



## **BMS 777607**

**Catalog No: tcsc0227** 

Available Sizes
Size: 5mg
Size: 10mg
Size: 50mg
Size: 100mg
Specifications
CAS No: 1025720-94-8
Formula: C <sub>25</sub> H <sub>19</sub> CIF <sub>2</sub> N <sub>4</sub> O <sub>4</sub>
Pathway: Protein Tyrosine Kinase/RTK;Protein Tyrosine Kinase/RTK
Target: c-Met/HGFR;TAM Receptor
Purity / Grade: >98%
<b>Solubility:</b> DMSO : ≥ 39 mg/mL (76.04 mM)
Alternative Names: BMS 817378
<b>Observed Molecular Weight:</b> 512.89



## **Product Description**

BMS 777607 is a **Met-related** inhibitor for **c-Met**, **AxI**, **Ron** and **Tyro3** with **IC**<sub>50</sub>s of 3.9 nM, 1.1 nM, 1.8 nM and 4.3 nM, respectively, and 40-fold more selective for Met-related targets than Lck, VEGFR-2, and TrkA/B, with more than 500-fold greater selectivity versus all other receptor and non receptor kinases.

IC50 & Target: IC50: 3.9 nM (c-Met), 1.1 nM (AxI), 1.8 nM (Ron), 4.3 nM (Tyro3)

In Vitro: BMS 777607 is a selective ATP-competitive Met kinase inhibitor which potently blocks the autophosphorylation of c-Met with IC $_{50}$  of 20 nM in GTL-16 cell lysates, and demonstrates selective inhibition of proliferation in Met-driven tumor cell lines, such as GTL-16 cell line, H1993 and U87 $^{[1]}$ . BMS 777607 inhibits hepatocyte growth factor (HGF)-triggered c-Met autophosphorylation with IC $_{50}$  of 50 [2]. Application of BMS 777607 (appr 10  $\mu$ M) to the highly metastatic murine KHT cells for 2 hours potently eliminates basal levels of autophosphorylated c-Met with IC $_{50}$  of 10 nM without affecting the total c-Met, leading to dose-dependent inhibition of phosphorylation of downstream signaling molecules including ERK, Akt, p7056K and S6. Treatment with BMS 777607 (appr 1  $\mu$ M) for 24 hours potently inhibits the KHT cell scatter, motility and invasion at doses in the nanomolar range which consists with MET gene knockdown, and modestly affects cell proliferation and colony formation [3].

*In Vivo:* Oral administration of BMS 777607 (6.25-50 mg/kg) significantly reduces tumor volumes of the GTL-16 human tumor xenografts in athymic mice with no observed toxicity<sup>[1]</sup>. Administration of BMS 777607 (25 mg/kg/day) decreases the number of KHT lung tumor nodules (28.3%), improves the morphological hemorrhage, and significantly impairs the metastatic phenotype in the 6-8 week-old female C3H/HeJ mice injected with rodent fibrosarcoma KHT cells without apparent systemic toxicity compared to the control treatment. A low dose of BMS 777607 (10 mg/kg) also offers a mild but not significant inhibition of lung nodule formation compared to the vehicle control<sup>[3]</sup>.

All products are for RESEARCH USE ONLY. Not for diagnostic & therapeutic purposes!