1:10 (v/v) PhosSTOP Phosphatase
	Inhibitor and 1:10 (v/v) Complete Protease Inhibitor. Antibodies used for immunoblot analysis include P-PLCy2, PLCy2 (3
	CST), Syk (2712; CST), P-Syk (2710; CST), Btk, P-Btk, and Tubulin. Membranes are scanned on a Li-Cor Odyssey scanner u
	infrared fluorescence detection ^[1] .
	MCE has not independently confirmed the accuracy of these methods. They are for reference only.

CUSTOMER VALIDATION

- Blood. 2016 Jun 23;127(25):3237-52.
- Br J Pharmacol. 2019 Dec;176(23):4491-4509.
- Stem Cell Reports. 2019 May 14;12(5):996-1006.
- Molecules. 2023, 28(1), 79.
- R Soc Open Sci. 2019 Jun 5;6(6):190434.

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REFERENCES

[1]. Evans EK, et al. Inhibition of Btk with CC-292 provides early pharmacodynamic assessment of activity in mice and humans. J Pharmacol Exp Ther. 2013 Aug;346(2):219-28.

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

 Tel: 609-228-6898
 Fax: 609-228-5909
 E-mail: tech@MedChemExpress.com

 Address: 1 Deer Park Dr, Suite Q, Monmouth Junction, NJ 08852, USA