



Grazoprevir potassium salt

Catalog No: tcsc1375

Available Sizes
Size: 5mg
Size: 10mg
Size: 50mg
Size: 100mg
Specifications
CAS No: 1206524-86-8
Formula: $C_{38}^{H}_{49}^{KN}_{6}^{O}_{9}^{S}$
Pathway: Metabolic Enzyme/Protease;Anti-infection
Target: HCV Protease;HCV
Purity / Grade: >98%
Solubility: 10 mM in DMSO
Alternative Names: MK-5172 (potassium salt)
Observed Molecular Weight: 804.99



Product Description

Grazoprevir potassium salt (MK-5172 potassium salt) is a selective inhibitor of **Hepatitis C virus NS3/4a** protease with broad activity across genotypes and resistant variants, with $\mathbf{K}_{\mathbf{i}}$ s of 0.01 nM (gt1b), 0.01 nM (gt1a), 0.08 nM (gt2a), 0.15 nM (gt2b), 0.90 nM (gt3a), respectively.

IC50 & Target: Ki: 0.01±[1]

In Vitro: In biochemical assays, Grazoprevir (MK-5172) is effective against a panel of major genotypes and variants engineered with common resistant mutations, with K_i of $0.01\pm R155K$), 0.14 ± 0.03 nM (gt1b^{D168V}), 0.30 ± 0.04 nM (gt1b^{D168Y}), 5.3 ± 0.9 nM (gt1b^{A156T}), and 12 ± 2 nM (gt1b^{A156V}), respectively. In the replicon assay, Grazoprevir demonstrates subnanomolar to low-nanomolar EC₅₀s against genotypes 1a, 1b, and 2a, with EC₅₀s of 0.5 ± 0.1 nM, 2 ± 1 nM, and 2 ± 1 nM for gt1b^{Con1}, gt1a, and gt2a, respectively. Grazoprevir is potent against a panel of HCV replication mutants NS5A (Y93H) (EC₅₀=0.7±0.3 nM), NS5B nucleosides (S282T) (EC₅₀=0.3±0.1 nM), and NS5B (C316Y) (EC₅₀=0.4±0.2)^[1]. Grazoprevir (MK-5172) maintains the excellent potency against the gt 3a enzyme as well as a broad panel of mutant enzymes, has excellent potency in the replicon system [gt1b IC₅₀(50% NHS)=7.4 nM; gt1a IC₅₀(40% NHS)=7 nM], and shows excellent rat liver exposure^[2].

In Vivo: Grazoprevir (MK-5172) demonstrates efficacy in vivo against chronic-HCV-infected chimpanzees [1]. When dosed to dogs, Grazoprevir (MK-5172) shows low clearance of 5 mL/min/kg and a 3 h half-life after iv dosing and has good plasma exposure (AUC=0.4 μ M h) after a 1 mg/kg oral dose. Dog liver biopsy studies showed that the liver concentration of Grazoprevir after the 1 mg/kg oral dose is 1.4 μ M at the 24 h time point. Similar to its behavior in rats, Grazoprevir demonstrates effective partitioning into liver tissue and maintains high liver concentration, relative to potency, 24 h after oral dosing in dogs [2].

All products are for RESEARCH USE ONLY. Not for diagnostic & therapeutic purposes!