



# **TG 100572**

**Catalog No: tcsc0783** 

7	

#### **Available Sizes**

Size: 5mg

Size: 10mg

Size: 50mg



# **Specifications**

#### CAS No:

867334-05-2

#### Formula:

 $C_{26}H_{26}CIN_5O_2$ 

### **Pathway:**

Protein Tyrosine Kinase/RTK; Protein Tyrosine Kinase/RTK; Protein Tyrosine Kinase/RTK; Protein Tyrosine Kinase/RTK

### **Target:**

Src;VEGFR;PDGFR;FGFR

## **Purity / Grade:**

>98%

## **Solubility:**

DMSO :  $\geq$  150 mg/mL (315.15 mM)

### **Observed Molecular Weight:**

475.97

# **Product Description**

TG 100572 is a multi-targeted kinase inhibitor which inhibits **receptor tyrosine kinases** and **Src kinases**; has  $IC_{50}$ s of 2, 7, 2, 16, 13, 5, 0.5, 6, 0.1, 0.4, 1, 0.2 nM for VEGFR1, VEGFR2, FGFR1, FGFR2, PDGFR $\beta$ , Fgr, Fyn, Hck, Lck, Lyn, Src, Yes, respectively.





IC50 & Target: IC50: 2 nM (VEGFR1), 7 nM (VEGFR2), 2 nM (FGFR1), 16 nM (FGFR2), 13 nM (PDGFRβ), 5 nM (Fgr), 0.5 nM (Fyn), 6 nM (Hck), 0.1 nM (Lck), 0.4 nM (Lyn), 1 nM (Src), 0.2 nM (Yes)<sup>[1]</sup>

In Vitro: TG 100572 shows sub-nanomolar activity against the Src family as well as RTK such as VEGFR1 and R2, FGFR1 and R2, and PDGFRβ. TG 100572 inhibits vascular endothelial cell proliferation ( $ED_{50}$ =610±71 nM) and blocks VEGF-induced phosphorylation of extracellular signal-regulated kinase. TG 100572 induces apoptosis in rapidly proliferating, but not quiescent, endothelial cell cultures [1]. TG 100572 is shown to inhibit hRMVEC cell proliferation, with an IC<sub>50</sub> of 610±72 nM. This suggests that TG 100572 has the therapeutic potential to inhibit VEGF function in ocular endothelial cells, a contributing factor to pathological angiogenesis in diseases such as AMD and PDR<sup>[2]</sup>.

In Vivo: Systemic delivery of TG 100572 in a murine model of laser-induced choroidal neovascularization (CNV) causes significant suppression of CNV, but with an associated weight loss suggestive of systemic toxicity<sup>[1]</sup>. A concentration of 23.4  $\mu$ M (C<sub>max</sub>) of TG 100572 is reached in 30 min (T<sub>max</sub>)=0.5 h) in the choroid and the sclera. However, the levels of TG 100572 in the retina are relatively low. The half-life of TG 100572 in ocular tissues is very short; hence, the compound is administered topically minimum t.i.d. to maintain appropriate drug levels in the eye. The maximum concentration one can achieve in formulations using TG 100572 is 0.7% w/v<sup>[2]</sup>.

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