

Produktinformation



Forschungsprodukte & Biochemikalien
Zellkultur & Verbrauchsmaterial
Diagnostik & molekulare Diagnostik
Laborgeräte & Service

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Zuschläge

- Mindermengenzuschlag
- Trockeneiszuschlag
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- Expressversand

SZABO-SCANDIC HandelsgmbH

Quellenstraße 110, A-1100 Wien T. +43(0)1 489 3961-0 F. +43(0)1 489 3961-7 <u>mail@szabo-scandic.com</u> www.szabo-scandic.com

Data Sheet (Cat.No.T0191)



Linagliptin

| Chemical Proper | ties |
|-------------------|---|
| CAS No. : | 668270-12-0 сн, сн, |
| Formula: | C25H28N8O2 |
| Molecular Weight: | |
| Appearance: | no data available |
| Storage: | keep away from direct sunlight,store under nitrogen Powder: -20°C for 3 years In solvent: -80°C for 1 year |

| Linagliptin (BI 1356) is a potent, orally bioavailable dihydropurinedione-based inhibitor of dipeptidyl peptidase 4 (DPP-4), with hypoglycemic activity. | | |
|--|--|--|
| Ferroptosis,Proteasome,DPP-4,Autophagy | | |
| Linagliptin shows a potent inhibition effect against DPP-4 in vitro and a low affinity for hERG channel and M1 receptor (IC50 295 nM). [1] Linagliptin acts as a competitive inhibitor with a Ki of 1 nM, and also shows 10,000-fold more selectivity for DPP-4 than DPP-8, DPP-9, amino-peptidases N and P, prolyloligopeptidase, trypsin, plasmin, and thrombin, and 90-fold more selectivity than fibroblast activation protein in vitro. [2] | | |
| In male Wistar rats, Beagle dogs, and Rhesus monkeys, Linagliptin shows a highly efficacious, long-lasting, and potent inhibitory activity against DPP-4 by more than 70% inhibition for all three species after oral administration of 1 mg/kg. Oral administration of Linagliptin to db/db mice 45 min before an oral glucose tolerance test reduces plasma glucose excursion in a dose-dependent manner from 0.1 mg/kg (15% inhibition) to 1 mg/kg (66% inhibition). [1] By inhibiting DPP-4 activity, Linagliptin reduces the expression of the proinflammatory markers cyclooxygenase-2 and macrophage inflammatory protein-2, and enhances the formation of myofibroblasts in healing wounds from ob/ob mice. [3] | | |
| EDTA plasma (20 μ L) is diluted with 30 μ L of DPP-4 assay buffer (100 mM Tris and 100 mM NaCl, adjusted to pH 7.8 with HCl) and mixed with 50 μ L of H-Ala-Pro-7-amido-4-trifluoromethylcoumarin. The 200 mM stock solution in dimethylformamide is diluted 1: 1000 with water to yield a final concentration of 100 μ M. The plate is incubated at room temperature for 10 min, and fluorescence in the wells is determined by using a Victor 1420 Multilabel Counter at an excitation wavelength of 405 nm and an emission wavelength of 535 nm. For the detection of DPP-4 activity in wound lysates, 100 μ g of protein from the respective wound lysates are used instead of 20 μ L of plasma. Active GLP-1 is also detected from 100 μ g of respective wound tissue samples and analyzed by using the Mouse/Rat Total Active GLP-1 Assay Kit. | | |
| A total of 4.0×107 keratinocytes per well are seeded into 24-well plates. After reaching 50% confluence, cells are starved for 24 h with DMEM. Proliferation of cells is assessed by using 1 µCi/mL of [3H]methyl-thymidine in DMEM in the presence of 10% fetal bovine serum and increasing concentrations of linagliptin (3, 30, 300, or 600 nM) for 24 h. Cells are then washed twice with phosphate-buffered saline and incubated in 5% | | |
| | | |

A DRUG SCREENING EXPERT

trichloroacetic acid at 4°C for 30 min, and the DNA is solubilized in 0.5mol/LNaOH for 30 min at 37°C. Finally, [3H]thymidine incorporation is determined.

| Solubility Information | | | | |
|---------------------------|--|------------|------------|--|
| Solubility | DMSO: 5.63 mg/mL (11.9 mM),Sonication is recommended. br/>H2O: < 1 mg/mL (insoluble or slightly soluble), Ethanol: 1 mg/mL (2.11 mM), (< 1 mg/ml refers to the product slightly soluble or insoluble) | | | |
| Preparing Stock Solutions | | | | |
| | 1mg | 5mg | 10mg | |
| 1 mM | 2.1162 mL | 10.5811 mL | 21.1622 mL | |
| 5 mM | 0.4232 mL | 2.1162 mL | 4.2324 mL | |
| 10 mM | 0.2116 mL | 1.0581 mL | 2.1162 mL | |
| 50 mM | 0.0423 mL | 0.2116 mL | 0.4232 mL | |

Please select the appropriate solvent to prepare the stock solution, according to the solubility of the product in different solvents. Please use it as soon as possible.

Reference

Eckhardt M, et al. J Med Chem. 2007, 50(26), 6450-6453. Thomas L, J Pharmacol Exp Ther. 2008, 325(1), 175-182.

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Tel:781-999-4286 E_mail:info@targetmol.com Address:36 Washington Street,Wellesley Hills,MA 02481