NXPZ-2

Cat. No.:	HY-149010	
CAS No.:	2254492-08-3	
Molecular Formula:	C ₂₇ H ₂₇ N ₅ O ₇ S ₂	
Molecular Weight:	597.66	
Target:	Keap1-Nrf2	, i i i i i i i i i i i i i i i i i i i
Pathway:	NF-κB	S NH2
Storage:	Please store the product under the recommended conditions in the Certificate of Analysis.	H ₂ N Ö

SOLVENT & SOLUBILITY

In Vitro

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Preparing Stock Solutions	Mass Solvent Concentration	1 mg	5 mg	10 mg
	1 mM	1.6732 mL	8.3660 mL	16.7319 m
	5 mM	0.3346 mL	1.6732 mL	3.3464 m
	10 mM	0.1673 mL	0.8366 mL	1.6732 ml

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

BIOLOGICAL ACTIV			
Description	NXPZ-2 is an orally active Keap1-Nrf2 protein–protein interaction (PPI) inhibitor with a K _i value of 95 nM, EC ₅₀ value of 120 and 170 nM. NXPZ-2 can dose-dependently ameliorate Aβ _[1-42] -Induced cognitive dysfunction, improve brain tissue pathological changes in Alzheimer's disease (AD) mouse by increasing neuron quantity and function. NXPZ-2 can inhibit oxidative stress by increasing Nrf2 expression levels and promoting its cytoplasm to nuclear translocation, which is helpful for Keap1-Nrf2 PPI inhibitors and AD associated disease research ^[1] .		
IC ₅₀ & Target	K _i : 95 nM (Keap 1); EC ₅₀ : 120 and 170 nM (Keap 1) $^{[1]}$		
In Vitro	NXPZ-2 (0-200 μM, 7 days) has no obvious toxicity on primary cortical neuron ^[1] . MCE has not independently confirmed the accuracy of these methods. They are for reference only. Cell Cytotoxicity Assay ^[1]		
	Cell Line:	primary cortical neuron	
	Concentration:	0, 1.6, 8, 40 and 200 μM	
	Incubation Time:	7 days	

Product Data Sheet

	Result:	Had no obvious toxicity on primary cortical neuron ^[1] .
In Vivo	NXPZ-2 (Male ICR mice, including increased spo increased the time sper NXPZ-2 (Male ICR mice, neuron numbers of AD NXPZ-2 (Male ICR mice, expression levels, and p Nrf2 in both the central MCE has not independe	, 52.5/105/210 mg/kg, p.o., once daily for 7 days) improves AD mice learning and memorizing function ontaneous alternation, increases number of active avoidance times, shortened escape latency, nt in the target quadrant and number of platform crossing ^[1] . , 52.5/105/210 mg/kg, p.o., once daily for 7 days) rescues the brain structure damage and lowers dead mice with no obvious toxicity on mouse organs ^[1] . , 52.5/105/210 mg/kg, p.o., once daily for 7 days) alleviates oxidative stress by increasing Nrf2 promotes Nrf2's cytoplasm to nuclear translocation, and improves cognitive dysfunction by elevating I nervous system and peripheral blood ^[1] . ently confirmed the accuracy of these methods. They are for reference only.
	Animal Model:	Male ICR mice, AD model ^[1] .
	Dosage:	52.5 mg/kg, 105 mg/kg, 210 mg/kg.
	Administration:	P.o., once daily for 7 days.
	Result:	Showed statistically increased spontaneous alternation and no influence on basal motivation, displayed an increased number of active avoidance times, which improved the learning and memory ability of AD mice. Cell number and morphology in NPZX-2-treated groups were restored, dead neuron numbers of AD mice was lowered. Increased serum Nrf2 level, displays more Nrf2 in the hippocampal and cortical nucleus and less expression level in the hippocampal and cortical cytoplasm. Increased Nrf2-ARE binding in both hippocampus and frontal cortex, dose-dependently restored SOD, GSH,

REFERENCES

[1]. Yi Sun, et al. Direct inhibition of Keap1-Nrf2 Protein-Protein interaction as a potential therapeutic strategy for Alzheimer's disease. Bioorg Chem. 2020 Oct;103:104172.

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

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