

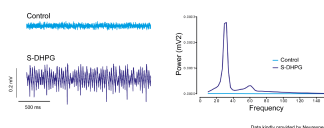
DATASHEET

(S)-3,5-DHPG

Product overview

Name	(S)-3,5-DHPG
Cat No	HB0045
Biological action	Agonist
Purity	>99%
Description	Selective group I mGlu receptor agonist. Induces LTD.

Images



Biological Data

Biological description	(S)-3,5-DHPG is a widely used selective group I mGlu receptor agonist (EC_{50} values are 2 and 6.6 μ M at mGlu ₅ and mGlu ₁ , 106 μ M at mGlu ₃ and >1000 μ M at mGlu _{2,4,7,8} receptors respectively).
	(S)-3,5-DHPG is the active enantiomer of DHPG. The racemic (R,S)-DHPG is also available.
	(S)-3,5-DHPG induces long term depression (LTD) (mGluR/DHPG-LTD). DHPG-LTD can temporarily be reversed by other mGluR antagonists such as (S)-MCPG.
	The CaMKII inhibitor KN-62 has also been shown to facilitate DHPG-LTD and (S)-3,5-DHPG-induced LTD is thought to involve rapid protein synthesis which can be antagonized by the protein synthesis inhibitor anisomycin.
Application notes	(S)-3,5-DHPG is active <i>in vivo</i> and shows a variety of biological actions.
	Figure 1: Gamma-frequency oscillations recorded in CA3 region of rat <i>in vitro</i> hippocampal slice preparation.

Increase in gamma frequency oscillatory activity (dark blue) following (S)-3,5-DHPG application. Under control conditions there is no gamma activity (light blue).

Data kindly provided by Neurexpert

Solubility & Handling

Storage instructions
Solubility overview
Handling continued..

-20 °C (desiccate). Protect from light.

Soluble in water (50mM)

The compound is of high purity, however it is air sensitive and light promoted oxidation may cause slight discoloration of the compound. Product performance should not be affected, and this decolourisation can be ignored.

It has been shown that this compound rapidly decomposes when dissolved in alkaline solution. We therefore recommend that you store the solid material at -20 °C and protect from light. The compound should be stable for a least 6 months.

When possible, you should make up solutions and use immediately. If storage of solutions is required, you should aliquot out the solution into tightly sealed vials and store at -20 °C and store these for up to one month. Allow the product to equilibrate to RT for at least one hour before opening and using.

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

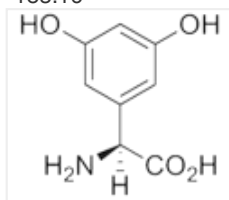
Important

Chemical Data

Chemical name
Molecular Weight
Chemical structure

(S)-3,5-Dihydroxyphenylglycine

183.16



Molecular Formula
CAS Number
PubChem identifier
SMILES
Source
InChi
InChiKey
MDL number
Appearance

C₈H₉NO₄

162870-29-3

443586

C1=C(C=C(C=C1O)O)[C@@H](C(=O)O)N

Synthetic

InChI=1S/C8H9NO4/c9-7(8(12)13)4-1-5(10)3-6(11)2-4/h1-3,7,10-11H,9H2,(H,12,13)/t7-/m0/s1

HOOWCUZPEFNHDT-ZETCQYMHSA-N

MFCD11044457

White solid

References

Behavioral and convulsant effects of the (S) enantiomer of the group I metabotropic glutamate receptor agonist 3,5-DHPG in mice.

Barton ME *et al* (2005) *Neuropharmacology* 48(6)

PubMedID [15829250](#)

Face-washing behavior induced by the group I metabotropic glutamate receptor agonist (S)-3,5-DHPG in mice is mediated by mGlu1 receptor.

Hikichi H *et al* (2008) *Eur J Pharmacol* 586(1-3)

PubMedID [18378225](#)

(S)-3,5-DHPG: a review.

Wiśniewski K *et al* (2002) *CNS Drug Rev* 8(1)

PubMedID [12070529](#)
