Screening Libraries

Product Data Sheet

Proflavine hemisulfate

Cat. No.: HY-B0883 CAS No.: 1811-28-5

Molecular Formula: $C_{13}H_{11}N_{3}\cdot 1/2H_{2}SO_{4}$

Molecular Weight: 258.28

Target: Bacterial; Autophagy; Potassium Channel

Pathway: Anti-infection; Autophagy; Membrane Transporter/Ion Channel

4°C, sealed storage, away from moisture and light Storage:

* In solvent: -80°C, 6 months; -20°C, 1 month (sealed storage, away from moisture

and light)

$$H_2N$$
 N N N

0.5H₂SO₄

SOLVENT & SOLUBILITY

In Vitro

 $H_2O : \ge 5 \text{ mg/mL } (19.36 \text{ mM})$

* "≥" means soluble, but saturation unknown.

	Solvent Mass Concentration	1 mg	5 mg	10 mg
Preparing Stock Solutions	1 mM	3.8718 mL	19.3588 mL	38.7177 mL
	5 mM	0.7744 mL	3.8718 mL	7.7435 mL
	10 mM	0.3872 mL	1.9359 mL	3.8718 mL

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

BIOLOGICAL ACTIVITY

 $Proflavine\ hemisulfate, an\ acridine\ dye, is\ a\ known\ DNA\ intercalating\ agent.\ Anti-microbial\ agent \ [1].\ Proflavine\ hemisulfate$ Description

behaves as a pore blocker for $K_{ir}3.2$. Proflavine hemisulfate is a potential lead compound for $K_{ir}3.2$ -associated neurological

diseases^[2].

Proflavine (0.1-10 μ M; 24 hours) inhibits the growth of $K_{ir}3.2$ -transformant cells and $K_{ir}3.2$ activity in a concentration-In Vitro

dependent manner^[1].

Proflavine (300 μ M) progressively reduces the current amplitude of K_{ir}3.2 mutant to 27.7±4.3% of the control^[2].

MCE has not independently confirmed the accuracy of these methods. They are for reference only.

Cell Viability Assay^[2]

Cell Line:	K _{ir} 3.2 [*] -transformant BYT123 cells	
Concentration:	0.1, 1, and 10 μM	
Incubation Time:	24 hours	

Result:	Dose-dependent inhibition of the growth of K _{ir} 3.2*-transformant cells.
	Attenuated the growth of K _{ir} 3.2*-transformant cells without affecting the growth of contro
	cells.

In Vivo

The concentrations of Proflavine (20 mg/kg) in whole blood after intravenous injection decreased rapidly at the beginning and remained stable from around 30 min after dosing [3].

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Animal Model:	Adult male Sprague Dawley rats (weighing approximately 200 g) $^{[3]}$	
Dosage:	20 mg/kg (Pharmacokinetic Analysis)	
Administration:	Intravenous injection; 2, 4, 5, 10, 15, 20, 25, and 30 min after dosing	
Result:	Concentration decreased rapidly from whole blood in the first 5 min after dosing, by a slower decrease.	

CUSTOMER VALIDATION

• EMBO Rep. 2022 Apr 11;e53932.

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REFERENCES

- [1]. Hitoshi Kawada, et al. Isolation of proflavine as a blocker of G protein-gated inward rectifier potassium channels by a cell growth-based screening system. Neuropharmacology. 2016 Oct;109:18-28.
- [2]. Mansour K.Gatasheh, et al. Proflavine an acridine DNA intercalating agent and strong antimicrobial possessing potential properties of carcinogen. Karbala International Journal of Modern Science. 2017 Dec, 3(4): 272-278.
- [3]. Jiaxin Chen, et al. Determination of proflavine in rat whole blood without sample pretreatment by laser desorption postionization mass spectrometry. Anal Bioanal Chem. 2017 Apr;409(11):2813-2819.

Caution: Product has not been fully validated for medical applications. For research use only.

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