NEW USE AND NOVEL N-AZABICYCLO-AMIDE DERIVATIVES

Abstract: This invention relates to new use of a compound of general formula (I) wherein: A represents (II), (III), (IV), (V), (VI); D represents oxygen, or sulfur; R¹ represents hydrogen or methyl; R² represents hydrogen, or C₁-C₄ alkyl; and R³ represents (A), (B), (C), or an enantiomer thereof, and pharmaceutically acceptable salts thereof, processes for preparing them, pharmaceutical compositions containing them and their use in therapy, especially in the treatment of prophylaxis of psychotic disorders and intellectual impairment disorders.
NEW USE AND NOVEL N-azaCyclo-AMIDE DERIVATIVES

Technical Field

This invention relates to new medical use of quinuclidine acrylamides or
5 pharmacologically acceptable salts thereof, processes for preparing them and pharmaceutical
compositions containing them. The present invention also relates to certain novel quinuclidine
acrylamides or pharmacologically acceptable salts thereof, processes for preparing them and
pharmaceutical compositions containing them. In particular the invention relates to the use of
quinuclidine acrylamides for the preparation of medicaments for the treatment or prophylaxis
10 of psychotic disorders or intellectual impairment disorders, as well as in the treatment and/or
prophylaxis of human diseases or conditions in which activation of the α7 nicotinic receptor
is beneficial.

Background of the Invention

The use of compounds which bind nicotinic acetylcholine receptors in the treatment of
15 a range of disorders involving reduced cholinergic function such as Alzheimer’s disease,
cognitive or attention disorders, anxiety, depression, smoking cessation, neuroprotection,
schizophrenia, analgesia, Tourette’s syndrome, and Parkinson’s disease has been discussed in
McDonald et al. (1995) “Nicotinic Acetylcholine Receptors: Molecular Biology, Chemistry
and Pharmacology”, Chapter 5 in Annual Reports in Medicinal Chemistry, vol. 30, pp. 41-50,
20 Academic Press Inc., San Diego, CA; and in Williams et al. (1994) “Neuronal Nicotinic

The use of certain quinuclidine acrylamide derivatives, as to which the present
invention has found a new pharmacological use, is known from EP 581165-A2 to have effect
25 as antitussives. The antitussive activity of the compounds was described as being "without
effects on the central nervous system", and the use of the compounds for the treatment of
diseases involving the central nervous system was not suggested.

Disclosure of the Invention

According to the present invention it has been found that compounds of the general
formula I:
wherein:

A represents:

II  III  IV  V  VI

D represents oxygen, or sulfur;
R¹ represents hydrogen or methyl;
R² represents hydrogen, or C₁–C₄ alkyl;
R³ represents:

R⁴, R⁵, and R⁶ are independently hydrogen, halogen, C₁–C₄ alkyl, C₂–C₄ alkenyl, C₂–
C₄ alkynyl, –CO₂R⁷, –CN, –CF₃, or Ar, provided that at least one of R⁴ and R⁵ represents Ar;
Ar represents a 5- or 6-membered aromatic or heteroaromatic ring containing zero to three
nitrogen atoms, zero or one oxygen atom, and zero or one sulfur atom, or an 8-, 9- or 10-
membered fused aromatic or heteroaromatic ring system containing zero to four nitrogen
atoms, zero to one oxygen atoms, and zero to one sulfur atoms which may optionally be
substituted with one or more substituents selected from the following: hydrogen, halogen, C₁–
C₄ alkyl, C₂–C₄ alkenyl, C₂–C₄ alkynyl, aryl, heteroaryl, –CO₂R⁷, –CN, –NO₂, –NR³R⁹, –
CF₃, –OR¹⁰;
R⁸, R⁹, and R¹⁰ are independently hydrogen, C₁–C₄ alkyl, aryl, heteroaryl, -C(O)R¹¹, -C(O)NHR¹², -C(O)R¹³, -SO₂R¹⁴ or R⁸ and R⁹ may together be (CH₂)₉Q(CH₂)₉ where Q is O, S, NR¹⁵, or a bond;

j is 2 to 4;
k is 0 to 2;

R⁷, R¹⁰, R¹¹, R¹², R¹³, R¹⁴, and R¹⁵, are independently C₁–C₄ alkyl, aryl, or heteroaryl; or an enantiomer thereof, and pharmaceutically acceptable salts thereof, are useful for the preparation of a medicament for the treatment or prophylaxis of psychotic disorders or intellectual impairment disorders.

Examples of such conditions, diseases or disorders are Alzheimer’s disease, learning deficit, cognition deficit, attention deficit, memory loss, Attention Deficit Hyperactivity Disorder, anxiety, schizophrenia, mania or manic depression, Lewy Body Dementia, Parkinson’s disease, Huntington’s disease, Tourette’s syndrome, neurodegenerative disorders in which there is loss of cholinergic synapse, jetlag, cessation of smoking, nicotinic addiction including that resulting from exposure to products containing nicotine, pain, and for ulcerative colitis.

Unless otherwise indicated, the C₁–C₄ alkyl groups referred to herein, e.g., methyl, ethyl, n-propyl, n-butyl, i-propyl, i-butyl, t-butyl, s-butyl, whether alone or part of another group, may be straight-chained or branched, and the C₃–C₄ alkyl groups may also be cyclic, e.g., cyclopropyl, cyclobutyl.

Unless otherwise indicated, aryl refers to a phenyl ring which may optionally be substituted with one to three of the following substituents chosen from among the following: halogen, C₁–C₄ alkyl, C₂–C₄ alkenyl, C₂–C₄ alkynyl, -CO₂R⁷, -CN, -NO₂, -NR₈R⁹, -CF₃, -OR¹⁰.

Unless otherwise indicated, heteroaryl refers to a 5- or 6-membered aromatic or heteroaromatic ring containing zero to three nitrogen atoms, zero or one oxygen atom, and zero or one sulfur atom, provided that the ring contains at least one nitrogen, oxygen, or sulfur atom, which may optionally be substituted with one or more substituents chosen from among the following: halogen, C₁–C₄ alkyl, C₂–C₄ alkenyl, C₂–C₄ alkynyl, -CO₂R⁷, -CN, -NO₂, -NR₈R⁹, -CF₃, -OR¹⁰.

Unless otherwise indicated, halogen refers to fluorine, chlorine, bromine, or iodine.
Pharmaceutically acceptable derivatives include solvates and salts. For example, the compounds of formula I can form acid addition salts with acids, such as the conventional pharmaceutically acceptable acids, for example, maleic, hydrochloric, hydrobromic, phosphoric, acetic, fumaric, salicylic, citric, lactic, mandelic, tartaric and methanesulfonic acids.

Preferred embodiments of the invention include compounds of formula I, wherein A represents:

![Diagram](image)

or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

Preferred embodiments of the invention also include compounds of formula I, wherein D represents oxygen; or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

Preferred embodiments of the invention also include compounds of formula I, wherein R² represents hydrogen; or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

Preferred embodiments of the invention also include compounds of formula I, wherein R³ represents:

![Diagram](image)

or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

Preferred embodiments of the invention also include compounds of formula I, wherein R³ represents:
when $R^4$ represents Ar; $R^5$ represents hydrogen, or $C_1$-$C_4$ alkyl; $R^6$ represents hydrogen, or halogen; or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

Preferred embodiments of the invention also include compounds of formula I, wherein

A represents:

![Diagram II](image-url)

$D$ represents oxygen;
$R^2$ represents hydrogen;
$R^3$ represents:

![Diagram Ar](image-url)

or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

Preferred embodiments of the invention also include compounds of formula I, wherein

A represents:

![Diagram II](image-url)

$D$ represents oxygen;
$R^2$ represents hydrogen;
$R^3$ represents:

![Diagram R6](image-url)

$R^4$ represents Ar;
R⁵ represents hydrogen, or C₁-C₄ alkyl;
R⁶ represents hydrogen, or halogen;
or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

Preferred embodiments of the invention also include compounds of formula I, wherein

Ar represents a 5- or 6-membered aromatic or heteroaromatic ring containing zero to three nitrogen atoms, zero or one oxygen atom, and zero or one sulfur atom, or an 8-, 9- or 10-membered fused aromatic or heteroaromatic ring system containing zero to four nitrogen atoms, zero to one oxygen atoms, and zero to one sulfur atoms which may optionally be substituted with one or more substituents selected from the following: hydrogen, halogen, C₁–C₄ alkyl, C₂–C₄ alkenyl, C₂–C₄ alkynyl, aryl, heteroaryl, −CO₂R⁷, −CN, −NO₂, −NR⁸R⁹, −CF₃, −OR¹⁰; or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

Preferred embodiments of the invention also include compounds of formula I, wherein

A represents:

D represents oxygen;
R¹ and R² are hydrogen;
R³ represents:

or R³ represents:

in which:
R⁴ represents Ar;
R⁵ represents hydrogen or C₁–C₄ alkyl;
R⁶ represents hydrogen or halogen;
Ar represents a 5- or 6-membered aromatic or heteroaromatic ring containing zero to
three nitrogen atoms, zero or one oxygen atom, and zero or one sulfur atom, including phenyl,
2-, 3-, or 4-pyridyl, 2- or 3-furanyl, and 2- or 3-thienyl, any of which may optionally be
substituted with one or more substituents chosen from among the following: hydrogen,
halogen, C₁–C₄ alkyl, C₂–C₄ alkenyl, C₂–C₄ alkynyl, –CO₂R₇, –CN, –NO₂, –NR₈R₉, –CF₃, –
OR¹⁰; or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

Preferred compounds of formula I include the following:
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(3-phenylpropynamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl) [E-3-(2-nitrophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl) [E-3-(2-nitrophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-aminophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[Z-3-(2-methoxyphenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)-N-methyl-(E-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-2-phenylcyclopropane-1-carboxamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-2-fluoro-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-formamidophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-nitrophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-aminophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-formamidophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-3-methyl-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-N-methylaminophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-N,N-dimethylaminophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-methyl-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2,3-diphenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-methoxyphenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2-methyl-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-methylphenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-methoxyphenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-fluorophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-fluorophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-chlorophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-chlorophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-chlorophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3,4-dichlorophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-bromophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-bromophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-iodophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-iodophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-trifluoromethylphenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-trifluoromethylphenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-furyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-furyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-pyridyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-pyridyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-pyridyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-thienyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-thienyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-nitro-2-furyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-methoxy-3-pyridyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-hydroxy-3-pyridyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-imidazolyl)propenamide];
N-(endo-8-Aza-8-methylbicyclo[3.2.1]oct-3-yl)(E-3-phenylpropenamide);
N-(exo-8-Aza-8-methylbicyclo[3.2.1]oct-3-yl)(E-3-phenylpropenamide);

or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

Particularly preferred compounds of formula I include the following:
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-phenylpropenamide);
(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-phenylpropenamide);
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-phenylpropenamide);
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(3-phenylpropynamide);
(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(3-phenylpropamide);
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-nitrophenyl)propenamide);
(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl) [E-3-(2-nitrophenyl)propenamide];
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl) [E-3-(2-nitrophenyl)propenamide];
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-aminophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-aminophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)][Z-3-(2-methoxyphenyl)propenamide];
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)-N-methyl-(E-3-phenylpropenamide);
(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-2-phenylcyclopropane-1-carboxamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2-phenylcyclopropane-1-carboxamide);
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[Z-2-fluoro-3-phenylpropenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-formamidophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)][E-3-(4-nitrophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-aminophenyl)propenamide);
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-formamidophenyl)propenamide);
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-3-methyl-3-phenylpropenamide);
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-N-methylaminophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-N,N-dimethylaminophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-3-phenylpropenamide);
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-methyl-3-phenylpropenamide);
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2,3-diphenylpropenamide);
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-methoxyphenyl)propenamide);
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-(2-chlorophenyl)propenamide];
(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-(2-chlorophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-chlorophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-chlorophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3,4-dichlorophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-bromophenyl)propenamide]
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-bromophenyl)propenamide]
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-iodophenyl)propenamide];
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-iodophenyl)propenamide];
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-trifluoromethylphenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-trifluoromethylphenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-furyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-furyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-thienyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-thienyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-nitro-2-furyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-methoxy-3-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-hydroxy-3-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-imidazolyl)propenamide];
N-(endo-8-Aza-8-methylbicyclo[3.2.1]oct-3-yl)(E-3-phenylpropenamide);
N-(exo-8-Aza-8-methylbicyclo[3.2.1]oct-3-yl)(E-3-phenylpropenamide),
or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

In another aspect of the present invention, there is provided novel compounds
according to formula I, with the additional proviso that Ar does not represent 2-, 3-, 4-pyridyl,
unsubstituted phenyl or phenyl substituted with one or more substituents selected from C1-C4
alkyl, C1-C4 alkoxy, halogen, phenoxy, hydroxy, OCOR11, NH2, NHCOR11, and nitro, when:
A represents:
R³ represents:

D represents oxygen;
R represents methyl;
R² and R⁵ both represent hydrogen;
R⁶ represents any of hydrogen, C₁–C₄ alkyl, phenyl, or cyano;

or an enantiomer thereof, and pharmaceutically acceptable salts thereof, which are potent ligands for nicotinic acetylcholine receptors.

In a preferred embodiment of this aspect of the invention, there is provided a compound according to formula I, as defined above, wherein A represents:

or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

In another preferred embodiment of this aspect of the invention, there is provided a compound according to formula I, as defined above, wherein A represents:
and $R^6$ represents any of 2-furyl; 3-furyl; 2-thienyl; 3-thienyl; or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

Among these compounds, the following compounds are preferred:

$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-trifluoromethylphenyl)propenamide];
$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-furyl)propenamide];
$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-furyl)propenamide];
$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-thienyl)propenamide];
$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-thienyl)propenamide];
$N$-(1-Aza-3-cyanobicyclo[2.2.2]oct-3-yl)[E-3-phenylpropenamide];
$N$-(1-Aza-3-methylbicyclo[2.2.2]oct-3-yl)[E-3-phenylpropenamide];
$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-nitro-2-furyl)propenamide];
$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-methoxy-3-pyridyl)propenamide];
$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-hydroxy-3-pyridyl)propenamide];
$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-imidazolyl)propenamide];
or an enantiomer and/or pharmaceutically acceptable salts thereof.

Among these compounds, the following compounds are particularly preferred:

$(R)$-$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-trifluoromethylphenyl)propenamide];
$(R)$-$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-furyl)propenamide];
$(R)$-$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-furyl)propenamide];
$(R)$-$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-thienyl)propenamide];
$(R)$-$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-thienyl)propenamide];
$N$-(1-Aza-3-cyanobicyclo[2.2.2]oct-3-yl)[E-3-phenylpropenamide];
$N$-(1-Aza-3-methylbicyclo[2.2.2]oct-3-yl)[E-3-phenylpropenamide];
$(R)$-$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-nitro-2-furyl)propenamide];
$(R)$-$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-methoxy-3-pyridyl)propenamide];
$N$-(1-Aza-3-methylbicyclo[2.2.2]oct-3-yl)[E-3-phenylpropenamide];
$(R)$-$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-nitro-2-furyl)propenamide];
$(R)$-$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-methoxy-3-pyridyl)propenamide];
$(R)$-$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-hydroxy-3-pyridyl)propenamide];
$(R)$-$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-imidazolyl)propenamide].

Pharmaceutically acceptable derivatives include solvates and salts. For example, the compounds of formula I can form acid addition salts with acids, such as the conventional pharmaceutically acceptable acids, for example, maleic, hydrochloric, hydrobromic, phosphoric, acetic, fumaric, salicylic, citric, lactic, mandelic, tartaric and methanesulfonic acids.
The compounds of the invention have the advantage that they may be less toxic, be more efficacious, be longer acting, have a broader range of activity, be more potent, produce fewer side effects, are more easily absorbed or have other useful pharmacological properties.

**Methods of Preparation**

In the reaction schemes and text that follow, A, R<sup>1</sup>, R<sup>2</sup>, and R<sup>3</sup>, unless otherwise indicated, are as defined above for formula I. The compounds of formula I may be prepared according to the methods outlined in Scheme 1.

![Diagram of Scheme 1](image)

**Scheme 1.**

Compounds of formula I wherein D represents oxygen may be prepared from compounds of formula VII by reaction with a compound of formula VIII, wherein Y represents a suitable leaving group, using a suitable acylation procedure. Suitable leaving groups Y include: OH, halogen, OAlkyl, OAryl, OCOAlkyl, OCOAryl, azide. A suitable acylation procedure involves treatment of a compound of formula I with a compound of formula VIII at 0-120°C in a suitable solvent. The presence of a base, or, when Y=OH, a coupling agent, may also be necessary for the reaction to occur. Suitable bases for the reaction include: 4-(N,N-dimethylamino)pyridine, pyridine, triethylamine, N,N-diisopropylethylamine. The preferred base is N,N-diisopropylethylamine. Suitable coupling agents when Y=OH include: carbodiimides, for example 1,3-dicyclohexylcarbodiimide or 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride; phosphonium reagents, for example benzotriazol-1-yloxytris(dimethylamino)phosphonium hexafluorophosphate or benzotriazol-1-yloxytripyrrolidinophosphonium hexafluorophosphate; and uronium reagents,
for example O-benzotriazol-1-yl-N,N,N',N'-tetramethyluronium tetrafluoroborate. The preferred coupling agent is O-benzotriazol-1-yl-N,N,N',N'-tetramethyluronium tetrafluoroborate. Suitable solvents for the reaction include N,N-dimethylformamide, dimethylsulfoxide, tetrahydrofuran, or chloroform. The preferred solvent is N,N-dimethylformamide. The reaction is preferably performed at a temperature of 0-50 °C, and most preferably at a temperature of 20-30 °C.

Compounds of formula I wherein D represents oxygen and R² represents hydrogen may alternatively be prepared from compounds of formula IX by treatment with a nitrile of formula X in a suitable solvent in the presence of a suitable acid. Suitable acids include sulfuric acid, and suitable solvents include acetic acid. The reaction is carried out at a temperature of 0-50 °C, and preferably at a temperature of 0-25 °C.

Compounds of formula I in which D represents sulfur may be prepared from compounds of formula I in which D represents oxygen by reaction with a suitable sulfide in a suitable solvent. The preferred sulfides are phosphorus sulfides, in particular 4-methoxyphenylthionophosphine sulfide dimer ("Lawesson's Reagent"), and diphosphorus pentasulfide. Suitable solvents for the reaction include aryl hydrocarbon solvents, for example toluene or xylene. The reaction is performed at a temperature of 0-200 °C, and preferably at a temperature of 50-180 °C.

Compounds of formula VII in which R² represents an alkyl group may be prepared from compounds of formula VII in which R² represents hydrogen by a suitable alkylation procedure. Typical alkylation procedures include treatment with an appropriate alkyl halide or sulfonate ester and base, for example sodium hydride, in a suitable solvent, for example DMF, or reductive alkylation using the appropriate aldehyde or ketone together with a suitable reducing agent in an inert solvent. The preferred method is reductive alkylation. Suitable reducing agents include sodium borohydride and sodium cyanoborohydride. The preferred reducing agent is sodium borohydride. Suitable inert solvents include water, methanol or ethanol. The preferred solvent is methanol. The reaction is usually conducted at a temperature of 0-100 °C, preferably at 20 °C - 65 °C.

Compounds of formula VII are commercially available or may be prepared by methods known to one skilled in the art. For example, certain compounds of formula VII may be prepared from compounds of formula IX via a Ritter reaction with a suitable nitrile, followed by hydrolysis of the resulting amide.
Compounds of formula VIII and X are commercially available or may be prepared by methods known to one skilled in the art. See, for example, the methods cited in "Comprehensive Organic Transformations" by R.C. Larock (VCH Publishers, 1989), pages 819-995, and the general references cited on page 823 of the same text.

It will be appreciated by one skilled in the art that certain optional aromatic substituents in the compounds of the invention may be introduced by employing aromatic substitution reactions, or functional group transformations to modify an existing substituent, or a combination thereof. Such reactions may be effected either prior to or immediately following the processes mentioned above, and are included as part of the process aspect of the invention. The reagents and reaction conditions for such procedures are known in the art.

Specific examples of procedures which may be employed include, but are not limited to, electrophilic functionalisation of an aromatic ring, for example via nitration, halogenation, or acylation; transformation of a nitro group to an amino group, for example via reduction, such as by catalytic hydrogenation; acylation, alkylation or sulfonylation of an amino or hydroxyl group; replacement of an amino group by another functional group via conversion to an intermediate diazonium salt followed by nucleophilic or free radical substitution of the diazonium salt; or replacement of a halogen by another functional group for example via nucleophilic or catalysed substitution reactions.

Where necessary, hydroxy, amino, or other reactive groups may be protected using a protecting group as described in the standard text "Protecting groups in Organic Synthesis", 3rd Edition (1999) by Greene and Wuts.

The above described reactions, unless otherwise noted, are usually conducted at a pressure of about one to about three atmospheres, preferably at ambient pressure (about one atmosphere). Unless otherwise stated, the above described reactions are conducted under an inert atmosphere, preferably under a nitrogen atmosphere.

The compounds of the invention and intermediates may be isolated from their reaction mixtures by standard techniques.

Acid addition salts of the compounds of formula I which may be mentioned include salts of mineral acids, for example the hydrochloride and hydrobromide salts; and salts formed with organic acids such as formate, acetate, maleate, benzoate, tartrate, and fumarate salts.

Acid addition salts of compounds of formula I may be formed by reacting the free base or a salt, enantiomer or protected derivative thereof, with one or more equivalents of the appropriate acid. The reaction may be carried out in a solvent or medium in which the salt is
insoluble or in a solvent in which the salt is soluble, e.g., water, dioxane, ethanol, tetrahydrofuran or diethyl ether, or a mixture of solvents, which may be removed in vacuum or by freeze drying. The reaction may be a metathetical process or it may be carried out on an ion exchange resin.

The compounds of formula I exist in tautomeric or enantiomeric forms, all of which are included within the scope of the invention. The various optical isomers may be isolated by separation of a racemic mixture of the compounds using conventional techniques, e.g. fractional crystallization, or chiral HPLC. Alternatively the individual enantiomers may be made by reaction of the appropriate optically active starting materials under reaction conditions which will not cause racemization.

**Pharmaceutical Compositions**

A further aspect of the invention relates to a pharmaceutical composition for treating or preventing a condition or disorder as exemplified below arising from dysfunction of nicotinic acetylcholine receptor neurotransmission in a mammal, preferably a human, comprising an amount of a compound of formula I, an enantiomer thereof or a pharmaceutically acceptable salt thereof, effective in treating or preventing such disorder or condition and an inert pharmaceutically acceptable diluent or carrier.

A condition or disorder arising from dysfunction of nicotinic acetylcholine receptor neurotransmission in a mammal, preferably a human, may be psychotic disorders or intellectual impairment disorders or diseases or conditions in which activation of the α7 nicotinic receptor is beneficial. Examples of such conditions, diseases or disorders include Alzheimer’s disease, learning deficit, cognition deficit, attention deficit, memory loss, Attention Deficit Hyperactivity Disorder, anxiety, schizophrenia, mania or manic depression, Lewy Body Dementia, Parkinson’s disease, Huntington’s disease, Tourette’s syndrome, neurodegenerative disorders in which there is loss of cholinergic synapse, jetlag, cessation of smoking, nicotine addiction including that resulting from exposure to products containing nicotine, pain, or ulcerative colitis.

For the above-mentioned uses the dosage administered will, of course, vary with the compound employed, the mode of administration and the treatment desired. However, in general, satisfactory results are obtained when the compounds of the invention are administered at a daily dosage of from about 0.1 mg to about 20 mg per kg of animal body weight, preferably given in divided doses 1 to 4 times a day or in sustained release form. For
man, the total daily dose is in the range of from 5 mg to 1,400 mg, more preferably from 10 mg to 100 mg, and unit dosage forms suitable for oral administration comprise from 2 mg to 1,400 mg of the compound admixed with a solid or liquid pharmaceutical carrier or diluent.

The compounds of formula I, or an enantiomer thereof, and pharmaceutically acceptable salts thereof, may be used on their own or in the form of appropriate medicinal preparations for enteral or parenteral administration. According to a further aspect of the invention, there is provided a pharmaceutical composition including preferably less than 80% and more preferably less than 50% by weight of a compound of the invention in admixture with an inert pharmaceutically acceptable diluent or carrier.

Examples of diluents and carriers are:
- for tablets and dragees: lactose, starch, talc, stearic acid; for capsules: tartaric acid or lactose;
- for injectable solutions: water, alcohols, glycerin, vegetable oils; for suppositories: natural or hardened oils or waxes.

There is also provided a process for the preparation of such a pharmaceutical composition, which comprises mixing the ingredients.

Utility

One aspect of the invention is the use of a compound according to the invention, an enantiomer thereof or a pharmaceutically acceptable salt thereof, in the manufacture of a medicament for the treatment or prophylaxis of one of the below mentioned diseases or conditions: and a method of treatment or prophylaxis of one of the above mentioned diseases or conditions, which comprises administering a therapeutically effective amount of a compound according to the invention, or an enantiomer thereof or a pharmaceutically acceptable salt thereof, to a patient.

Compounds to be used according to the invention are agonists of nicotinic acetylcholine receptors. While not being limited by theory, it is believed that agonists of the $\alpha_7$ nAChR (nicotinic acetylcholine receptor) subtype should be useful in the treatment or prophylaxis of psychotic disorders and intellectual impairment disorders, and have advantages over compounds which are or are also agonists of the $\alpha_4$ nAChR subtype. Therefore, compounds which are selective for the $\alpha_7$ nAChR subtype are preferred. The use of compounds of the invention is indicated as pharmaceuticals, in particular in the treatment or prophylaxis of psychotic disorders and intellectual impairment disorders. Examples of
psychotic disorders include schizophrenia, mania or manic depression, and anxiety. Examples of intellectual impairment disorders include Alzheimer’s disease, Lewy Body Dementia, learning deficit, cognition deficit, attention deficit, memory loss, and Attention Deficit Hyperactivity Disorder. The compounds of the invention may also be useful as analgesics in the treatment of pain (including chronic pain) and in the treatment or prophylaxis of Parkinson’s disease, Huntington’s disease, Tourette’s syndrome, and neurodegenerative disorders in which there is loss of cholinergic synapses. The compounds may further be indicated for the treatment or prophylaxis of jetlag, for use in inducing the cessation of smoking, and for the treatment or prophylaxis of nicotine addiction (including that resulting from exposure to products containing nicotine).

It is also believed that compounds according to the invention are useful in the treatment and prophylaxis of ulcerative colitis.

**Pharmacology**

The pharmacological activity of the compounds of the invention may be measured in the tests set out below:

**Test A - Assay for affinity at \( \alpha_7 \) nAChR subtype**

\( ^{125} \text{I}-\alpha\)-Bungarotoxin (BTX) binding to rat hippocampal membranes. Rat hippocampi were homogenized in 20 volumes of cold homogenization buffer (HB: concentrations of constituents (mM): tris(hydroxymethyl)aminomethane 50; MgCl\(_2\) 1; NaCl 120; KCl 5; pH 7.4). The homogenate was centrifuged for 5 minutes at 1000 g, the supernatant was saved and the pellet re-extracted. The pooled supernatants were centrifuged for 20 minutes at 12000 g, washed, and resuspended in HB. Membranes (30–80 \( \mu \)g) were incubated with 5 nM \( ^{125} \text{I}\)-BTX, 1 mg/mL BSA (bovine serum albumin), test drug, and either 2 mM CaCl\(_2\) or 0.5 mM EGTA [ethylene glycol-bis(\( \beta \)-aminoethylether)] for 2 hours at 21 °C, and then filtered and washed 4 times over Whatman glass fibre filters (thickness C) using a Brandel cell harvester. Pretreating the filters for 3 hours with 1% (BSA/0.01% PEI (polyethyleneimine) in water was critical for low filter blanks (0.07% of total counts per minute). Nonspecific binding was described by 100 \( \mu \)M (−)-nicotine, and specific binding was typically 75%.

**Test B - Assay for affinity to the \( \alpha_4 \) nAChR subtype**

\( ^{3} \text{H}(\text{−})\)-nicotine binding. Using a procedure modified from Martino-Barrows and Kellar (Mol Pharm (1987) 31:169-174), rat brain (cortex and hippocampus) was homogenized as in the \( ^{125} \text{I}\)-BTX binding assay, centrifuged for 20 minutes at 12,000 x g, washed twice, and
then resuspended in HB containing 100 μM diisopropyl fluorophosphate. After 20 minutes at 4°C, membranes (approximately 0.5 mg) were incubated with 3 nM [3H]-(−)-nicotine, test drug, 1 μM atropine, and either 2 mM CaCl₂ or 0.5 mM EGTA for 1 hour at 4°C, and then filtered over Whatman glass fibre filters (thickness C) (pretreated for 1 hour with 0.5% PEI) using a Brandel cell harvester. Nonspecific binding was described by 100 μM carbachol, and specific binding was typically 84%.

**Binding data analysis for Tests A and B**

IC₅₀ values and pseudo Hill coefficients (nₜ) were calculated using the non-linear curve fitting program ALLFIT (DeLean A, Munson P J and Rodbard D (1977) Am. J. Physiol., 235:E97-E102). Saturation curves were fitted to a one site model, using the non-linear regression program ENZFITTER (Leatherbarrow, R.J. (1987)), yielding Kᵋ values of 1.67 and 1.70 nM for the ¹²⁵I-α-BTX and [³H]-(−)-nicotine ligands respectively. Kᵋ values were estimated using the general Cheng-Prusoff equation:

$$Kᵋ = \frac{[IC₅₀]}{(2+([ligand]/[K_D])^n)^{1/n} - 1}$$

where a value of n=1 was used whenever nₕ<1.5 and a value of n=2 was used when nₕ≥1.5.

Samples were assayed in triplicate and were typically ±5%. Kᵋ values were determined using 6 or more drug concentrations. The compounds of the invention are compounds with binding affinities (Kᵋ) of less than 10 μM in either Test A or Test B, indicating that they are expected to have useful therapeutic activity.

The use of compounds of the invention have the advantage that they may be less toxic, be more efficacious, be longer acting, have a broader range of activity, be more potent, produce fewer side effects, are more easily absorbed or have other useful pharmacological properties.

**General Experimental Procedures**

Commercial reagents were used without further purification. Mass spectra were recorded using either a Hewlett Packard 5988A or a MicroMass Quattro-1 Mass Spectrometer and are reported as m/z for the parent molecular ion. Room temperature refers to 20–25°C.

**Examples**

The following examples are preferred non-limiting examples embodying preferred aspects of the invention.
Example 1

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-phenylpropenamide)

(R)-1-Azabicyclo[2.2.2]oct-3-ylamine dihydrochloride (7.5 g, 0.038 moles) and E-3-phenylpropenoic acid (5.6 g, 0.038 moles) were combined in anhydrous N,N-dimethylformamide (570 mL) under nitrogen atmosphere. The resulting mixture was stirred while adding 1-hydroxybenzotriazole monohydrate (5.8 g, 0.0378 moles) O-benzotriazol-1-yl-N,N,N′,N′-tetramethyluronium tetrafluoroborate (12.1 g, 0.038 moles) and N,N-diisopropylethylamine (26.4 mL) were added. The solution was stirred at ambient temperature for 6 h and then placed in a freezer overnight. The solution was concentrated in vacuo and the residue was taken up in chloroform (360 mL) and washed with aqueous sodium hydroxide (1 M, 360 mL). The basic layer was extracted twice more with chloroform and the organic layers were combined, dried (MgSO₄), and concentrated in vacuo. The compound was then purified on a flash silica gel column with a 5-20% methanol/chloroform/ammonium hydroxide gradient. The hydrochloride salt was prepared from isopropanol and ethyl acetate, giving 9.5 g of slightly off-white powder; MS (ES⁺) 257 (MH⁺).

Example 2

(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-phenylpropenamide)

Prepared as free base by a method analogous to that described in Example 1 from (S)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-phenylpropenoic acid; MS (ES⁺) 257 (MH⁺).

Example 3

(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-phenylpropenamide)

Prepared as free base by a method analogous to that described in Example 1 from (RS)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-phenylpropenoic acid; MS (ES⁺) 257 (MH⁺).

Example 4

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(3-phenylpropynamide)

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and 3-phenylpropynoic acid; MS (ES⁺) 255 (MH⁺).
Example 5

(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(3-phenylpropynamide)

Prepared as a free base by a method analogous to that described in Example 1 from (S)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and 3-phenylpropynoic acid; MS (ES⁺) 255 (MH⁺).

Example 6

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-nitrophenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(2-nitrophenyl)propenoic acid; MS (ES⁺) 302 (MH⁺).

Example 7

(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl) [E-3-(2-nitrophenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (S)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(2-nitrophenyl)propenoic acid as a solid; MS (ES⁺) 302 (MH⁺).

Example 8

(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl) [E-3-(2-nitrophenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (RS)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(2-nitrophenyl)propenoic acid as a solid; MS (ES⁺) 302 (MH⁺).

Example 9

(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-aminophenyl)propenamide]

(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-nitrophenyl)propenamide] (0.42 g, 0.014 moles) prepared was suspended in acetone (4.2 mL) and ammonium chloride (0.15 g, 0.028 moles) in water (2.1 mL) was added. The clear solution was brought to reflux using an oil bath, and then removed from heating while zinc dust (0.42 g, 0.063 moles) was added. The reaction was held at reflux for one hour, at which point the conversion was complete by thin layer chromatography. The suspension was diluted with chloroform and saturated sodium bicarbonate, and decanted from the solid zinc. The layers were separated and the aqueous layer was extracted twice more with chloroform. The combined layers were dried (MgSO₄) and evaporated. The maleate salt was prepared in isopropanol. The solvent was decanted and
the orange deliquescent solid was transferred with methanol and stripped to give a foamy orange glass; MS (ES⁺) 272 (MH⁺)

**Example 10**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-aminophenyl)propenamide]

Prepared by a method analogous to that described in Example 9 from (R)-N-(1-azabicyclo[2.2.2]oct-3-yl)[E-3-(2-nitrophenyl)propenamide]; MS (ES⁺) 272 (MH⁺).

**Example 11**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[Z-3-(2-methoxyphenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and Z-3-(2-methoxyphenyl)propenoic acid; MS (ES⁺) 287 (MH⁺).

**Example 12**

(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)-N-methyl-(E-3-phenylpropenamide)

(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-phenylpropenamide) (0.15 g, 0.059 moles)

was suspended in anhydrous tetrahydrofuran under nitrogen atmosphere. The reaction mixture was cooled with an ice bath while commercial 1.0M borane/tetrahydrofuran complex (0.59 mL) was added dropwise by syringe. The suspension dissolved immediately and the solution was stirred at 0°C for one hour. The reaction was carefully quenched by adding water (2 mL) with cooling. The reaction was then diluted with brine and methylene chloride.

The layers were separated and the aqueous layer was extracted twice more with methylene chloride. The combined organic layers were dried (MgSO₄) and evaporated in vacuo. Sodium hydride (28 mg, 0.00071 moles) was suspended in anhydrous N,N-dimethylformamide (2 mL) under nitrogen. The crude borane complex was dissolved in N,N-dimethylformamide (1 mL) and added dropwise. The reaction was stirred at room temperature for 30 minutes and then the methyl iodide (55 µL, 0.000885 moles) was added by syringe. After stirring 2½ h the reaction was cooled with an ice bath and quenched with water (1 mL). The suspension was diluted with saturated sodium bicarbonate and ethyl acetate and the layers were separated. The aqueous layer was extracted twice more and the organic layers were combined, dried (MgSO₄), and concentrated in vacuo. The crude complex was dissolved in acetone (1.6 mL) and water (0.27 mL) was added. The reaction was cooled with an ice bath and aqueous hydrogen bromide (0.27 mL) was added dropwise. The reaction was stirred at 0°C. Additional hydrogen bromide (0.27 mL) was added after 5 h but no further change was detected by thin
layer chromatography. The acetone was removed in vacuo and chased with one portion of methanol and three of ethanol. The crude was triturated first with ethanol and then ether. Two phases resulted, and the lower was separated and washed with more ether, which was decanted. The remainder of the solvent was removed in vacuo. The yellow semi-solid was transferred using methanol and this was removed in vacuo and chased with two portions if ether. Yielded 0.15 g; MS (ES⁺) 271 (MH⁺).

**Examples 13 and 14**

(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2-phenylcyclopropane-1-carboxamide)

Prepared as free bases by a method analogous to that described in Example 1 from (S)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and trans-2-phenylcyclopropane-1-carboxylic acid. The diastereomers could be separated by chromatography on silica gel using a 10-30% methanol/chloroform/ammonium hydroxide gradient; both MS (ES⁺) 271 (MH⁺).

**Examples 15 and 16**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2-phenylcyclopropane-1-carboxamide)

Prepared as free bases by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and trans-2-phenylcyclopropane-1-carboxylic acid. The diastereomers could be separated by chromatography on silica gel using a 10-30% methanol/chloroform/ammonium hydroxide gradient; both MS (ES⁺) 271 (MH⁺).

**Example 17**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-2-fluoro-3-phenylpropenamide)

Prepared as free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and Z-2-fluoro-3-phenylpropenoic acid; MS (ES⁺) 275 (MH⁺).

**Example 18**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-formimidophenyl)propenamide)

Formic acid (98%, 2.9 mL) and acetic anhydride (1.0 mL) were combined under an inert atmosphere while cooling with a cold water bath. (R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-aminophenyl)propenamide) (0.16 g, 0.00059 moles) was added and the reaction was stirred for three days at room temperature. The solution was poured into saturated sodium carbonate and more solid carbonate was added until the solution was basic. The aqueous layer was extracted four times with chloroform. The organic extracts were combined, dried (MgSO₄), and concentrated in vacuo. The product was transferred using chloroform and this
was stripped and chased with two portions of ether. The white solid was dried at room
temperature with high vacuum, giving 64 mg; MS (ES⁺) 300 (MH⁺).

**Example 19**

\[(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-nitrophenyl)propenamide]\]

Prepared as free base by a method analogous to that described in Example 1 from \((R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(4-nitrophenyl)propenoic acid; MS (ES⁺) 302 (MH⁺).

**Example 20**

\[(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-aminophenyl)propenamide]\]

Prepared by a method analogous to Example 9 from \((R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-nitrophenyl)propenamide]; MS (ES⁺) 272 (MH⁺).

**Example 21**

\[(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-formamidophenyl)propenamide]\]

Prepared as free base by a method analogous to that described in Example 18 from \((R)-N-(1-azabicyclo[2.2.2]oct-3-yl)(E-3-(4-aminophenyl)propenamide); MS (ES⁺) 300 (MH⁺).

**Example 22**

\[(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-3-methyl-3-phenylpropenamide)\]

**(A) Z-3-Methyl-3-phenylpropenoic acid**

Copper(I) iodide (1.1 g, 0.006 moles) was suspended in anhydrous ether (20 mL) and the resulting suspension was kept below -10 °C and stirred protected from light while commercial 1.0M methyl lithium solution (6 mL) was slowly added dropwise. The white suspension was stirred at -10 °C for 30 minutes while the solid dissolved. 3-Phenylpropenoic acid (0.44 g, 0.003 moles) was added in three portions at -60 °C. The reaction mixture was then stirred at -10 °C for 90 min and was then poured into dilute aqueous hydrochloric acid. Chloroform was added and the resulting emulsion was filtered through diatomaceous earth, and the phases were then separated. The aqueous layer was extracted three more times with chloroform. The organic layers were combined, dried (MgSO₄), and concentrated. The crude product was dried under high vacuum and used without further purification.

**(B) (R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-3-methyl-3-phenylpropenamide)**

Prepared as a free base by a procedure analogous to that described in Example 1 from \((R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and Z-3-methyl-3-phenylpropenoic acid; MS (ES⁺) 271 (MH⁺).
Example 23

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-N-methylaminophenyl)propenamide]

Sodium metal (0.1 g, 0.0043 moles) was added in two portions to dry methanol (2 mL) stirred at 0°C under nitrogen. After 20 min, (R)-N-(1-azabicyclo[2.2.2]oct-3-yl)[E-3-(4-aminophenyl)propenamide] (0.27 g, 0.001 moles) in methanol was added followed by paraformaldehyde (0.18 g, 0.006 moles), and the resulting solution was stirred at room temperature overnight. The reaction was heated at 50°C for 4 h then sodium borohydride (0.10 g, 0.0028 moles) was added and the solution was heated under reflux for 2 h. The reaction mixture was allowed to cool and aqueous potassium hydroxide (1 M, 0.8 mL) was added. After the resulting mixture had been stirred at room temperature for 1 h, the solution was concentrated in vacuo, and the residue was partitioned between water and chloroform. The aqueous layer was extracted three times more with chloroform. The organic layers were combined, dried (MgSO₄), and concentrated in vacuo. The material was purified first by solid phase extraction on silica using a 10-40% ammoniated methanol/chloroform gradient, and then by reverse phase HPLC on a Waters Bondapak® C₁₈ column using a gradient of 10-60% acetonitrile and 0.1% aqueous trifluoroacetic acid as the eluent. The free base was recovered by neutralization and extraction, yielding 45 mg of a yellow powder; MS (ES⁺) 286 (MH⁺).

Example 24

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-N,N-dimethylaminophenyl)propenamide]

Sodium cyanoborohydride (0.19 g, 0.003 mol) and anhydrous zinc chloride (0.21 g, 0.0015 moles) were combined in anhydrous methanol. The mixture was stirred for 5 min and (R)-N-(1-azabicyclo[2.2.2]oct-3-yl)[E-3-(4-aminophenyl)propenamide] (0.27 g, 0.001 moles) was dissolved in methanol (3 mL) was added, followed by paraformaldehyde (0.18 g, 0.006 moles). The reaction mixture was stirred at room temperature overnight. The methanol was concentrated in vacuo and the residue was taken up in 1 M sodium hydroxide (20 mL). The aqueous layer was extracted three times with chloroform and these layers were combined, dried (MgSO₄) and evaporated in vacuo. The material was purified first by solid phase extraction on silica using a 10-40% ammoniated methanol/chloroform gradient, and then by reverse phase HPLC on a Waters Bondapak® C₁₈ column using a gradient of 10-60% acetonitrile and 0.1% aqueous trifluoroacetic acid as the eluent. The free base was recovered by neutralization and extraction, yielding 12 mg of yellow powder; MS (ES⁺) 300 (MH⁺).
**Example 25**

**(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-3-phenylpropenamide)**

**(A) Z-3-Phenylpropenoic acid**

Nickel (II) acetate tetrahydrate (0.16 g, 0.00063 moles) was dissolved in ethanol (6.3 mL) and placed under hydrogen atmosphere. The green solution was stirred rapidly while sodium borohydride (0.024 g, 0.00063 moles) in ethanol (0.63 mL) was added. The deep purple solution was then treated with ethylene diamine (0.42 mL, 0.0063 moles) followed by 3-phenylpropenoic acid (0.73 g, 0.005 moles). The reaction was stirred under hydrogen for 5 hours. The hydrogen was displaced with nitrogen and the suspension was filtered through diatomaceous earth to remove the catalyst. The filtrate was concentrated *in vacuo*. The residue was taken up in chloroform and water, and concentrated hydrochloric acid was added to acidify the aqueous layer. The layers were separated and the acidic layer was extracted three times more with chloroform. The organic layers were combined and concentrated. The material was purified by reverse phase HPLC on a Waters Bondapak® C_{18} column using a gradient of 10-40% acetonitrile and 0.25% aqueous trifluoroacetic acid as the eluent to give a white solid (0.51 g).

**(B) (R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-3-phenylpropenamide)**

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride, and Z-3-phenylpropenoic acid; MS (ES^+) 257 (MH^+).

**Example 26**

**(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-methyl-3-phenylpropenamide)**

**(A) E-3-Methyl-3-phenylpropenoic acid**

A complex of copper (I) bromide (1.72 g, 0.12 moles) and lithium bromide (1.0 g, 0.012 moles) was prepared in anhydrous tetrahydrofuran and cooled to -50 °C. Commercial methyl magnesium bromide (3.0 M in ether; 4 mL) was added dropwise. The suspension was stirred at -50 °C for 15 min, then 3-phenylpropenoic acid (0.44 g, 0.003 moles) was added in three portions. The mixture was stirred at -50 °C for 15 minutes and then was directly warmed to 30 °C with a warm water bath. After stirring for one hour the reaction mixture was poured into saturated ammonium chloride. The solution was acidified by slowly adding concentrated hydrochloric acid and then was extracted four times with chloroform. The organic layers were combined, dried (MgSO_4_), and concentrated in vacuo. The material was purified by reverse
phase HPLC on a Waters Bondapak® C₁₈ column using a gradient of 20-60% acetonitrile and
0.25% aqueous trifluoroacetic acid as the eluent to give a colourless solid (0.14 g).
(B) (R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-methyl-3-phenylpropenamide)
Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-
azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-methyl-3-phenylpropenoic acid,
giving a light tan solid; MS (ES⁺) 271 (MH⁺).

Example 27
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2,3-diphenylpropenamide)
Prepared as free base by a method analogous to that described in Example 1 from (R)-
1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-2,3-diphenylpropenoic acid; MS
(ES⁺) 333 (MH⁺).

Example 28
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-methoxyphenyl)propenamide)
Prepared as free base by a method analogous to that described in Example 1 from (R)-
1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(4-methoxy)phenylpropenoic acid;
MS (ES⁺) 287 (MH⁺).

Example 29
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2-methyl-3-phenylpropenamide)
Prepared as free base by a method analogous to that described in Example 1 from (R)-
1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-2-methyl-3-phenylpropenoic acid;
MS (ES⁺) 271 (MH⁺).

Example 30
(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2-methyl-3-phenylpropenamide)
Prepared as free base by a method analogous to that described in Example 1 from (S)-
1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-α-methyl-3-phenylpropenoic acid;
MS (ES⁺) 271 (MH⁺).

Example 31
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-(2-methylphenyl)propenamide)
Prepared as a free base by a method analogous to that described in Example 1 from
(R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(2-methylphenyl)propenoic
acid; MS (ES⁺) 271 (MH⁺).
Example 32

$(S)-N-(1$-Azabicyclo[2.2.2]oct-3-yl)$-[E-(2-methylphenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from $(S)-1$-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and $E$-3-(2-methylphenyl)propenoic acid; MS $(ES^+) 271 (MH^+)$.

Example 33

$(RS)-N-(1$-Azabicyclo[2.2.2]oct-3-yl)$-[E-(2-methoxyphenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from racemic 1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and $E$-3-(2-methoxyphenyl)propenoic acid; MS $(ES^+) 287 (MH^+)$.

Example 34

$(R)-N-(1$-Azabicyclo[2.2.2]oct-3-yl)$-[E-(3$-methoxyphenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from $(R)$-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and $E$-3-(3-methoxyphenyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS $(ES^+) 287 (MH^+)$.

Example 35

$(RS)-N-(1$-Azabicyclo[2.2.2]oct-3-yl)$-[E-(4$-methoxyphenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from racemic 1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and $E$-3-(4-methoxyphenyl)propenoic acid; MS $(ES^+) 287 (MH^+)$.

Example 36

$(RS)-N-(1$-Azabicyclo[2.2.2]oct-3-yl)$-[E-(2$-fluorophenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from racemic 1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and $E$-3-(2-fluorophenyl)propenoic acid; MS $(ES^+) 275 (MH^+)$.

Example 37

$(R)-N-(1$-Azabicyclo[2.2.2]oct-3-yl)$-[E-(3$-fluorophenyl)propenamide]

Prepared by a method analogous to that described in Example 1 from $(R)$-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and $E$-3-(3-fluorophenyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol / chloroform mixtures as the eluent; MS $(ES^+) 275 (MH^+)$. 
Example 38

\[(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)][E-3-(4-fluorophenyl)propenamide]\]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(4-fluorophenyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS (ES⁺) 275 (MH⁺).

Example 39

\[(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)][E-3-(2-chlorophenyl)propenamide]\]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(2-chlorophenyl)propenoic acid; MS (ES⁺) 291 and 293 (MH⁺).

Example 40

\[(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)][E-3-(2-chlorophenyl)propenamide]\]

Prepared as a free base by a method analogous to that described in Example 1 from (S)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(2-chlorophenyl)propenoic acid; MS (ES⁺) 291 and 293 (MH⁺).

Example 41

\[(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)][E-3-(3-chlorophenyl)propenamide]\]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(3-chlorophenyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol / chloroform mixtures as the eluent; MS (ES⁺) 291, 293 (MH⁺).

Example 42

\[(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)][E-3-(4-chlorophenyl)propenamide]\]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(4-chlorophenyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol / chloroform mixtures as the eluent; MS (ES⁺) 291, 293 (MH⁺).

Example 43

\[(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)][E-3-(3,4-dichlorophenyl)propenamide]\]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(3,4-dichlorophenyl)propenoic
acid; the compound was purified by chromatography on silica gel using ammoniated methanol / chloroform mixtures as the eluent; MS (ES$^+$) 325, 327, 329 (MH$^+$).

**Example 44**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-bromophenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(3-bromophenyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol / chloroform mixtures as the eluent; MS (ES$^+$) 335, 337 (MH$^+$).

**Example 45**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-bromophenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(4-bromophenyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol / chloroform mixtures as the eluent; MS (ES$^+$) 335, 337 (MH$^+$).

**Example 46**

(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-iodophenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from racemic 1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(3-iodophenyl)propenoic acid; MS (ES$^+$) 357 (MH$^+$).

**Example 47**

(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-iodophenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from racemic 1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(4-iodophenyl)propenoic acid; MS (ES$^+$) 357 (MH$^+$).

**Example 48**

(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-trifluoromethylphenyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from racemic 1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(2-trifluoromethylphenyl)propenoic acid; MS (ES$^+$) 325 (MH$^+$).

**Example 49**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)-[E-3-(3-trifluoromethylphenyl)propenamide]
Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(3-trifluoromethyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS (ES⁺) 271 (MH⁺).

**Example 50**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-furyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(2-furyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS (ES⁺) 247 (MH⁺).

**Example 51**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-furyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(3-furyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS (ES⁺) 247 (MH⁺).

**Example 52**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-pyridyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(2-pyridyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS (ES⁺) 258 (MH⁺).

**Example 53**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-pyridyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(3-pyridyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS (ES⁺) 258 (MH⁺).

**Example 54**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-pyridyl)propenamide]

Prepared by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(3-pyridyl)propenoic acid. The
compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent. The product was dissolved in methanol, excess hydrogen chloride (1M in ether) was added. The solution was then evaporated, and recrystallisation from methanol/t-butyl methyl ether gave the dihydrochloride salt of the title compound as a colourless solid; MS (ES⁺) 258 (MH⁺).

**Example 55**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-thienyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(2-thienyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS (ES⁺) 263 (MH⁺).

**Example 56**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-thienyl)propenamide]

Prepared by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(3-thienyl)propenoic acid; the compound was purified by reverse phase HPLC on a Waters Bondapak® C₁₈ column using a gradient of acetonitrile and 0.1% aqueous trifluoroacetic acid as the eluent. The solvent was evaporated from the product-containing fractions, then the product was dissolved in aqueous hydrochloric acid and the solution was evaporated again to give the hydrochloride salt of the title compound as a colourless solid; MS (ES⁺) 282 (MH⁺).

**Example 57**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-nitro-2-furyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(5-nitro-2-furyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS (ES⁺) 293 (MH⁺).

**Example 58**

(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-methoxy-3-pyridyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(5-methoxy-3-pyridyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS (ES⁺) 288 (MH⁺).
Example 59

\((R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-hydroxy-3-pyridyl)propenamide]\)

Prepared as a free base by a method analogous to that described in Example 1 from \((R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(5-hydroxy-3-pyridyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS (ES\(^+\)) 274 (MH\(^+\)).

Example 60

\((R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-imidazolyl)propenamide]\)

Prepared as a free base by a method analogous to that described in Example 1 from \((R)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(4-imidazolyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent; MS (ES\(^+\)) 247 (MH\(^+\)).

Example 61

\(N-(endo-8-Aza-8-methylbicyclo[3.2.1]oct-3-yl)(E-3-phenylpropenamide)\)

A mixture of 3\(\alpha\)-aminotropane dihydrochloride (3.6 g), E-phenylpropenoic acid (2.5 g), 1-[3-(dimethylamino)propyl]-3-ethyl-carbodiimide hydrochloride(3.2 g), and triethylamine (12 mL) in N,N-dimethylformamide (30 mL) was stirred at room temperature overnight. The solution was evaporated, and the residue was partitioned between aqueous sodium hydroxide and chloroform. The solution was evaporated and the residue was purified by reverse phase HPLC on a Waters Bondapak\textsuperscript{®} C\textsubscript{18} column using a gradient of acetonitrile and 0.1% aqueous trifluoroacetic acid as the eluant. The hydrochloride salt was prepared by evaporation of the product-containing fractions, dissolution of the residue in methanol, addition of excess hydrogen chloride solution (1M in diethyl ether) and evaporation. After drying under vacuum, the dihydrochloride salt of the title compound was obtained as a colourless solid (236 mg); MS (ES\(^+\)) 307 (MH\(^+\)).

Example 62

\(N-(exo-8-Aza-8-methylbicyclo[3.2.1]oct-3-yl)(E-3-phenylpropenamide)\)

A mixture of 3\(\beta\)-aminotropane dihydrochloride (1.83 g), E-3-phenylpropenoic acid (1.57 mg), 1-hydroxybenzotriazole hydrate (1.16 g), O-benzotriazol-1-yl-N,N,N\(^{\prime}\),N\(^{\prime}\)-tetramethyluronium tetrafluoroborate (2.76 g), and N,N-diisopropylethylamine (6.0 mL) in N,N-dimethylformamide (15 mL) was stirred at room temperature overnight. The solution was evaporated, and the residue was partitioned between aqueous sodium hydroxide and
chloroform. The solution was evaporated and the residue was purified by reverse phase HPLC on a Waters Bondapak® C₁₈ column using a gradient of acetonitrile and 0.1% aqueous trifluoroacetic acid as the eluant. The hydrochloride salt was prepared by evaporation of the product-containing fractions, dissolution of the residue in methanol, addition of excess hydrogen chloride solution (1M in diethyl ether) and evaporation. After drying under vacuum, the dihydrochloride salt of the title compound was obtained as a colorless solid (243 mg); MS (ES⁺) 307 (MH⁺).

**Example 63**

\((S)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-furyl)propenamide]

Prepared as a free base by a method analogous to that described in Example 1 from (S)-1-azabicyclo[2.2.2]oct-3-ylamine dihydrochloride and E-3-(2-furyl)propenoic acid; the compound was purified by chromatography on silica gel using ammoniated methanol/chloroform mixtures as the eluent, followed by recrystallization from ethyl acetate / hexane; MS (ES⁺) 247 (MH⁺).
CLAIMS

1. The use of a compound of the general formula I

\[
\begin{array}{c}
\text{II} \\
\text{III} \\
\text{IV} \\
\text{V} \\
\text{VI}
\end{array}
\]

wherein:

A represents:

D represents oxygen, or sulfur;
R\(^1\) represents hydrogen or methyl;
R\(^2\) represents hydrogen, or C\(_1\)–C\(_4\) alkyl;
R\(^3\) represents:

R\(^4\), R\(^5\), and R\(^6\) are independently hydrogen, halogen, C\(_1\)–C\(_4\) alkyl, C\(_2\)–C\(_4\) alkenyl, C\(_2\)–C\(_4\) alkynyl, –CO\(_2\)R\(^7\), –CN, –CF\(_3\), or Ar, provided that at least one of R\(^4\) and R\(^5\) represents Ar;
Ar represents a 5- or 6-membered aromatic or heteroaromatic ring containing zero to three nitrogen atoms, zero or one oxygen atom, and zero or one sulfur atom, or an 8-, 9-, or 10-membered fused aromatic or heteroaromatic ring system containing zero to four nitrogen atoms, zero to one oxygen atoms, and zero to one sulfur atoms which may optionally be substituted with one or more substituents selected from the
following: hydrogen, halogen, C₁–C₄ alkyl, C₂–C₄ alkenyl, C₂–C₄ alkynyl, aryl, heteroaryl, –CO₂R, –CN, –NO₂, –NR₈R₉, –CF₃, –OR¹⁰; R¹, R⁹, and R¹₀ are independently hydrogen, C₁–C₄ alkyl, aryl, heteroaryl, –C(O)R¹¹, –C(O)NHR¹², –C(O)R¹³, –SO₂R¹⁴ or R⁸ and R⁹ may together be (CH₂)₁Q(CH₂)ₖ where Q is O, S, NR¹⁵, or a bond; j is 2 to 4; k is 0 to 2; R⁷, R¹⁰, R¹¹, R¹², R¹³, and R¹⁴, are independently C₁–C₄ alkyl, aryl, or heteroaryl; or an enantiomer thereof, and pharmaceutically acceptable salts thereof, for the preparation of a medicament for the treatment or prophylaxis of psychotic disorders or intellectual impairment disorders.

2. The use according to Claim 1, wherein:
A represents:

![Chemical Structure](image)

3. The use according to Claim 1 or Claim 2, wherein D represents oxygen.

4. The use according to any one of Claims 1 through 3, wherein R² represents hydrogen.

5. The use according to any one of Claims 1 through 4, wherein R³ represents

![Chemical Structure](image)

6. The use according to any one of Claims 1 through 4, wherein:

R³ represents
R\(^4\) represents Ar;
R\(^5\) represents hydrogen, or C\(_1\)-C\(_4\) alkyl; and
R\(^6\) represents hydrogen, or halogen.

7. The use according to Claim 1, wherein:
A represents

![Diagram](image)

D represents oxygen;
R\(^2\) represents hydrogen; and
R\(^3\) represents

![Diagram](image)

8. The use according to Claim 1, wherein the compound of formula I is a compound wherein:
A represents

![Diagram](image)

D represents oxygen;
R\(^2\) represents hydrogen:
R^3 represents

\[ \begin{array}{c}
\text{R}^4 \\
\text{R}^5 \\
\text{R}^6 \\
\end{array} \]

R^4 represents Ar;
R^5 represents hydrogen, or C_1-C_4 alkyl; and
R^6 represents hydrogen, or halogen.

9. The use according any one of Claims 1 to 8, wherein Ar represents a 5- or 6-membered aromatic or heteroaromatic ring containing zero to three nitrogen atoms, zero or one oxygen atom, and zero or one sulfur atom, or an 8-, 9- or 10-membered fused aromatic or heteroaromatic ring system containing zero to four nitrogen atoms, zero to one oxygen atoms, and zero to one sulfur atoms which may optionally be substituted with one or more substituents selected from the following: hydrogen, halogen, C_1-C_4 alkyl, C_2-C_4 alkenyl, C_2-C_4 alkynyl, aryl, heteroaryl, -CO_2R^7, -CN, -NO_2, -NR^8R^9, -CF_3 and -OR^{10}.

10. The use according to Claim 1, wherein said compound is:
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(3-phenylpropynamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl) [E-3-(2-nitrophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl) [E-3-(2-nitrophenyl)propynamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-aminophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[Z-3-(2-methoxyphenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)-N-methyl-(E-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-2-phenylcyclopropane-1-carboxamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-2-fluoro-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-formamidophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-nitrophenyl)propenamide];
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-aminophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-formamidophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-methyl-3-phenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-N-methylaminophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-N,N-dimethylaminophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-methyl-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2,3-diphenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-methoxyphenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2-methyl-3-phenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-methylphenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(3-methoxyphenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-fluorophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-fluorophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-2-chlorophenylpropenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(3-chlorophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-chlorophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)-(E-3-(3,4-dichlorophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)-(E-3-(3-bromophenyl)propenamide)
N-(1-Azabicyclo[2.2.2]oct-3-yl)-(E-3-(4-bromophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(3-iodophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-iodophenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-trifluoromethylphenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(3-trifluoromethylphenyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-furyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(3-furyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-pyridyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(3-pyridyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-pyridyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-thienyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(3-thienyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(5-nitro-2-furyl)propenamide);
N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(5-methoxy-3-pyridyl)propenamide);
11. The use according to claim 1, wherein said compound is:

\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(5-hydroxy-3-pyridyl)propenamide);
\((S)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-imidazolyl)propenamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-nitrophenyl)propenamide);
\((S)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-formamidophenyl)propenamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-formamidophenyl)propenamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-N,N-dimethylamino-phenyl)propenamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(Z-3-methyl-3-phenylpropenamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-aminophenyl)propenamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-formamidophenyl)propenamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-N-methylaminophenyl)propenamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-fluoro-3-phenylpropenamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-phenylcyclopropane-1-carboxamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(2-phenylcyclopropene-1-carboxamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-nitrophenyl)propenamide);
\((R)\)-N-(1-Azabicyclo[2.2.2]oct-3-yl)(E-3-(4-nitrophenyl)propenamide).

or an enantiomer thereof, and pharmaceutically acceptable salts thereof.
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-methoxyphenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-2-methyl-3-phenylpropenamide];
(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-2-methyl-3-phenylpropenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-methylphenyl)propenamide];
(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-methylphenyl)propenamide];
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-methoxyphenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-methoxyphenyl)propenamide];
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-methoxyphenyl)propenamide];
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-fluorophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-fluorophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-fluorophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-2-chlorophenylpropenamide];
(S)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-2-chlorophenylpropenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-chlorophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-chlorophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3,4-dichlorophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-bromophenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-bromophenyl)propenamide];
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-iodophenyl)propenamide];
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-iodophenyl)propenamide];
(RS)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-trifluoromethylphenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-trifluoromethylphenyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-furyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-furyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-thienyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-thienyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-nitro-2-furyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-methoxy-3-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-hydroxy-3-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-imidazolyl)propenamide];
N-(endo-8-Aza-8-methylbicyclo[3.2.1]oct-3-yl)[E-3-phenylpropenamide];
N-(exo-8-Aza-8-methylbicyclo[3.2.1]oct-3-yl)[E-3-phenylpropenamide];
or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

12. A compound of the general formula I

\[
\begin{array}{c}
\text{A} \\
\text{N} \\
\text{D} \\
\text{R}^2 \\
\text{R}^3 \\
\end{array}
\]

wherein:

- A represents:
  - [Structures II, III, IV, V, VI]

- D represents oxygen, or sulfur;
- R\(^1\) represents hydrogen or methyl;
- R\(^2\) represents hydrogen, or C\(_1\)–C\(_4\) alkyl;
- R\(^3\) represents:
  - [Structures for R\(^3\)]

- R\(^4\), R\(^5\), and R\(^6\) are independently hydrogen, halogen, C\(_1\)–C\(_4\) alkyl, C\(_2\)–C\(_4\) alkenyl, C\(_2\)–C\(_4\) alkynyl, –CO\(_2\)R\(^7\), –CN, –CF\(_3\), or Ar, provided that at least one of R\(^4\) and R\(^5\)
  represents Ar;
- Ar represents a 5- or 6-membered aromatic or heteroaromatic ring containing zero to three nitrogen atoms, zero or one oxygen atom, and zero or one sulfur atom, or an 8-,
9- or 10-membered fused aromatic or heteroaromatic ring system containing zero to four nitrogen atoms, zero to one oxygen atoms, and zero to one sulfur atoms which may optionally be substituted with one or more substituents selected from the following: hydrogen, halogen, C1–C4 alkyl, C2–C4 alkenyl, C2–C4 alkynyl, aryl, heteroaryl, –CO₂R⁷, –CN, –NO₂, –NR⁸R⁹, –CF₃, –OR¹⁰, R⁸, R⁹, and R¹⁰ are independently hydrogen, C₁–C₄ alkyl, aryl, heteroaryl, –C(O)R¹¹, –C(O)NHR¹², –C(O)R¹³, –SO₂R¹⁴ or R⁸ and R⁹ may together be (CH₂)₂Q(CH₂)ₖ where Q is O, S, NR¹⁵, or a bond; j is 2 to 4; k is 0 to 2; R⁷, R¹⁰, R¹¹, R¹², R¹³, R¹⁴, and R¹⁵ are independently C₁–C₄ alkyl, aryl, or heteroaryl; or an enantiomer thereof, and pharmaceutically acceptable salts thereof, with the additional proviso that Ar does not represent 2-, 3-, 4-pyridyl, unsubstituted phenyl or phenyl substituted with one or more substituents selected from C₁–C₄ alkyl, C₁–C₄ alkoxy, halogen, phenoxy, hydroxy, OCOR¹¹, NH₂, NHCOR¹¹, and nitro, when:
A represents

\[
\begin{align*}
\text{II} & \quad \text{or} \quad \text{VI} \\
\end{align*}
\]

R³ represents

\[
\begin{align*}
\text{D represents oxygen;} \\
\text{R represents methyl;} \\
\text{R² and R⁵ both represent hydrogen; and} \\
\text{R⁶ represents any of hydrogen, C₁–C₄ alkyl, phenyl, or cyano.}
\end{align*}
\]
13. A compound according to Claim 12, wherein A represents:

![Diagram](image)

II

or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

14. A compound according to Claim 13, wherein $R^6$ represents any of 2-furyl; 3-furyl; 2-thienyl; 3-thienyl; or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

15. A compound according to Claim 12, said compound being:

$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-trifluoromethylphenyl)propenamide];

$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-furyl)propenamide];

$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-furyl)propenamide];

$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-thienyl)propenamide];

$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-thienyl)propenamide];

$N$-(1-Aza-3-cyanobicyclo[2.2.2]oct-3-yl)[E-3-phenylpropenamide];

$N$-(1-Aza-3-methylbicyclo[2.2.2]oct-3-yl)[E-3-phenylpropenamide];

$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-nitro-2-furyl)propenamide];

$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-methoxy-3-pyridyl)propenamide];

$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-hydroxy-3-pyridyl)propenamide];

$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-imidazolyl)propenamide];

or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

16. A compound according to Claim 12, said compound being:

(R)$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-trifluoromethylphenyl)propenamide];

(R)$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-furyl)propenamide];

(R)$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-furyl)propenamide];

(R)$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(2-thienyl)propenamide];

(R)$N$-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(3-thienyl)propenamide];
N-(1-Aza-3-cyanobicyclo[2.2.2]oct-3-yl)[E-3-phenylpropenamide];
N-(1-Aza-3-methylbicyclo[2.2.2]oct-3-yl)[E-3-phenylpropenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-nitro-2-furyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-methoxy-3-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(5-hydroxy-3-pyridyl)propenamide];
(R)-N-(1-Azabicyclo[2.2.2]oct-3-yl)[E-3-(4-imidazolyl)propenamide];
or an enantiomer thereof, and pharmaceutically acceptable salts thereof.

17. A compound according to any one of Claims 12 to 16, for use in therapy.

18. The use of a compound as defined in any one of Claims 12 to 16, in the manufacture of a medicament for the treatment or prophylaxis of psychotic disorders or intellectual impairment disorders.

19. The use of a compound as defined in any one of Claims 12 to 16, in the manufacture of a medicament for the treatment or prophylaxis of human diseases or conditions in which activation of the α7 nicotinic acetylcholine receptor is beneficial.

20. The use of a compound as defined in any one of Claims 12 to 16, in the manufacture of a medicament for the treatment or prophylaxis of Alzheimer’s disease, learning deficit, cognition deficit, attention deficit, memory loss, Attention Deficit Hyperactivity Disorder, anxiety, schizophrenia, mania or manic depression, Lewy Body Dementia, Parkinson’s disease, Huntington’s disease, Tourette’s syndrome, neurodegenerative disorders in which there is loss of cholinergic synapses, jetlag, cessation of smoking, nicotine addiction including that resulting from exposure to products containing nicotine, pain, or ulcerative colitis.

21. The use according to Claim 20, wherein the condition or disorder is Alzheimer’s disease, learning deficit, cognition deficit, attention deficit, memory loss, Lewy Body Dementia, or Attention Deficit Hyperactivity Disorder.
22. The use according to Claim 20, wherein the disorder is anxiety, schizophrenia, or mania or manic depression.

23. The use according to Claim 20, wherein the disorder is Parkinson's disease, Huntington's disease, Tourette's syndrome, or neurodegenerative disorders in which there is loss of cholinergic synapses.

24. The use according to Claim 20, wherein the condition or disorder is jetlag, nicotine addiction including that resulting from exposure to products containing nicotine, pain, or ulcerative colitis.

25. The use according to Claim 20, wherein the condition or disorder is Alzheimer's disease.

26. A pharmaceutical composition including a compound as defined in any one of Claims 12 to 16, in admixture with an inert pharmaceutically acceptable diluent or carrier.

27. The pharmaceutical composition according to Claim 26, for use in the treatment or prophylaxis of psychotic disorders or intellectual impairment disorders.

28. The pharmaceutical composition according to Claim 26, for use in the treatment or prophylaxis of human diseases or conditions in which activation of the α7 nicotinic receptor is beneficial.

29. The pharmaceutical composition according to Claim 26, for use in the treatment or prophylaxis of Alzheimer's disease, learning deficit, cognition deficit, attention deficit, memory loss, Attention Deficit Hyperactivity Disorder, anxiety, schizophrenia, or mania or manic depression, Lewy Body Dementia, Parkinson's disease, Huntington's disease, Tourette's syndrome, neurodegenerative disorders in which there is loss of cholinergic synapse, jetlag, cessation of smoking, nicotine addiction including that resulting from exposure to products containing nicotine, pain, or ulcerative colitis.
30. The pharmaceutical composition according to Claim 29, for use in the treatment or prophylaxis of Alzheimer's disease, learning deficit, cognition deficit, attention deficit, memory loss, Lewy Body Dementia, or Attention Deficit Hyperactivity Disorder.

31. The pharmaceutical composition according to Claim 29, for use in the treatment or prophylaxis of anxiety, schizophrenia, or mania or manic depression.

32. The pharmaceutical composition according to Claim 29, for use in the treatment or prophylaxis of Parkinson's disease, Huntington's disease, Tourette's syndrome, or neurodegenerative disorders in which there is loss of cholinergic synapses.

33. The pharmaceutical composition according to Claim 29, for use in the treatment or prophylaxis of jetlag, nicotine addiction including that resulting from exposure to products containing nicotine, pain, or ulcerative colitis.

34. The pharmaceutical composition according to Claim 29, for use in the treatment or prophylaxis of Alzheimer's disease.

35. A method of treatment or prophylaxis of psychotic disorders or intellectual impairment disorders, which comprises administering a therapeutically effective amount of a compound as defined in any one of Claims 12 to 16.

36. A method of treatment or prophylaxis of human diseases or conditions in which activation of the α7 nicotinic receptor is beneficial which comprises administering a therapeutically effective amount of a compound as defined in any one of Claims 12 to 16.

37. The method according to Claim 35 or Claim 36, wherein the disorder is Alzheimer's disease, learning deficit, cognition deficit, attention deficit, memory loss, Attention Deficit Hyperactivity Disorder, anxiety, schizophrenia, or mania or manic depression, Lewy Body Dementia, Parkinson's disease, Huntington's disease, Tourette's syndrome, neurodegenerative disorders in which there is loss of cholinergic synapse,
jetlag, cessation of smoking, nicotine addiction including that resulting from exposure to products containing nicotine, pain, or ulcerative colitis.

38. The method according to Claim 37, wherein the disorder is Alzheimer’s disease, learning deficit, cognition deficit, attention deficit, memory loss, Lewy Body Dementia, or Attention Deficit Hyperactivity Disorder.

39. The method according to Claim 37, wherein the disorder is Parkinson’s disease, Huntington’s disease, Tourette’s syndrome, or neurodegenerative disorders in which there is loss of cholinergic synapses.

40. The method according to Claim 37, wherein the disorder is anxiety, schizophrenia or mania or manic depression.

41. The method according to Claim 37, wherein the disorder is jetlag, nicotine addiction, pain, and for ulcerative colitis.

42. The method according to Claim 37, wherein the disorder is Alzheimer’s disease.
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

IPC7: C07D 451/04, C07D 453/02, C07D 453/06, C07D 487/08, A61K 31/439, A61K 31/46, A61K 31/40, A61P 25/00

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC7: C07D, A61K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

SE, DK, FI, NO classes as above

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
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<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tr>
<td>X</td>
<td>EP 0581165 A2 (DOMPE'FARMACEUTICI S.P.A.), 2 February 1994 (02.02.94), see esp. page 7, example 8, line 30</td>
<td>12-17, 26-34</td>
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<td>X</td>
<td>WO 9420465 A1 (GLAXO S.P.A.), 15 Sept 1994 (15.09.94), see example 4 and page 5 lines 20-32</td>
<td>1-42</td>
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Further documents are listed in the continuation of Box C. See patent family annex.

Date of the actual completion of the international search: 27 February 2001

Date of mailing of the international search report: 02.03.2001

Name and mailing address of the ISA/Swedish Patent Office:
Box 5055, S-102 42 STOCKHOLM
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Authorized officer:
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Form PCT/ISA/210 (second sheet) (July 1998)
INTERNATIONAL SEARCH REPORT

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. □ Claims Nos.: 35–42
   because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

2. □ Claims Nos.: 12, 15–16 (partly) and 14
   because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

   See extra sheet*

3. □ Claims Nos.: 35–42
   because they relate to subject matter not required to be searched by this Authority, namely:

   See extra sheet**

This International Searching Authority found multiple inventions in this international application, as follows:

1. □ As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.

2. □ As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.

3. □ As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:

4. □ No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

Remark on Protest

□ The additional search fees were accompanied by the applicant's protest.

□ No protest accompanied the payment of additional search fees.
*Claims 35-42 relate to methods of treatment of the human or animal body by surgery or by therapy/diagnostic methods practised on the human or animal body/ Rule. 39.1.(iv). Nevertheless, a search has been executed for these claims. The search has been based on the alleged effects of the compound(s)/composition(s).

**In claim 12 the definition R in the "proviso part" is believed to stand instead of R². The definition of R⁶ in claim 14 seems to be irrelevant with regard to the examples, as R⁶ is never a heterocyclic group in the examples. Some of the compounds of claim 15 and 16 are not within the scope of claim 12 or not even of claim 1.
<table>
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<tr>
<td>EP 0581165 A2</td>
<td>02/02/94</td>
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