# LY 294002

#### PI3 kinase inhibitor

Potent, cell permeable, inhibitor of PI(3)K (phosphoinositide 3-kinase) that acts on the ATP-binding site of the enzyme. Displays an IC $_{50}$  of 1.4µM and inhibits all isoforms equally in in vitro assays using purified PI 3-kinase (p110 $\alpha$  IC $_{50}$ =0.63µM). Inhibits Pim-1 kinase with IC $_{50}$ =4.0µM. Other kinases such as PKC, PKA, MAP kinase, S6 kinase, EGF receptor tyrosine kinase, c-src kinase, PI 4-kinase, diacylglycerol kinase or rabbit kidney ATPase are not inhibited at 50µM.

Induces apoptosis in many cell types by blocking the PI3-K/Akt antiapoptotic pathway. Useful tool for identifying cellular events which are regulated by the PI3-K/Akt axis. Sensitizes tumor cells to drug-induced apoptosis independent of its PI3K inhibitory activity. Inhibits proliferation and induces apoptosis in human colon cancer cells. Suppresses proliferation of mouse embryonic stem cells. Furthermore, inhibits autophagic sequestration in rat hepatocytes.

Citations: 51

View Online »

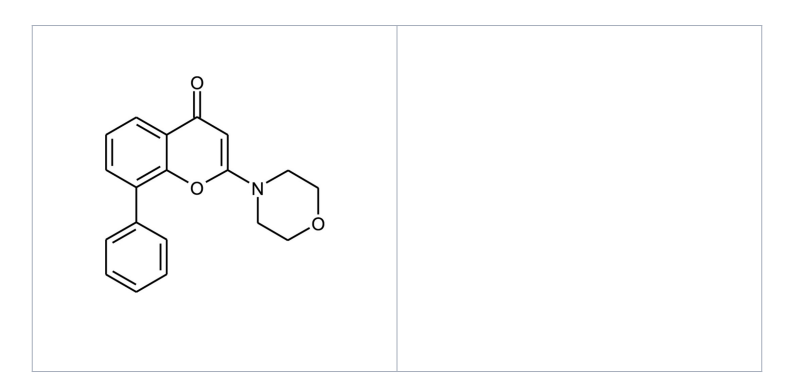
## **Ordering Information**

**Order Online** »

BML-ST420-0005	5mg
BML-ST420-0025	25mg

Manuals, SDS & CofA

View Online »



## **Handling & Storage**

**Use/Stability** As indicated on product label or CoA when stored as recommended. Stable for at least

1 year after receipt when stored at -20°C. Stock solutions are stable for up to 3 months

at -20°C.

Long Term Storage -20°C

**Shipping** Ambient Temperature

# Regulatory Status RUO - Research Use Only

### **Product Details**

Alternative Name 2-(4-Morpholinyl)-8-phenyl-4H-1-benzopyran-4-one

**Appearance** White to pale yellow solid.

**CAS** 154447-36-6

Couple Target PI3 kinase, Pim

Couple Type Inhibitor

Formula  $C_{19}H_{17}NO_3$ 

**MW** 307.3

Purity ≥98% (HPLC)

Soluble in DMSO (25mg/ml, warm), 100% ethanol

(25mg/ml, warm) or methanol.

Technical Info / Product Notes Replacement for ADI-HPK-106.

Last modified: May 29, 2024

