Product Data Sheet

PMX-53

Cat. No.: HY-106178 CAS No.: 219639-75-5 Molecular Formula: C47H65N11O7 Molecular Weight: 896.09

Sequence Shortening: F-{Orn}-P-{d-Cha}-WR (Lactam bridge: Orn2- Arg6)

Target: **Complement System**

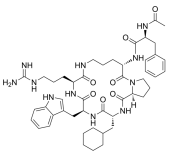
Pathway: Immunology/Inflammation

Sealed storage, away from moisture and light, under nitrogen Storage:

> Powder -80°C 2 years -20°C 1 year

* In solvent: -80°C, 2 years; -20°C, 1 year (sealed storage, away from moisture and

light, under nitrogen)



SOLVENT & SOLUBILITY

In Vitro

DMSO: 200 mg/mL (223.19 mM; Need ultrasonic)

H₂O: 2.5 mg/mL (2.79 mM; ultrasonic and warming and heat to 60°C)

Preparing Stock Solutions	Solvent Mass Concentration	1 mg	5 mg	10 mg
	1 mM	1.1160 mL	5.5798 mL	11.1596 mL
	5 mM	0.2232 mL	1.1160 mL	2.2319 mL
	10 mM	0.1116 mL	0.5580 mL	1.1160 mL

Please refer to the solubility information to select the appropriate solvent.

In Vivo

- 1. Add each solvent one by one: 10% DMSO >> 40% PEG300 >> 5% Tween-80 >> 45% saline Solubility: ≥ 2.08 mg/mL (2.32 mM); Clear solution
- 2. Add each solvent one by one: 10% DMSO >> 90% (20% SBE-β-CD in saline) Solubility: ≥ 2.08 mg/mL (2.32 mM); Clear solution
- 3. Add each solvent one by one: 10% DMSO >> 90% corn oil Solubility: ≥ 2.08 mg/mL (2.32 mM); Clear solution

BIOLOGICAL ACTIVITY

Description

PMX-53 (3D53) is a synthetic peptidic and a potent and orally active complement C5a receptor (CD88) antagonist with an IC 50 of 20 nM. PMX-53 is also a low-affinity MrgX2 agonist that stimulates MrgX2-mediated mast cell degranulation. PMX-53 specifically binds to C5aR1 and does not bind to the second C5aR (C5L2) and C3aR. PMX-53 has anti-inflammatory, anticancer and antiatherosclerotic effects^{[1][2][3][4][5][6]}.

IC ₅₀ & Target	IC50: 20 nM (Complement C5a receptor) ^[4] MrgX2 ^[1]		
In Vitro	PMX-53 is a potent CD88 antagonist and inhibits C5a-induced neutrophil myeloperoxidase release and chemotaxis with IC ₅₀ values of 22 nM and 75 nM, respectively ^[1] . ?PMX-53 (10 nM) inhibits C5a-induced Ca ²⁺ mobilization in HMC-1 cells, but at higher concentrations(≥30 nM) it causes degranulation in LAD2 mast cells, CD34 ⁺ cell-derived mast cells, and RBL-2H3 cells stably expressing MrgX2. Replacement of Trp with Ala and Arg with dArg abolishes the ability of PMX-53 to inhibit C5a-induced Ca ²⁺ mobilization in HMC-1 cells and to cause degranulation in RBL-2H3 cells expressing MrgX2 ^[1] . MCE has not independently confirmed the accuracy of these methods. They are for reference only.		
In Vivo	PMX-53 (0.3-3?mg/kg; subcutaneous injection; once; male Wistar rats) treatment inhibits the hypernociception induced by zymosan-activated serum and C5a but not by the direct-acting hypernociceptive mediators, prostaglandin E2 and dopamine [2]. Local pretreatment of rats with PMX-53 (60-180?μg per paw) inhibits zymosan-, carrageenan-, lipopolysaccharide (LPS)- and antigen-induced hypernociception ^[2] . Pharmacokinetic analyses have shown that PMX-53 (3D53) appears in the plasma within 5 min of oral administration (3 mg/kg) to rats, with peak blood levels of approximately 0.3 μM being reached within 20 min The plasma elimination half-life was approximately 70 min in this case ^[3] . The non-acetylated version of PMX-53 (3D53) binds to isolated mouse neutrophils with a K _d value of 30 nM (mouse C5a binds with a K _d value of 0.3 nM) and inhibits mouse C5a-induced chemotaxis with an IC ₅₀ value of 0.5 nM ^[3] . MCE has not independently confirmed the accuracy of these methods. They are for reference only. Animal Model: Adult male Wistar rats (weighing 180-200 g) injected with zymosan ^[2] Dosage: 0.3 mg/kg, 1 mg/kg or 3 mg/kg Administration: Subcutaneous injection; once Result: Inhibited the hypernociception induced by zymosan-activated serum and C5a.		

CUSTOMER VALIDATION

- Mol Ther. 2023 May 3;S1525-0016(23)00256-3.
- Research Square Print. November 28th, 2022.

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REFERENCES

- [1]. Subramanian H, et al. PMX-53 as a dual CD88 antagonist and an agonist for Mas-related gene 2 (MrgX2) in human mast cells. Mol Pharmacol. 2011 Jun;79(6):1005-13.
- [2]. Ting E, et al. Role of complement C5a in mechanical inflammatory hypernociception: potential use of C5a receptor antagonists to control inflammatory pain. Br J Pharmacol. 2008 Mar;153(5):1043-53.
- [3]. Holland MC, et al. Synthetic small-molecule complement inhibitors. Curr Opin Investig Drugs. 2004 Nov;5(11):1164-73.
- [4]. Finch AM, et al. Low-molecular-weight peptidic and cyclic antagonists of the receptor for the complement factor C5a. J Med Chem. 1999 Jun 3;42(11):1965-74.
- $[5]. \ Manthey\ HD,\ et\ al.\ Complement\ C5a\ inhibition\ reduces\ atherosclerosis\ in\ ApoE-/-\ mice.\ FASEB\ J.\ 2011\ Jul; 25(7): 2447-55.$
- [6]. Vadrevu SK, et al. Complement c5a receptor facilitates cancer metastasis by altering T-cell responses in the metastatic niche. Cancer Res. 2014 Jul 1;74(13):3454-65.

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 $\label{lem:caution:Product} \textbf{Caution: Product has not been fully validated for medical applications. For research use only.}$

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