# Data Sheet (Cat.No.T18205)



### m-PEG48-Mal

### **Chemical Properties**

CAS No.:

Formula: C104H202N2O51

Molecular Weight: 2296.7

Appearance: no data available

Storage: Powder: -20°C for 3 years | In solvent: -80°C for 1 year

## **Biological Description**

Description	m-PEG48-Mal is a PEG-based linker for PROTACs which joins two essential ligands, crucial for forming PROTAC molecules. This linker enables selective protein degradation by leveraging the ubiquitin-proteasome system within cells.
Targets(IC50)	Others
In vitro	PROTACs contain two different ligands connected by a linker; one is a ligand for an E3 ubiquitin ligase and the other is for the target protein. PROTACs exploit the intracellular ubiquitin-proteasome system to selectively degrade target proteins[1].

## **Preparing Stock Solutions**

	1mg	5mg	10mg	
1 mM	0.4354 mL	2.177 mL	4.3541 mL	
5 mM	0.0871 mL	0.4354 mL	0.8708 mL	
10 mM	0.0435 mL	0.2177 mL	0.4354 mL	
50 mM	0.0087 mL	0.0435 mL	0.0871 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

An S, et al. Small-molecule PROTACs: An emerging and promising approach for the development of targeted therapy drugs. EBioMedicine. 2018 Oct;36:553-562

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