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DATASHEET

Salvinorin B (SALB)

Product overview

| | |
|--------------------------|---|
| Name | Salvinorin B (SALB) |
| Cat No | HB4887 |
| Description | Potent, selective KORD DREADD activator |
| Alternative names | SALB, Divinorin B |
| Biological action | Activator |
| Purity | >98% |
| Customer comments | <i>High quality with better price. I have compared SalB (Salvinorin B) from different producer, Hello Bio really provide a high quality compound with a cheaper price. Would definitely order again from here. Verified customer, Stony Brook University</i> |

Biological Data

| | |
|-------------------------------|--|
| Biological description | <p>Salvinorin B (SALB) is a pharmacologically inert ligand that potently and selectively activates the KORD (the κ-opioid designer receptor (DREADD)) ($EC_{50} = 11.8$ nM).</p> <p>Salvinorin B (SALB) is ~100-fold selective for the KORD DREADD over human κ opioid receptor and other targets and shows good CNS penetrability.</p> <p>Activation of KORD by Salvinorin B (SALB) induces neuronal inhibition and modifies behaviour in vivo.</p> <p>Salvinorin B (SALB) can be used in mice also expressing Clozapine N-Oxide (CNO) responsive DREADDs, to allow bi-directional manipulation of neural circuits.</p> <p>CNO dihydrochloride (water soluble), Clozapine N-oxide (CNO) freebase, Compound 21 and perlapine freebase also available.</p> |
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Solubility & Handling

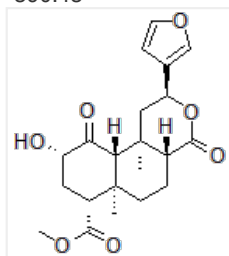
| | |
|-----------------------------|---|
| Storage instructions | -20°C |
| Solubility overview | Soluble in DMSO (20 mM) |
| Handling | <ul style="list-style-type: none">This compound is light sensitive; we therefore recommend protecting the solid and solutions from exposure to light.Salvinorin B (SalB) is unstable in solution and we recommend that solutions are stored at -20°C and used within 24 hours. |
| Important | This product is for RESEARCH USE ONLY and is not intended for therapeutic or diagnostic use. Not for human or veterinary use. |

Chemical Data

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|----------------------|--|
| Chemical name | (2S,4aR,6aR,7R,9S,10aS,10bR)-2-(3-Furanyl)dodecahydro-9-hydroxy-6a,10b-dimethyl-4,10-dioxo-2H-naphtho[2,1-c]pyran-7-carboxylic acid methyl ester |
|----------------------|--|

Molecular Weight
Chemical structure

390.43



Molecular Formula
CAS Number
PubChem identifier
SMILES

C₂₁H₂₆O₇
92545-30-7
11440685
C[C@@]12CC[C@H]3C(=O)O[C@@H](C[C@@]3([C@H]1C(=O)[C@H](C[C@H]2C(=O)OC)O)C)C4=COC=C4

Source

Extracted from salvia divinorum

InChi

InChI=1S/C21H26O7/c1-20-6-4-12-19(25)28-15(11-5-7-27-10-11)9-21(12,2)17(20)16(23)14(22)8-13(20)18(24)26-3/h5,7,10,12-15,17,22H,4,6,8-9H2,1-3H3/t12-,13-,14-,15-,17-,20-,21-/m0/s1

InChiKey

BLTMVAIOAAGYAR-CEFSSPBYSAN

MDL number

MFCD16036232

Appearance

Off-white solid

References

A New DREADD Facilitates the Multiplexed Chemogenetic Interrogation of Behavior.

Vardy et al (2015) Neuron. 86(4)

PubMedID [25937170](#)

DREADDS: Use and application in behavioral neuroscience.

Smith et al (2016) Behav Neurosci 130(2)

PubMedID [26913540](#)

Behavioral and Physiological Effects of a Novel Kappa-Opioid Receptor-Based DREADD in Rats.

Marchant et al (2016) Neuropsychopharmacology 41(2)

PubMedID [26019014](#)

Antinociceptive and hypothermic effects of Salvinorin A are abolished in a novel strain of kappa-opioid receptor-1 knockout mice.

Ansonoff MA et al (2006) J Pharmacol Exp Ther 318 (2):

PubMedID [16672569](#)

DREADDs: The Power of the Lock, the Weakness of the Key. Favoring the Pursuit of Specific Conditions Rather than Specific Ligands.

Goutaudier et al (2019) eNeuro 6

PubMedID [31562177](#)
