8-(4-Chlorophenylthio)-2'-O-methyladenosine 3',5'-cyclic

receptor. Activates Epac1 and Epac2, the guanine nucleotide-exchange factors (PFS) for the small GTPases Rap1 and Rap2. Does not activate protein knase A (PKA), rietiner *in vitro* nor *in vivo* and is therefore a valuable tool to specifically discriminate between PKA- and Epac-mediated effects. Since a free 2'-ribose hydroxyl group in cyclic AMP is essential for stimulation of PKA, the methylated structure of 8-CPT-2'-O-Me-cAMP is an extremely poor PKA activator. Its high lipophilicity allows for good membrane permeability in most biosystems and its increased resistance towards phosphodiesterases prevents rapid hydrolysis.

Citations: 3

View Online »

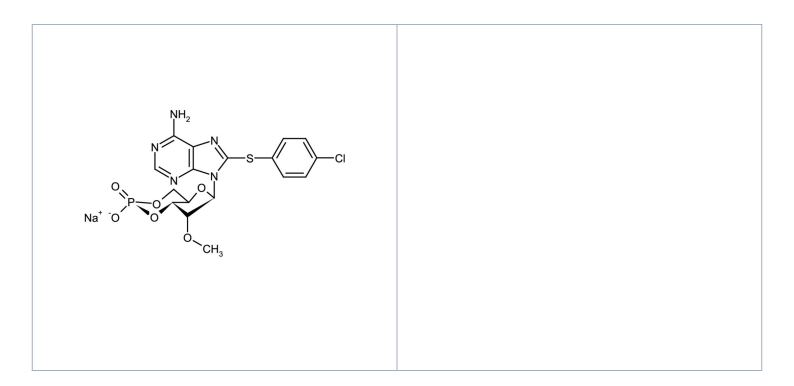
Ordering Information

Order Online »

ALX-480-093-M001	1mg
ALX-480-093-M005	5mg

Manuals, SDS & CofA

View Online »



Handling & Storage

Use/Stability As indicated on product label or CoA when stored as recommended.

Handling Protect from light. Hygroscopic.

Long Term Storage -20°C

Shipping Ambient Temperature

Regulatory Status RUO - Research Use Only

Product Details

Alternative Name 8-CPT-2'-O-Me-cAMP . Na

Appearance White to off-white crystalline powder.

CAS 510774-50-2

Couple Target CAMP receptor, GEF

Couple Type Activator

Formula $C_{17}H_{16}CIN_5O_6PS$. Na

Identity Determined by ESI-MS and UV.

MW 484.8 . 23.0

Purity ≥98% (HPLC)

Purity Detail Product is not sterile.

Solubility Soluble in water, aqueous buffers, DMSO, dimethyl formamide or methanol.

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