



Lodelaben

Catalog No: tcsc0018391

且	Available Sizes
Size: 1	mg
Size: 5	mg
Size: 1	0mg
	Specifications
CAS No 111149	
Formu	
Pathw a Metabo	ay: lic Enzyme/Protease
Target Elastas	
Purity >98%	/ Grade:
Solubi 10 mM	l ity: in DMSO
	ative Names: 26;Declaben
Observ 425.04	ved Molecular Weight:
Produ	uct Description





Lodelaben is a human neutrophil **elastase** inhibitor with an IC_{50} and K_i of 0.5 and 1.5 μ M, respectively.

IC50 & Target: IC50: 0.5 μ M (elastase)^[1]

Ki: 1.5 μ M (elastase)^[1]

In Vitro: Lodelaben is a human neutrophil elastase inhibitor with an IC $_{50}$ and K $_{i}$ of 0.5 and 1.5 μ M, respectively. Results indicate that the inhibition of human neutrophil elastase (HNE) by Lodelaben is non-competetive. Lodelaben is not inhibitory at 10 μ M with the synthetic substrates or at 5 μ M vith Azocoll. Pseudomonas aeruginosa elastase, a metallo-protease is not inhibited by Lodelaben. Cathepsin G activity, however, is inhibited by Lodelaben, with an IC $_{50}$ of approximately 2.5 μ M, with Azocoll as substrate^[1].

In Vivo: The mean pulmonary artery pressures of the saline/vehicle and saline/Lodelaben groups are similar, 16.4 ± 1.1 and 17.4 ± 0.9 mm Hg, respectively. Although, mean pulmonary artery pressure in the monocrotaline/vehicle group is 27.5 ± 0.8 mm Hg, treatment of monocrotaline rats with Lodelaben results in significantly lower values (21.00 ± 1.6 mm Hg, p[2].

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