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DATASHEET

Cmpd101

Product overview

 Name
 Cmpd101

 Cat No
 HB2840

Alternative names Compound 101; Takeda compound 101

Biological action Inhibitor >98%

Customer comments We would recommend Cmpd 101 from Hello Bio – it performs exactly as expected in assays looking

at MOPr desensitisation, phosphorylation and internalisation. Dr Chris Bailey, University of Bath,

UK and author on Mol Pharmacol paper, PubMed ID 26013542

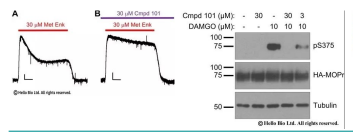
Your Cmpd101 - worked great! Dr Steven Gee, Pfizer Neuroscience, USA

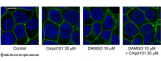
Your Cmpd101 behaved as expected. Verified customer, Monash University

Novel, potent and selective GRK2/GRK3 inhibitor

Images

Description







Biological Data

Biological description

Cmpd101 (Compound 101) is a novel, potent and selective G-protein coupled receptor kinase 2 and 3 (GRK2/GRK3) inhibitor (IC $_{50}$ values are 35 and 32 nM at GRK2 and GRK3 respectively).

Shows no activity at GRK5 at concentrations up to 125 μ M and shows little activity at a broad range of other kinases.

Membrane permeable.

Cmpd101 can be used to study roles of GRK2/3 in GPCR desensitization and other functions.

Shown to potentiate phosphatidylinositol 4,5-bisphosphate (PIP2) depletion and slow agonist-induced desensitization of protease-activated receptor 2 (PAR2).

Solubility & Handling

Solubility overview

Handling

Important

Soluble in DMSO (100mM)

Hydroscopic solid, contact with air may cause material to become sticky. Product performance should

not be affected but we recommend storing the material in a sealed jar.

This product is for RESEARCH USE ONLY and is not intended for therapeutic or diagnostic use. Not

for human or veterinary use.

Chemical Data

Chemical name 3-[(4-methyl-5-pyridin-4-yl-1,2,4-triazol-3-yl)methylamino]-N-[[2-(trifluoromethyl)phenyl]methyl]benzam

ide hydrochloride

Molecular Weight

Chemical structure

502.92

SMILES CN1C(=NN=C1C2=CC=NC=C2)CNC3=CC=CC(=C3)C(=O)NCC4=CC=CC=C4C(F)(F)F

Source Synthetic

InChl=1S/C24H21F3N6O/c1-33-21(31-32-22(33)16-9-11-28-12-10-16)15-29-19-7-4-6-17(13-19)23(

34)30 - 14 - 18 - 5 - 2 - 3 - 8 - 20(18)24(25, 26)27/h2 - 13, 29H, 14 - 15H2, 1H3, (H, 30, 34)24(25, 26)27/h2 - 13, 29H, 14 - 15H2, 1H3, (H, 30, 34)24(25, 26)27/h2 - 13, 29H, 14 - 15H2, 1H3, (H, 30, 34)24(25, 26)27/h2 - 13, 29H, 14 - 15H2, 1H3, (H, 30, 34)24(25, 26)27/h2 - 13, 29H, 14 - 15H2, 1H3, (H, 30, 34)24(25, 26)27/h2 - 13, 29H, 14 - 15H2, 1H3, (H, 30, 34)24(25, 26)27/h2 - 13, 29H, 14 - 15H2, 1H3, (H, 30, 34)24(25, 26)27/h2 - 13, 29H, 14 - 15H2, 1H3, (H, 30, 34)24(25, 26)24

InChiKey WFOVEDJTASPCIR-UHFFFAOYSA-N

Appearance Yellow solid

References

Molecular mechanism of selectivity among G protein-coupled receptor kinase 2 inhibitors.

Thal et al (2011) Mol Pharmacol 80

PubMedID 21596927

Role of G Protein-Coupled Receptor Kinases 2 and 3 in μ-Opioid Receptor Desensitization and Internalization.

Lowe et al (2015) Mol Pharmacol 88(2)

PubMedID 26013542

Contributions of protein kinases and β-arrestin to termination of protease-activated receptor 2 signaling.

Jung et al (2016) J Gen Physiol 147(3)

PubMedID 26927499

Distinct cortical and striatal actions of a β-arrestin-biased D2 receptor ligand reveal unique antipsychotic-like properties.

Urs et al (2016) Proc Natl Acad Sci U S A 113(50) **PubMedID**27911814

Agonist-selective recruitment of engineered protein probes and of GRK2 by opioid receptors in living cells

Stoeber et al (2019) bioRxiv https://doi.org/10.1101/866780