# **Product** Data Sheet

# **UNC0379 TFA**

Cat. No.: HY-12335A CAS No.: 1620401-83-3 Molecular Formula:  $C_{25}H_{36}F_3N_5O_4$ Molecular Weight: 527.58

Target: Histone Methyltransferase

Pathway: **Epigenetics** 

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

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

#### **SOLVENT & SOLUBILITY**

In Vitro

H<sub>2</sub>O: 100 mg/mL (189.54 mM; Need ultrasonic)

Preparing Stock Solutions	Solvent Mass Concentration	1 mg	5 mg	10 mg
	1 mM	1.8954 mL	9.4772 mL	18.9545 mL
	5 mM	0.3791 mL	1.8954 mL	3.7909 mL
	10 mM	0.1895 mL	0.9477 mL	1.8954 mL

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

### **BIOLOGICAL ACTIVITY**

Description UNC0379 TFA is a selective, substrate-competitive inhibitor of lysine methyltransferase SETD8 (KMT5A) with an IC50 of 7.3  $\mu$ 

M, K<sub>D</sub> value of 18.3 μM. UNC0379 TFA can be used in the research of inflammation and cancers, such as pulmonary fibrosis,

ovarian cancer, neuroblastoma<sup>[1][2][3]</sup>.

SETD8 (KMT5A)[1] IC<sub>50</sub> & Target

UNC0379 TFA (1-10 μM, 9 days) inhibits HGSOC cells proliferation<sup>[2]</sup>. In Vitro

UNC0379 TFA (10  $\mu$ M, 96 h) increases in the proportion of sub-G1 phase cells in HGSOC cells [2].

UNC0379 TFA (10  $\mu$ M, 48 h) induces myofibroblast de-differentiation and inhibits additional fibroblast to myofibroblast

differentiation<sup>[3]</sup>.

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

Cell Viability Assay<sup>[1]</sup>

Cell Line:	JHOS2, JHOS3, JHOS4, OVCAR3, OVCAHO, OVKATE, KURAMOCHI, TYKnu
Concentration:	1-10 μΜ

Incubation Time:	9 days	
Result:	Inhibited HGSOC cells proliferation with IC $_{\!50}\text{s}$ ranging from 0.39 to 3.20 $\mu\text{M}.$	
Cell Cycle Analysis <sup>[1]</sup>		
Cell Line:	JHOS3, OVCAR3	
Concentration:	10 μΜ	
Incubation Time:	96 h	
Result:	Arrested cells in sub-G1 phase.	

#### In Vivo

 $\label{lem:condition} UNC0379\ TFA\ (intratracheal\ administration, 1\ mg/kg/day, 1\ mg/kg/day, on\ day 7, 8,\ and\ 9)\ ameliorates\ the\ lung\ fibrosis\ in\ Bleomycin\ (BLM)-induced\ lung\ fibrosis\ mouse^{\left[3\right]}.$ 

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

Animal Model:	Bleomycin (BLM)-induced lung fibrosis mouse model <sup>[3]</sup>	
Dosage:	1 mg/kg/day	
Administration:	Intratracheal administration, on day7, 8, and 9.	
Result:	Ameliorated BLM-induced lung fibrosis (supported by the evaluation of the Ashcroft score and changes in the collagen content in the lung samples) without affecting pulmonary inflammation.	

## **CUSTOMER VALIDATION**

- Cell Metab. 2021 Jan 5;33(1):160-173.e6.
- Proc Natl Acad Sci U S A. 2019 Feb 19;116(8):2961-2966.
- Cell Death Dis. 2018 Jan 26;9(2):129.
- Sci Rep. 2020 Mar 11;10(1):4490.
- J Gastroenterol Hepatol. 2021 May 14.

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#### **REFERENCES**

- $[1]. \ Miku \ Wada, et al. \ Epigenetic \ Modifier \ SETD8 \ as \ a \ The rapeutic \ Target \ for \ High-Grade \ Serous \ Ovarian \ Cancer. \ Biomolecules. \ 2020 \ Dec \ 16;10(12):1686.$
- [2]. Keita Ugai, et al. Inhibition of the SET8 Pathway Ameliorates Lung Fibrosis Even Through Fibroblast Dedifferentiation. Front Mol Biosci. 2020 Aug 5;7:192.
- [3]. Ma A, et al. Discovery of a Selective, Substrate-Competitive Inhibitor of the Lysine Methyltransferase SETD8. J Med Chem. 2014 Aug 14;57(15):6822-33.

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

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