

Produktinformation



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Diagnostik & molekulare Diagnostik



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SZABO-SCANDIC HandelsgmbH

Quellenstraße 110, A-1100 Wien

T. +43(0)1 489 3961-0

F. +43(0)1 489 3961-7

mail@szabo-scandic.com

www.szabo-scandic.com

linkedin.com/company/szaboscandic in



PRODUCT INFORMATION



Iloperidone

Item No. 22957

CAS Registry No.: 133454-47-4

1-[4-[3-[4-(6-fluoro-1,2-benzisoxazol-Formal Name:

3-yl)-1-piperidinyl]propoxy]-3-

methoxyphenyl]-ethanone

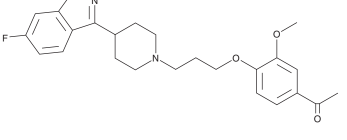
MF: $C_{24}H_{27}FN_2O_4$ FW: 426.5

Purity: ≥98%

 λ_{max} : 229, 275, 304 nm UV/Vis.: Supplied as: A crystalline solid

Storage: -20°C Stability: ≥2 years

Information represents the product specifications. Batch specific analytical results are provided on each certificate of analysis.



Laboratory Procedures

lloperidone is supplied as a crystalline solid. A stock solution may be made by dissolving the iloperidone in the solvent of choice. Iloperidone is soluble in organic solvents such as DMSO and dimethyl formamide, which should be purged with an inert gas. The solubility of iloperidone in these solvents is approximately 0.1 and 0.2 mg/ml, respectively.

Description

lloperidone is an atypical antipsychotic and adrenergic, dopamine, and serotonin (5-HT) receptor antagonist.¹ It binds to several receptors, including the α_1 -adrenergic receptor (α_1 -AR), α_2 -AR, and dopamine D₂ receptor (K_i s = 0.31, 3, and 3.3 nM, respectively), as well as the 5-HT_{1D}, 5-HT_{1D}, 5-HT_{2D}, and 5-HT_{2C} receptors (K_is = 33, 15, 0.2, and 14 nM, respectively) in radioligand binding assays using human post-mortem brain tissue.² Iloperidone also binds to human D_1 , D_3 , D_4 , D_5 , and rat 5-HT₂ receptors $(K_i s = 216, 7.1, 25, 319, and 3.1 nM, respectively, in CHO cells) and the histamine <math>H_1$ receptor $(K_i = 12.3 nM)$ in human post-mortem brain tissue).^{2,3} lloperidone (1-3 mg/kg) prevents the reduction in prepulse inhibition induced by apomorphine (Item No. 16094), phencyclidine (PCP), and cirazoline (Item No. 21791) in rats.¹ It also increases the time rats spend in the open arms of the elevated plus maze and the number of social interactions when administered at a dose of 0.5 mg/kg.4 Formulations containing iloperidone have been used in the treatment of schizophrenia.

References

- 1. Barr, A.M., Powell, S.B., Markou, A., et al. lloperidone reduces sensorimotor gating deficits in pharmacological models, but not a developmental model, of disrupted prepulse inhibition in rats. Neuropharmacology **51(3)**, 457-465 (2006).
- 2. Richelson, E. and Souder, T. Binding of antipsychotic drugs to human brain receptors focus on newer generation compounds. Life Sciences 68(1), 29-39 (2000).
- Kongsamut, S., Roehr, J.E., Cai, J., et al. lloperidone binding to human and rat dopamine and 5-HT receptors. Eur. J. Pharmacol. 317(2-3), 417-423 (1996).
- Szewczak, M.R., Corbett, R., Rush, D.K., et al. The pharmacological profile of iloperidone, a novel atypical antipsychotic agent. J. Pharmacol. Exp. Ther. 274(3), 1404-1413 (1995).

WARNING
THIS PRODUCT IS FOR RESEARCH ONLY - NOT FOR HUMAN OR VETERINARY DIAGNOSTIC OR THERAPEUTIC USE.

This material should be considered hazardous until further information becomes available. Do not ingest, inhale, get in eyes, on skin, or on clothing. Wash thoroughly after handling. Before use, the user must review the complete Safety Data Sheet, which has been sent via email to your institution.

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CAYMAN CHEMICAL

1180 EAST ELLSWORTH RD ANN ARBOR, MI 48108 · USA PHONE: [800] 364-9897

[734] 971-3335

FAX: [734] 971-3640 CUSTSERV@CAYMANCHEM.COM WWW.**CAYMANCHEM**.COM