## BGT226 maleate

MedChemExpress

®

Cat. No.:	HY-13334	
CAS No.:	1245537-68-1	
Molecular Formula:	$C_{32}H_{29}F_{3}N_{6}O_{6}$	
Molecular Weight:	650.6	
Target:	PI3K; mTOR; Autophagy; Apoptosis	N N-OH
Pathway:	PI3K/Akt/mTOR; Autophagy; Apoptosis	N
Storage:	4°C, sealed storage, away from moisture	
	* In solvent : -80°C, 6 months; -20°C, 1 month (sealed storage, away from moisture)	

### SOLVENT & SOLUBILITY

In Vitro	DMSO : 62.5 mg/mL (96.07 mM; ultrasonic and warming and heat to 60°C)				
	Preparing Stock Solutions	Solvent Mass Concentration	1 mg	5 mg	10 mg
		1 mM	1.5370 mL	7.6852 mL	15.3704 mL
		5 mM	0.3074 mL	1.5370 mL	3.0741 mL
		10 mM	0.1537 mL	0.7685 mL	1.5370 mL
	Please refer to the sol	ubility information to select the app	propriate solvent.		
In Vivo	1. Add each solvent o Solubility: ≥ 2.5 mg	one by one: 10% DMSO >> 40% PEC g/mL (3.84 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: 2.5 mg/mL (3.84 mM); Suspended solution; Need ultrasonic				
	3. Add each solvent o Solubility: ≥ 2.5 mg	one by one: 10% DMSO >> 90% cor g/mL (3.84 mM); Clear solution	n oil		

Description	BGT226 (NVP-BGT226 maleate inhibitor which displays poter	e) is a PI3K (with IC <sub>50</sub> s of 4 nM, 63 nt growth-inhibitory activity agai	nM and 38 nM for ΡΙ3Κα, ΡΙ3Κβ a nst human head and neck cance	and PI3Ky) /mTOR dual r cells <sup>[1][2]</sup> .
IC <sub>50</sub> & Target	ΡΙ3Κα 4 nM (IC <sub>50</sub> )	ΡΙ3Κγ 38 nM (IC <sub>50</sub> )	ΡΙ3Κβ 63 nM (IC <sub>50</sub> )	mTOR
	Autophagy			
In Vitro	BGT226 shows significant grou	wth inhibition or signal blockage	profiles compared with LY29400	2 and Rapamycin. BGT226

# Product Data Sheet

(10-10000 nM) inhibits FaDu and OECM1 cells growth with  $IC_{50}$ s of 23.1±7.4 and 12.5±5.1 nM, respectively <sup>[2]</sup>. The expression levels of p-mTOR Ser2481 are decreased in BGT226-treated cell lines (200 nM; 24 hours) and both p-AKT Ser473 and p-mTOR Ser2448 are also decreased in BGT226-treated cell lines<sup>[2]</sup>.

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

#### Cell Viability Assay<sup>[2]</sup>

FaDu cells; OECM1 cells	
10, 100, 1000, 10000 nM	
Inhibited FaDu and OECM1 cells growth with $\rm IC_{50}s$ of 23.1±7.4 and 12.5±5.1 nM, respectively.	
FaDu cells; OECM1 cells	

Cell Line:	Fadu cells; OECM1 cells
Concentration:	200 nM
Incubation Time:	24 hour
Result:	p-mTOR Ser2481 expression levels decreased, and both p-AKT Ser473 and p-mTOR Ser2448 expression levels also decreased.

#### In Vivo

BGT226 (2.5 and 5 mg/kg; oral administration for 21 days in male athymic mice) causes 34.7% and 76.1% reduction of the tumor growth on day 21 compared with control<sup>[2]</sup>.

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

Animal Model:	Male athymic mice (strain BALB/cAnN.Cg-Foxn1nu/CrlNarl) with FaDu cell xenografted mouse model <sup>[2]</sup>
Dosage:	2.5 and 5 mg/kg
Administration:	Oral administration; 21 days
Result:	Caused 34.7% and 76.1% reduction of the tumor growth.

#### **CUSTOMER VALIDATION**

- Sci Transl Med. 2018 Jul 18;10(450):eaaq1093.
- Front Pharmacol. 2020 Nov 11;11:580407.
- Molecules. 2020 Apr 23;25(8):1980.
- Research Square Print. 2023 Mar 9.
- Harvard Medical School LINCS LIBRARY

See more customer validations on www.MedChemExpress.com

#### REFERENCES

[1]. Markman B, et al. Phase I safety, pharmacokinetic, and pharmacodynamic study of the oral phosphatidylinositol-3-kinase and mTOR inhibitor BGT226 in patients with advanced solid tumors. Ann Oncol. 2012 Sep;23(9):2399-408.

[2]. Chang KY, et al. Novel phosphoinositide 3-kinase/mTOR dual inhibitor, NVP-BGT226, displays potent growth-inhibitory activity against human head and neck cancer cells in vitro and in vivo. Clin Cancer Res. 2011 Nov 15;17(22):7116-26.

#### 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