

Guanylyl cyclase inhibitor

ODQ is a potent and selective, direct-acting inhibitor of guanylate cyclase ($IC_{50}=20$ nM). In cerebellar slices ODQ reversibly inhibited cGMP generation in response to endogenous NO or exogenously added NO-donors. ODQ does not inhibit NO-mediated macrophage toxicity (a phenomenon unrelated to cGMP) and does not inhibit particulate guanylyl cyclase or adenylyl cyclase. ODQ is a unique and selective tool to elucidate the physiological importance of the NO-cGMP pathway.

Citations: 19

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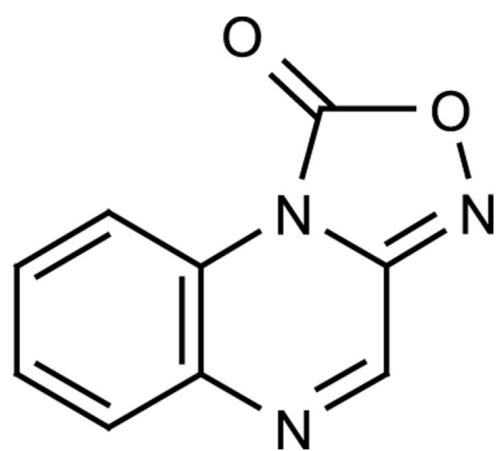
Ordering Information

ALX-270-034-M050

50mg

Manuals, SDS & CofA

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Handling & Storage

Use/Stability	As indicated on product label or CoA when stored as recommended. Solutions may decompose slowly.
Handling	Protect from light.
Short Term Storage	+4°C
Long Term Storage	-20°C
Shipping	Ambient Temperature

Regulatory Status

RUO - Research Use Only

Product Details

Appearance	Off-white to yellow powder.
CAS	41443-28-1
Couple Target	Guanlyl cyclase
Couple Type	Inhibitor
Formula	C ₉ H ₅ N ₃ O ₂
MW	187.2
Purity	≥98% (TLC)
Solubility	Soluble in DMSO (5mg/ml), ethyl acetate or 100% ethanol (1.2mg/ml); insoluble in water.

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ENZO LIFE SCIENCES,
INC.
Phone: 800.942.0430
[info-
usa@enzolifesciences.com](mailto:info-usa@enzolifesciences.com)

European Sales Office
ENZO LIFE SCIENCES
(ELS) AG
Phone: +41 61 926 8989
[info-
eu@enzolifesciences.com](mailto:info-eu@enzolifesciences.com)

Belgium, The Netherlands
& Luxembourg
Phone: +32 3 466 0420
[info-
fr@enzolifesciences.com](mailto:info-fr@enzolifesciences.com)

France
Phone: +33 472 440 655
[info-
fr@enzolifesciences.com](mailto:info-fr@enzolifesciences.com)

Germany
Phone: +49 7621 5500 526
[info-
de@enzolifesciences.com](mailto:info-de@enzolifesciences.com)

UK & Ireland
Phone (UK customers):
0845 601 1488
Phone: +44 1392 825900
[info-
uk@enzolifesciences.com](mailto:info-uk@enzolifesciences.com)