# **Orlistat**

#### Fatty acid synthase inhibitor

Cell permeable, irreversible inhibitor of gastric and pancreatic lipases. Shows only minimal activity against amylase, trypsin, chymotrypsin, or phospholipase  $A_2$  (PLA $_2$ ). Partially inhibits the hydrolysis of triglycerides and lowers the absorption of dietary fat and promotes weight loss. Antiobesity drug. Exhibits antitumor activity by inhibition of the thioesterase domain of fatty acid synthase (FAS) both *in vitro* and *in vivo*.

Citations: 8

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**Ordering Information** 

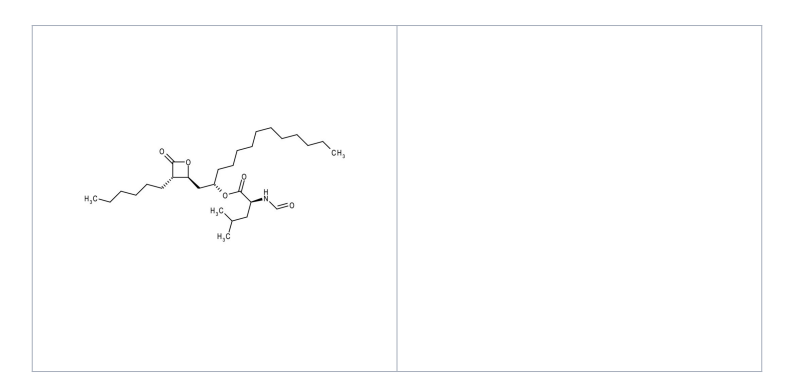
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ALX-350-152-M250

250mg

Manuals, SDS & CofA

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### **Handling & Storage**

**Use/Stability** As indicated on product label or CoA when stored as recommended. Stock solutions are

stable for up to 6 weeks when stored at -20°C.

**Handling** Protect from light and moisture. After reconstitution, prepare aliquots and store at -20°C.

Long Term Storage -20°C

**Shipping** Ambient Temperature

## Regulatory Status RUO - Research Use Only

#### **Product Details**

Alternative Name Tetrahydrolipstatin, N-Formyl-L-leucine-(1S)-1-(((2S,3S)-3-hexyl-4-oxo-2-

oxetanyl)methyl)dodecyl ester

**Appearance** White to off-white solid.

**CAS** 96829-58-2

Couple Target Fatty acid synthase

Couple Type Inhibitor

Formula  $C_{29}H_{53}NO_5$ 

MI 14: 6869

**MW** 495.7

Purity ≥97%

**Soluble** in DMSO or 100% ethanol.

**Source** Synthetic. Originally isolated from *Streptomyces* sp.



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