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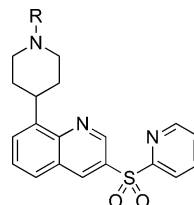
Quinoline Derivatives as 5-HT₆ Receptor PET Ligands

Gerard Rosse*

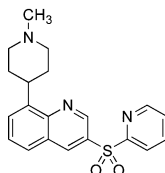
Structure Guided Chemistry, Dart Neuroscience LLC, 7473 Lusk Boulevard, San Diego, California 92121, United States, and Adjunct Associate Professor, Department of Pharmacology and Physiology, College of Medicine, Drexel University, New College Building, 245 North 15th Street, Philadelphia, Pennsylvania 19102, United States

Title: Quinoline Derivatives as 5-HT₆ Receptor PET Ligands
Patent/Patent Application Number: US-20130343993-A1 **Publication date:** December 26, 2013
Priority Application: US-20130343993-A1 **Priority date:** January 08, 2012
Inventors: Black, L. A.
Assignee Company: AbbVie Inc. USA
Disease Area: Alzheimer's Disease, deficits in memory, cognition, and learning **Biological Target:** 5-HT₆ receptor
Summary: This application claims a series of quinolines for treating or preventing a condition or disorder related to memory deficits such as Parkinson's disease, Alzheimer's disease, mild cognitive impairment, depression, and anxiety. The invention claims also radiolabeled quinolines useful as diagnostic tools as 5-HT₆ receptor PET ligands.

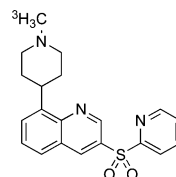
Important Compound Classes:



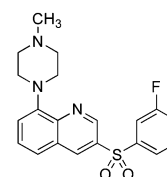
Key Structures:



Example 1



Example 2



GSK-215083

Biological Assay:

Compounds were evaluated in 5-HT₆, 5-HT_{2A}, and 5-HT_{2B} receptor binding assays and against a panel of 78 receptors/drug targets.

Biological Data:

Compounds binding affinities

	Example 1	GSK-215083
Human 5-HT ₆ K _i (nM)	0.22	0.34
Human 5-HT _{2A} K _i (nM)	123 (559x)	0.39 (1.16x)
Human 5-HT _{2B} K _i (nM)	144 (654x)	

Rat PK (0.05 mg/kg, iv) for Example 1

Minutes after dose	Plasma conc. (ng/mL)	Free brain conc. (ng/g)	Free B/P ratio
3	104.5	21.5	0.21
5	83.5	23.1	0.28

Brain Distribution for Tritium Labeled Example 2

Minutes after dose	Stratium/ Cer.	Hippocampus/ Cer.	Cortex/Cer.
5	0.81	1.06	0.9
40	1.31	1.07	1.20

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Notes

The authors declare no competing financial interest.