# Data Sheet (Cat.No.T2605)



## Fostamatinib Disodium

Chemical Propert	ies	
CAS No. :	1025687-58-4	o~ <sup>CH</sup> 3
Formula:	C23H24FN6O9P·2Na	HNC HN CH3
Molecular Weight:	624.42	
Appearance:	no data available	
Storage:	Powder: -20°C for 3 years   In solvent: -80°C for 1 year	1 0° Nđ

## **Biological Description**

Description	Fostamatinib Disodium (R788 Disodium) is an orally available Syk kinase inhibitor with potential anti-inflammatory and immunomodulating activities.
Targets(IC50)	FLT,Syk
In vitro	In mouse tumor models, daily administration of R935788 (80 mg/kg) effectively inhibited the growth of TCL1-002, TCL1-551, and TCL1-870 tumors. In Eµ-TCL1 transgenic mice, R935788 suppressed leukemia cell proliferation and survival by blocking antigen-dependent B-cell receptor signaling.
In vivo	R406, within a range of EC50 values (0.8-8.1 μM) across various diffuse large B-cell lymphoma cell lines, reduces the phosphorylation of BLNK, Akt, GSK-3, FOXO, and ERK, thereby inhibiting cell proliferation.
Kinase Assay	In vitro fluorescence polarization kinase assays: R406 (in vitro active form of R935788) is serially diluted in DMSO and then diluted to 1% DMSO in kinase buffer (20 mM HEPES, pH 7.4, 5 mM MgCl2, 2 mM MnCl2, 1 mM DTT, 0.1 mg/mL acetylated BGG). ATP and substrate in kinase buffer are added at room temperature, resulting in a final DMSO concentration on 0.2%. The kinase reactions are performed in a final volume of 20 µL containing 5 µM HS1 peptide substrate and 4 µM ATP and started by addition of 0.125 ng of Syk in kinase buffer. The reaction is allowed to proceed for 40 minutes at room temperature. The reaction is stopped by the addition of 20 µL of PTK quench mix containing EDTA/anti-phosphotyrosine antibody (1× final)/fluorescent phosphopeptide tracer (0.5× final) diluted in FP Dilution Buffer. The plate is incubated for 30 minutes in the dark at room temperature and then read on a Polarion fluorescence polarization plate reader. Data is converted to determine the amount of phosphopeptide present using a calibration curve generated by competition with the phosphopeptide competitor provided in the Tyrosine Kinase Assay Kit. For IC50 determination, R406 is tested at eleven concentrations in duplicate and curve-fitting is performed by non-linear regression analysis using Prism GraphPad Software.
Cell Research	Cells are exposed to increasing concentrations of R406 (in vitro active form of R935788) for 48 hours. The percentage of apoptotic cells is determined by double staining with propidium iodide (PI) and annexin-A5–FITC conjugate. Ki-67 staining is performed with the FITC mouse anti–Ki-67 set. Samples are analyzed on a FACSCalibur flow cytometer with CellQuest Version 3.3 software. (Only for Reference)

## A DRUG SCREENING EXPERT

### **Solubility Information**

Solubility

Ethanol: < 1 mg/mL (insoluble or slightly soluble),<br/>H2O: < 1 mg/mL (insoluble or slightly soluble),<br/>DMSO: 50 mg/mL (80.07 mM),<br/>(&lt; 1 mg/ml refers to the product slightly soluble or insoluble)

## **Preparing Stock Solutions**

	1mg	5mg	10mg	
1 mM 📀	1.6015 mL	8.0074 mL	16.0149 mL	
5 mM	0.3203 mL	1.6015 mL	3.203 mL	
10 mM	0.1601 mL	0.8007 mL	1.6015 mL	
50 mM	0.032 mL	0.1601 mL	0.3203 mL	

Please select the appropriate solvent to prepare the stock solution, according to the solubility of the product in different solvents. Please use it as soon as possible.

#### Reference

Braselmann S, et al. J Pharmacol Exp Ther, 2006, 319(3), 998-1008. Chen L, et al. Blood, 2008, 111(4), 2230-2237.

Inhibitor • Natural Compounds • Compound Libraries • Recombinant Proteins This product is for Research Use Only• Not for Human or Veterinary or Therapeutic Use Tel:781-999-4286 E\_mail:info@targetmol.com Address:36 Washington Street,Wellesley Hills,MA 02481