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(56) Documents cited GB A 2153821 EP A1 0214772 EP A1 0013138 GB A 2125398 EP A1 0099789 The Extra Pharmacopoeia 28th Edition, The Pharmaceutical Press (1982), p. 362 p. 367 pp. 1445-1446 p. 1504

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(54) Therapeutic use of serotonin antagonists

(57) Use of a mono or bicyclic carbocylic, or heterocyclic carboxylic, acid ester or amide or an imidazolyl carbazol in the manufacture of a medicament suitable for the treatment of stress-related psychiatric disorders, for increasing vigilance, for the treatment of rhinitis or serotonin-induced disorders and/or coadministration with another active agent to increase the bioavailability thereof, or for nasal administration.

2 193 633

SPECIFICATION

Therapeutic use of serotonin antagonists

5	This invention relates to new uses and modes of administration of serotonin 5HT ₃ antagonists and also to mono- or bicyclic carbocylic, or heterocyclic carboxylic, acid esters, and amides and imidazolyl carbazols, e.g. imidazoylmethylcarbazols.	5
10	The compounds may be used in any pharmaceutically acceptable form, including the free base and at least for the esters and amides, in acid addition and quaternary ammonium salt forms. These compounds are referred to hereinafter as compounds of the invention. The above mentioned esters, and amides and imidazolyl carboxols have in general serotonin 5HT ₃ antagonist activity which may have not been previously recognized. These esters, amides and carbazols are in general known for example from Belgian patents 897117, 900425 and	10
15	901274. These compounds are described therein as being serotonin 5HT ₃ receptor antagonists or serotonin M receptor antagonists (serotonin M receptors have been reclassified as 5HT ₃ receptors).	15
20	The compounds are for treatment of anti-migraine agents, anti-arrhythmics, and serotonin-induced gastro-intestinal disorders including emesis, e.g. induced by anti-cancer agents. Other classes of the compounds of the invention are known from e.g. European patent publications 13138A, 250444A, and 214772A and British Patent publication 2153821A. We have now discovered that these compounds have interesting new uses and modes of administration which have been hitherto unrecognized. In a first aspect the present invention provides use of a mono- or bicyclic carbocylic, or cyclic	20
25	heterocyclic carboxylic, acid ester or amide, e.g. of a cyclic alcohol or amine, containing nitrogen as a ring atom, or a serotonin rHT ₃ antagonist or an imidazolyl carbazol, for the treatment of stress-related psychiatric disorders, rhinitis, or serotonin-induced nasal disorders or lung embolism, or coadministration with another active agent to increase the biovailability thereof, or for	25
30	nasal administration or for the manufacture of a medicament suitable therefor. The invention also provides a method of treating a subject with any of the uses which comprises administering to a subject in need of such treatment a compound of the invention. The invention also provides:	30
35	(i) a process for the production of a pharmaceutical composition adapted for the treatment of stress-related psychiatric disorders, for increasing viligance, for the treatment of rhinitis, or for serotonin-induced nasal disorders or lung embolism which comprises working up a compound of the invention with pharmaceutical carrier s and diluents to manufacture unit dosage formulations	35
40	bility which comprises working up a compound of the invention, e.g. a compound of formula I or la as defined hereinafter, with another active agent, e.g. a peptide, and if desired formulating as	40
45	a unit dosage form. In a group of compounds the ester or amide is an ester or amide of a cyclic alcohol, or amine, containing nitrogen as a ring atom. In a sub-group of compounds the heterocycle is a bicyclic heterocycle preferably aromatic.	45
50	In another group of compounds the ester or amide or serotonin HT ₃ antagonist is a dicarbocylic or heterocyclic carboxylic acid ester, or carboxylic acid amide, of a piperidinol containing an alkylene bridge, or of piperidylamine containing an alkylene bridge, or of an ester, or amide, of a substituted benzoic acid, or of a piperidinol or piperidylamine containing an alkylene bridge, with the proviso that in each benzoic acid amide the alkylene bridge of the piperidyl ring is bonded to	50
•	the nitrogen atom and to a cyclic carbon atom. In another group of compounds the ester or amide or serotonin 5HT ₃ antagonist or imidazol-carbazol is a compound of formula l	

carbazol is a compound of formula I

A-B-C-D (I)

$$R_1$$
 R_2

$$R_1$$

5

(IIa)

10

15

(IIc)

20

(IId)

20

(IIe)

25 .

30

35 (111)

(IV)

35

40

$$R_1 - \left(\begin{array}{c} \\ N \end{array}\right)$$

(Y)

40

wherein the free valence is attached to either fused ring in formula II,IIa,IIb,IIe or IV.

- X-Y is -CH=CH-, $-O-CH_2-$ or -N=CH-,
- Z is $-CH_2-$, $-NR_3-$, -O- or -S-,

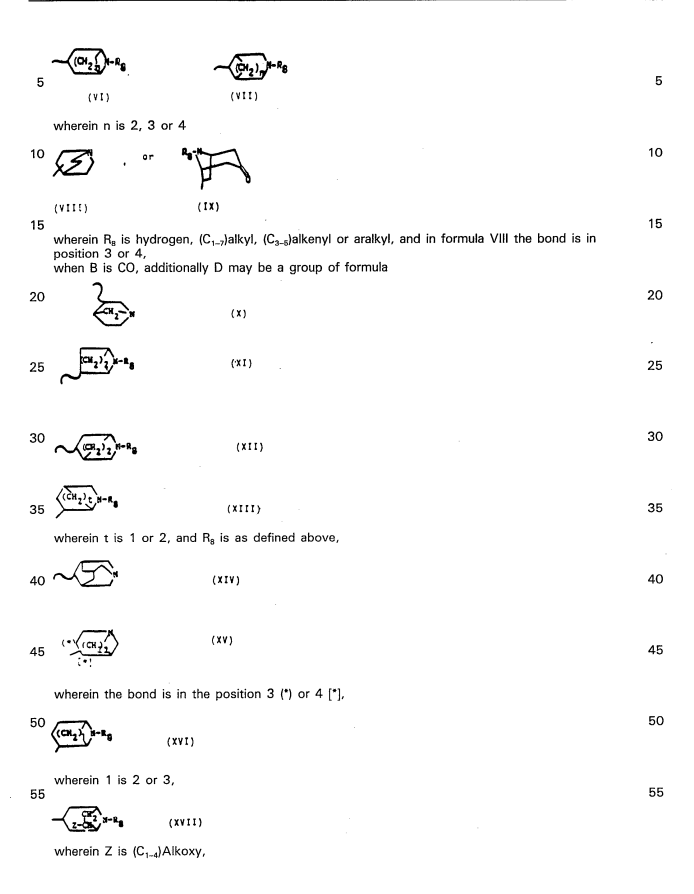
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a,

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- R_1 and R_2 are independently hydrogen, halogen, (C_{1-4}) alkyl, (C_{1-4}) alkoxy, hydroxy, amino, (C_{1-4}) alkylamino, di (C_{1-4}) alkylamino, mercapto or (C_{1-4}) alkylthio,
 - R_3 is hydrogen, (C_{1-4}) alkyl, acyl, (C_{3-5}) alkenyl, aryl or arylalkyl, and
- R_4 to R_7 are, independently, hydrogen, amino, nitro, (C_{1-4}) alkylamino, di (C_{1-4}) alkylamino, halo-
- 50 gen, (C_{1-4}) alkoxy, (C_{1-4}) alkyl, (C_{1-4}) alkanoylamino, pyrrolyl, sulfamoyl, or carbamoyl,

- B is -CO- or $-SO_2-$,
- C is -O- or -NH- or a bond,
- D is a group of formula



25

wherein R_9 to R_{12} are independently hydrogen or (C_{1-4}) , alkyl, m is 0, 1 or 2 and 15 n, o, p independently are 0 or 1,

$$_{20}$$
 -(CH₂)_q -N $_{R_{14}}$ (XIX)

wherein q is 2 or 3, $R_{13} \mbox{ and } R_{14} \mbox{ independently are } (C_{1-4}) \mbox{alkyl}, \mbox{ 25}$

30 wherein the bond is in position 3 or 4,



(XXI)

5

5

(XXII)

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(XXIII)

15

20

(XXIVa)

25

25

(XXIVb)

30

30

35



(XXY)

35 -

65

and R₈ is as defined above,

in free base form, acid addition salt form or quaternary ammonium salt form, or a compound of formula la

10 10

wherein R_{15} is hydrogen, (C_{1-10}) alkyl, $(C_{3-9}$ cycloalkyl, (C_{3-6}) alkenyl, phenyl or phenyl (C_{1-3}) alkyl and one of the groups R_{16} , R_{17} and R_{18} is hydrogen, $(C_{1-6}alkyl, (C_{3-7})cycloalkyl, (C_{2-6})alkenyl or$ phenyl(C_{1-3})alkyl and the others independently are hydrogen or (C_{1-4})alkyl.

In a sub-group the compound is of formula I wherein A chosen from formula II, III, IV and V, 15 R₃ is other than acyl, C is -O- or -NH- and D is chosen from formula VI to XVIII, 15 with the proviso that, when A is formula III, B is CO and C is NH, D is not a group of formula VI, in free form, in acid addition salt form or in quaternary ammonium salt form.

In a further group of compounds of formula I' A is a group of formula II'

$$\mathbf{R}_{1}^{\prime} - \mathbf{R}_{2}^{\prime} \qquad \mathbf{I}\mathbf{I}^{\prime}$$

wherein the free bond may be situated in any of the rings,

25 X' is
$$-CH_2$$
, $-NR_3$, $-O$, $-S$, R'₁ and R'₂ independently of one another, are hydrogen, halogen, (C_{1-4}) alkyl, (C_{1-4}) alkoxy, hydroxy, amino, (C_{1-4}) alkylamino, di (C_{1-4}) alkylamino, mercapto or (C_{1-4}) alkylthio, and

R's is hydrogen, (C₁₋₄)alkyl, (C₃₋₅)alkenyl, aryl or aralkyl, or a group of formula III' 30 30

35 35

 R'_4 to R'_2 independently of one another, are hydrogen, amino, nitro, (C_{1-4}) alkylamino, di (C_{1-4}) lamino, halogen, (C_{1-4}) alkoxy, (C_{1-4}) alkyl, (C_{1-4}) alkanoylamino or pyrrolyl,

B signifies -O- or -NH-, D signifies a group of formula IV', 40

wherein

40

n is 2, 3 or 4 and

$$R_8$$
 is hydrogen, (C_{1-7}) alkyl, (C_{3-5}) alkenyl or aralkyl, or a group of formula V', 50 (C_{1-7}) R_8 is hydrogen, (C_{1-7}) alkyl, (C_{3-5}) alkenyl or aralkyl, or a group of formula V', 50

55 55 with the proviso that when A is a group of formula III and B is -NH-, then D signifies a group

of formula V'. A further group comprises formula I wherein A is a group of formula II or III wherein

B=-CO-, C=-O- or -NH- and D is a group of formula VII, IX, X, XI or one of XIII to XXV or 60 A is group of formula IIa, IIb, IIc, IId, IIe, IV or V and B, C and D are as defined above. 60 Another group of compounds of formula I comprises compounds wherein A is formula II or III, wherein R4 to R7 are other than sulfamoyl or carbamoyl, D is VI or VIII with the proviso when A is III and C is -NH- and D is VIII.

Preferred compounds include:

Indole-3-yl-carboxylic acid endo-8-methyl-8-aza-bicyclo 3,2,1]oct-3-yl ester, (hereinafter com-

	pound E)	
5	benzo[b]thiophen-3-ylcarboxylic acid endo-9-methyl-9-azabicylo[3,3,1]-non-3-yl ester, 5-fluoro-1-methyl-indol-3-yl carboxylic acid endo-9-methyl-9-aza-bicyclo[3,3,1]-non-3-yl ester, 1,2,3,9-tetrahydro-9-methyl-3-[(2-methyl-1H-imidazol-1-yl)methyl-4H-carbazol-4-one (hereinafter compound H) and 1-methyl-indazol-3-yl carboxylic acid 9-methyl-9-azabicyclo[3,3,1]non-3α-yl-amide. In one group of compounds the compounds of formula I is a group of formula II, in particular	5
10	Z is NR ₃ , O, or S. In another group the compound of formula I has a group of formula III. In a sub-group D is VI. In a 2nd sub-group D is VII. In a 3rd sub-group D is VIII. In a 4th sub-group D is IX. In a 5th sub-group D is X. In a 6th sub-group D is XI. In a 7th sub-group D is XIII. In a 9th sub-group D is XIV. In a 10th sub-group D is XV. In a 11th sub-group D is XVI. In a 12th sub-group D is XVIII. In a 13th sub-group D is XVIII.	10
15	Preferably D is VI or VIII. A preferred group of compounds comprises a compound wherein A is a group of formula II wherein R_1 and R_2 are independently hydrogen, halogen, (C_{1-4}) alkyl or alkoxy; R_1 is in position 4 or 5;	15
20	R_3 is hydrogen, acetyl or (C_{1-4}) alkyl and the corresponding bond is in position 3, 4 or 5. The present invention also provides novel compounds of formula lb $A'-CO-C'-D'$ lb	20
25	wherein (1) A' is a group of formula II, wherein R_1 , R_2 and Z are as defined above, C' is $-O-$ or $-NH-$ and D' is a group of formula XIX, wherein q, R_{13} and R_{14} are as defined above, (2) A' is a group of formula III, wherein R_4 and R_6 are each hydrogen, and R_5 and R_7 each chlorine, C' is $-O-$ and D' is one of the groups of formula VI, wherein n is 3 and R_8 as defined above or VIII.	25
30	 (3) A' is a group of formula II, wherein R₁, R₂ and Z are each as defined above, C' is -O- and D' is a group of formula XX, wherein the bond is in position 3 or 4, (4) A' is a group of formula IIa, wherein R₁, R₂ and Z are as defined above, C' is -NH- or -CH₂- and D is a group of formula VI, wherein R₈ is as defined above or VIII, wherein the bond 	30
35	is in position 3 or 4, (5) A' is a group of formula IIb, wherein R ₁ , R ₂ and Z are as defined above, C' is -O- or D is a group of formula VI, wherein R ₈ is as defined above, (6) A' is a group of formula IIc, wherein R ₁ , R ₂ and Z are as defined above, C' is -O- and D' is a group of formula VI, wherein R ₈ is as defined above,	35
40	(7) A' is a group of formula IId, C' is $-O-$ and D' is a group of formula VI, (8) A' is a group of formula II, wherein R_1 , R_2 and Z are as defined above, C' $-NCH_3-$, and D' is a group of formula VI, wherein R_3 is as defined above, (9) A' is a group of formula II, wherein R_1 , R_2 and Z are as defined above, C' is $-O-$ and D' is a group of formula XXI,	40
45	(10) A' is a group of formula III, wherein R_4 — R_7 are as defined above, C' is —NH— and D' is a group of formula XXII, wherein R_8 is as defined above, (11) A' is a group of formula II, wherein R_1 is 6-hydroxy- or 6-methoxy- or 5-methyl, R_2 is hydrogen and Z—NH—, C' is —O— and in a group of formula VI, wherein R_8 is as defined above, (12) A' in a group of formula II, wherein R_1 , R_2 and Z are as defined above, C' is a bond and	45
50	D' is a group of formula VI, wherein R ₈ is as defined above, (13) A' is a group of formula II, wherein R ₁ , R ₂ and Z are as defined above, C' is -O- and D' is a group of formula XXIII, wherein R ₈ is as defined above, (14) A' is a group of formula II, wherein R ₁ , R ₃ and Z are as defined above, C' is -O- and D'	50
55	D' is a group of formula XXIVa, wherein R _s is as defined above,	55
60	(17) A' is a group of formula II, wherein R_1 , R_2 and Z are as defined above, C' is a bond and D' is a group of formula XXIVb, wherein R_8 is as defined above, (18) A' is a group of formula II, wherein R_1 , R_2 are as defined above, Z is N-acyl, C' is $-O-$ and D is a group of formula VI, wherein R_8 is as defined in above, (19) A' is a group of formula II, wherein R_1 , is hydroxy, R_2 is hydrogen and Z is $-NH-$, C' is	60
65	-CH ₂ - and D' is a group of formula VI, wherein R ₈ is as defined in above. in free base form or in acid addition salt form or quaternary ammonium salt form. The present invention also provides a process for the production of a compound of formula Ib A'-CO-C'-D' as defined above in free base form or in acid addition salt form or in quaternary	: 65

65 nary ammonium salt form.

ammonium salt form, which includes the step of:-(a) for the production of a compound of formula lb wherein C' is -O- or -NH- reacting a compound of formula XXX XXX5 5 A'-COOH wherein A' is as defined above, or a reactive derivative thereof, or a precursor of the acid or reactive derivative with an appropriate compound of formula 10 10 XXXI H-C'-D' XXXI wherein C' and D' are as defined above, 15 or a precursor of this compound, or 15 (b) for the production of a compound of formula lb wherein C' is −CH₂-, reacting a compound of formula XXXII A'-Mg-Hal XXXII 20 20 wherein A' is as defined above, and Hal is chlorine, bromine or iodine, with an appropriate compound of formula XXXIII CIOC-CH2-D' XXXIII, or 25 25 wherein D' is as defined above, in free base or protected form under conditions of a Grignard reaction, and removing any protecting group present, (c) for the production of a compound of formula lb wherein C' is a direct bond, reacting a 30 compound of formula XXXII as defined above, with an appropriate compound of formula XXXIV 30 CI-OC-D' **XXXIV** wherein D' is as defined above, 35 35 under conditions of a Grignard reaction, (d) for the production of a compound of formula lb wherein C' is a -NCH₃- group, reacting a compound of formula XXX as defined above with an appropriate compound of formula XXXI wherein C' is -NCH₃-, (e) for the production of a compound of formula lb wherein Z is N-acyl- wherein an appropri-40 ate compound of formula I, wherein A is a compound of formula II and Z is -NH--, B is -CO-, C 40 is -O- and D is as defined above is acylated, and recovering the compound of formula lb in free base form, acid addition salt form or quaternary ammonium salt form. The reactions may be effected in conventional manner, e.g. as described in the patent publica-45 tions referred to above or in analogous manner for known compounds. The processes may be 45 generally effected in an inert solvent at e.g from about -30°C to about 200°C using conventional reagents. Compounds of formula lb wherein C' is -O- or -NH- (i.e. groups 1 to 7, 9 to 11, 13 to 15 and 18) are conveniently produced by process (a), e.g. as described in Belgian Patent No. 50 50 897117. Compounds of formula lb wherein C' is -CH2- (i.e. groups 4 and 19) are conveniently produced by process b), e.g. as described in Belgian Patent No. 903,984. Compounds of formula Ib wherein C' is a direct bond (i.e. groups 12, 16 and 17) may be conveniently produced by process c), e.g. as described in Belgian Patent No. 903,984. Compounds of formula lb wherein C' is -NCH₃- (i.e. group 8) may be conveniently produced 55 according to process (d) which may be effected conveniently as for process (a). Starting materials may be conveniently obtained by reacting the corresponding free amine with chloroformic acid ethyl ester and reduction of the product with lithium aluminium hydride. Compounds of formula Ib wherein Z is N-acyl (i.e. group 18) may be conveniently produced 60 according to process (e), in conventional manner e.g. by acylating a compound as disclosed e.g. 60 in Belgian Patent No. 897,117 with an acylating agent such as acetic acid anhydride, benzyl chloride. The resultant compounds may be converted in conventional manner into acid addition salt form and back into free base forms and also converted in conventional manner into the quater-

Examples of each of these groups are described hereinafter. The antagonistic action against 5-HT₃ receptors of the preferred compound ICS 205-930 (indol-3-yl carboxylic acid endo-8-methyl-8-aza-bicyclo[3,2,1]oct-3-yl ester) on the rabbit vagus, rabbit heart and guinea pig ileum has been described (P.Donatsch et al., Br.J. Pharmacol. 1984, 5 5 81, 348), and also its topical application to humans as the first 5-HT₃ antagonist. In the Belgian Patent No. 897,117 it was also stated that the compounds disclosed therein are indicated as anti-psychotics. We have now found e.g. from ethological and endocrinological tests that the compounds of the invention are useful for the treatment of stress-related psychiatric disorders, including stress 10 related-social phobias and social withdrawal, affective disorders, psychoses, especially maniac 10 depressive disorders and promote approach-oriented behaviour in behaviour perturbed by stress. The compounds of the invention also increase vigilance, e.g. in geriatrics. Trials have been carried out on the effects of the compounds of the invention on approachoriented behaviour in mice. The compounds increase approach-oriented behaviour in stress-15 15 situations normally inhibitory to social responses. Thus situations involving unfamiliar surroundings, contact with foreign aggressive opponents, competition for food, were created under reversed lighting conditions whereby observations were performed during the darkphase. Under these conditions, untreated rodents exihibit high levels of flight, particularly depressive ambivalence and escape behaviour, but low levels of social behaviour involving approach activity. 20 Anxiolytics like diazepam reduce the ambivalent behaviour and can increase social behaviour. In 20 food deprived mice under white light, certain antidepressants e.g. bupropion and impramine, but also a typical anxiolytics e.g. bupropion as well as the compounds of the invention, increase approach-oriented social behaviours. 25 25 Intruder test Study A A foreign male mouse (intruder), placed for 6 minutes, into the cage of an isolated male is attacked and responds with defensive escape patterns collectively known as "flight". Flight and associated defensive activities override the intruder's tendency to approach the attacker so that 30 30 much of the behaviour includes social forms of ambivalence. Intruders receiving oral doses of benzodiazepines show less of these defensive activities and more approach-oriented behaviour (A.K. Dixon Triangle 1982, 21, 95-105; M.Krisak, Br.J. Pharmacol. 1975, 5-5, 141-150), e.g. investigation, aggression and sexual activity. The compounds of the invention are administered 1 hour before the encounter per-orally at 35 from about 0.1 to about 10 mg/kg. Groups of 8 mice pairs are used. The frequency and 35 duration of social and non-social behaviour of the mice were recorded using ethological techniques. The compounds increase social oriented activity. In this test compound E at a dose of 1 mg/kg increased the frequency (from ca 60 to 80) 40 40 and duration of social interaction (from ca 90 to 120 seconds). Study B Study A was modified using a large cage (59×38.5×20cm) which allowed greater freedom of movement so that elements of defense could be separated more clearly from the approach-45 oriented social activities. The compounds of the invention were given i.p. at doses from about 45 0.01 to about 100 microgon/kg 45 minutes before the encounter. Indruders received the drug. It was found that the compounds of the invention clearly promoted the approach-oriented social activities. They also reduced escape behaviour, a special form of flight. In this test at a dose of 1 microgram/kg compound E promoted 8 elements of 50 approach oriented social activities by about 40 per cent. Compound E also reduced escape 50 behaviour. Compound H at the same dose promoted 3 elements of approach-oriented social activities by about 40 per cent. It's effect on escape behaviour was less. It did however increase cage exploratory behaviour indicative of a stimulant effect. 55 55 Competitive feeding Study C In this test 8 pairs of male OF-1 mice deprived of food for 6 hours are forced to compete over 6 minutes for a single food pellet. Because of the close proximity of the mice the tendency to eat is competitively offset by the tendency to interact socially. One partner receives a 60 60 compound of the invention at an oral dose from about 0.01 to about 1 mg/kg 1 or 2 hour before the encounter. Frequencies and durations of social activities were recorded for both animals with the aid of a posture and timer machine (K.Hausamann, A.K.Dixon, Physiol. Behav. 1982, 28, 743-745).

In this test benzodiazepines increase eating behaviour more than social interactions. Antide-

65 pressants may increase such social activities. The compounds of the invention increase ap-

proach-oriented social behaviour, more than eating.

Compound E at a dose of 0.1 mg/kg p.o. given 2 hours before the encounter increased, relative to controls, the frequency (54 percent) and duration (321 per cent) of social interactions. In contrast frequency of feeding increased by only 4 per cent and their duration 55 per cent.

Stretched Attend Postures (SAP) (No conflict)

Mice placed upon an unfamiliar elevated platform in a novel environment display characteristic stretched body postures called SAP's which signify ambivalence. Drugs which have putative anxiolytic reactions, e.g. benzodiazepines, barbiturates, and buspirone reduce the incidence of 10 SAP's (H.P. Käsermann, Psychopharmacology 1986, 89, 31-37).

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The compounds of the invention administered p.o. at a dose of from about 0.1 to about 10 mg/kg. 2 hours before the test reduce the duration of SAP when placed on the platform for 2 minutes under conditions where no other mice are present.

The durations in the case of compound E at 0.1 mg/kg were similar to that observed with 15 dobazam at the same dose and at higher doses the effect was less.

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Corticosterone levels (Endocrinological profile)

Mice subjected to a novel environment, e.g. on transferring them from one room to another via a trolley, exhibit a rise in plasma corticosterone typical of stress related disturbances which 20 are reduced by benzodiazepines and barbiturates (Lahti R.A., Borsulm C., Res. Comm.Chem.Path. Pharm.11: 596-603; G.Le Fur et al., J.Pharm.exp.Ther. 211: 305-308), reduction is observed with the compounds of the invention at from about 0.1 to 10 mg/kg p.o. compound E reduces such stress-induced corticosterone at about 1 to 10 mg/kg p.o whilst lower doses from about 0.1 to about 0.3 mg/kg increase basal plasma levels of this hormone. This profile is analogous 25 to that observed with diazepam.

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Taken together, the results of these studies also show that compounds of the invention promote approach-oriented social behaviour in stressful situations. This suggests that the compounds of invention are of use in stress-related psychiatric disorders, e.g. where the treatment of social withdrawal, affective disorders, and other stress-related illnesses is desired.

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The increases in corticosterone also suggest that compounds of invention increase vigilance, thus indicating a potential use for the compounds in disorders of vigilance e.g. geriatric illnesses.

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The compounds of the invention may be administered in similar manner to known standards. e.g. bupropion. The preferred compound E and bupropion have been found to have a pronounced effect on promoting social interaction under stress related conditions. For example, 35 compound E increased the mean frequency of social interactions in the competitive feeding experiment study (c) by 54 per cent of 0.01 mg/kg compared to bupropion provoking an increase of 179 per cent at 2.5 mg/kg. It is indicated that compound E will be useful in the treatment of stress-related psychiatric disorders at a daily oral dose of from 0.1 mg which could

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Compounds of the invention including and excluding (i) compounds of formula I wherein A is a group of formula II or III, B is CO, C is -O- or -NH- and D is a group of formula VI, VIII and XII and (ii) compounds of formula lb also have an anxiolytic effect which more general indication is also indicated by the above testing.

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The use of the compounds of formula I wherein (i) A is a group of formula II or III and B is 45 -CO-, C is -O- or -NH- and D is a group of formula VII, IX, X, XI and one of XIII to XXV, and (ii) A is a group of formula IIa, IIb, IIc, IId, IIe, IV or V as an anxiolytic or in the manufacture of a medicament available therefor also forms part of the present invention and is also shown by the above testing.

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For these indications, the appropriate dosage will, of course, vary depending upon, for 50 example, the compound employed, the host, the mode of administration and the nature and severity of the condition being treated. However, in general, satisfactory results are indicated to be obtained at daily dosages from about 0.001 mg/kg to about 50 mg/kg animal body weight. In humans, an indicated daily dosage is in the range from about 0.1 mg of about 50 mg of a compound of the invention conveniently administered, for example, in divided doses up to four 55 times a day.

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In a further aspect the present invention provides use of a mono or bicylic carboxylic or heterocyclic carboxylic acid esters or amides of a cyclic alcohol or amine containing nitrogen as a ring atom in free base form or in acid addition salt or quaternary ammonium salt form as a 5-HT₃ antagonists in the manufacture of a medicament suitable for the treatment of serotonin

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60 induced psychiatric disorders, e.g. when chosen from one of the following: anxiety, social withdrawal, affective disorders, psychoses and other stress-related illnesses, disorders of vigilance, e.g. geriatic illnesses

60

Preferred compounds include:-

be increased up to 50 mg.

Indol-3-yl carboxylic acid endo-8-methyl-8-aza-bicyclo[3,2,1]oct-3-yl ester [hereinafter com-65 pound E].[ICS]

Benzo[b]thiophen-3-yl carboxylic acid endo-9-methyl-aza-bicyclo[3,3,1]non-3-yl ester [hereinafter compound F]. 5-fluoro-1-methyl-indol-3-yl carboxylic acid endo-9-methyl-9-aza-bicyclo[3,3,1]non-3-yl ester [hereinafter compound G]. 1,2,3,9-tetrahydro-9-methyl-3-[(2-methyl-1H-imidazol-1-yl)methyl]-4H-carbazol-4-one (hereinafter 5 compound H).[GR 38032F] 1-methyl-indazol-3-yl-carboxylic acid 9-methyl-9-aza-bicyclo[3,3,1]non-3 α -yl-amide (hereinafter compound I), and especially compound E. The compounds of the invention may be administered in free base form or, when they can be 10 10 formed, in pharmaceutically acceptable acid addition salt form or in a quaternary ammonium salt form. Such salts may be prepared in conventional manner and are in general known. They exhibit the same order of activity as the free base form and pharmaceutical compositions comprise a compound of the invention in free base or pharmaceutically acceptable acid addition salt form or quaternary ammonium salt form in association with pharmaceutical carrier or diluent. 15 15 Such compositions may be manufactured in conventional manner. The compounds may be administered by any conventional route, in particular enterally, preferably orally, e.g. in the form of tablets or capsules or parenterally, e.g. in the form of injectable solutions or suspensions. Suitable pharmaceutical carriers and diluents for oral administration include polyethylene glycol, 20 polyvinylpyrrolidone, mannitol, lactose etc. granulating agents, and disintegrating agents such as 20 starch and algenic acid, binding agents such as stearic and gelatine, lubricating agents such as magnesium stearate, stearic acid and talc. Suspensions may contain conserving agents like ethyl p-hydroxy-benzoate, suspending agents such as methyl-cellulose, tenside etc. For parenteral forms the compositions are preferably bufferred, aqueous solutions (pH between 4 and 5). We have moreover found that the compounds of the invention may be administered nasally 25 and have an especially interesting resorption profile. Furthermore the compounds of the invention increase the nasal resorption or bioavailability of other active agents such as peptides particularly when administered by the nasal route. The compounds of the invention moreover are useful in the treatment of rhinitis and serotonin-30 induced nasal disorders as indicated by an inhibition of nasal secretions on administration of the 30 compounds of the invention. The testing may be effected as follows:-The bioavailability and pharmacokinetic profile of the compounds of the invention may be determined in conventional manner, e.g. in mammals including rhesus monkeys and humans. The 35 35 concentrations of the compounds of the invention in the blood plasma after administration of from about 0.01 to about 10 mg/kg to each nostril, e.g. 7.5 mg in the case of compound E, locally to the nasal mucous membrane, e.g. as a spray, may be determined in conventional manner by e.g. radioimmunoassay or HPLC methods. The compounds of the invention are rapidly absorbed, e.g. over about 10 minutes. Even after ca. 5 to 10 minutes following nasal administration, 200 ng of the compound indol-40 3-yl-carboxylic acid-endo-8-methyl-8-aza-bicyclo[3,2,1]oct-3-yl ester may be detected in 1 ml of plasma. Upon oral administration, this concentration of active ingredient in the plasma is reached only after ca. 30 to 40 minutes. The general bioavailability of the compounds of the invention over a period of 6 hours is the same for nasal administration as for oral administration. 45 Nasal secretions are also inhibited. Additionally, the compounds of the invention when administered, e.g. at a dose of from 0.01 to 10 mg/kg with a therapeutically effective dossage of another compound, e.g. a peptide, such as salmon calcitonin increases the absorption thereof. For example in the case of compound E (15 mg) and salmon calcitonin (100 lU) half of which is applied to each nastril, the bioavailability of salmon calcitonin (AUC up to 2 hours) is 50 increased from 0.08 IV)/ml/hr plasma to 1.632 m IU/ml/hr/plasma in the rhesus monkey. 50 For the rhinitis and nasal serotonin-induced disorder indications, the appropriate dosage will, of course, vary depending upon, for example, the compound of invention employed, the host, the mode of administration and the nature and severity of the condition being treated. However, in general, satisfactory results are indicated to be obtained at daily dosages from about 0.01 55 mg/kg to about 10 mg/kg animal body weight. In humans, an indicated daily dosage for oral 55 administration is in the range from about 5 mg to about 300 mg of a compound of formula I conveniently administered, for example, in divided doses up to four times a day, e.g. in the range of about 40 mg p.o. in the case of compound E. When a compound of the invention is co-administered with another active agent, the appropri-60 ate dosage will, of course, vary depending upon, for example, the active agent of the compound 60 of the invention and other active agent employed, the host, the mode of administration and the nature and severity of the condition being treated. However, in general, satisfactory results are indicated to be obtained at daily dosages from about one half to one tenth the usual dose of the

other active agent. The compound of the invention is indicated to be administered at about one

65 half to one tenth the usual dose.

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	The compounds of the invention may be administered for the rhinitis and nasal serotonin-	
	induced disorders and for co-administration with another active agent, e.g. a peptide, by any conventional route, in particular enterally, preferably orally, e.g., in the form of tablets or capsules, or parenterally, e.g., in the form of injectable solutions or suspensions. The local application but the parent route is preferred.	_
5	tion by the nasal route is preferred. For the nasal administration route, the appropriate dosage will, of course, vary depending upon, for example, the compound of the invention employed, the host, and the nature and	5
10	severity of the condition being treated. However, in general, satisfactory results in animals are indicated to be obtained at daily dosages from about 0.001 mg/kg to about 10 mg/kg animal body weight. In humans, an indicated dosage per actuation is in the range from about 0.01 mg to about 1 mg of a compound of the invention conveniently administered, for example, in doses up to four times a day.	10
15	Thus for gastrointestinal disorders or for migraine prophylaxis a compound according to the invention is indicated to be administered to the body nasally in a dosage of 0.13 to 0.4 mg kg body weight, i.e. ca. 10 to 30 mg or 1 to 3 pumps of the nasal spray per patient, and in order to control arrhythmia, it should be given in a dosage which is ca. 10 times higher, i.e. from 1.3 to 4 mg per kg body weight or 100 to 300 mg, or resp. 10 to 30 pumps of the nasal spray	15
20	per patient. The compound E is the preferred compound for the rhinitis and nasal route administration. It is indicated that the compound of example may be administered at daily dosages of about 0.1 mg nasally to humans.	20
	The compounds of the invention may be administered nasally in any pharmacologically active form, e.g. in free base form, in acid addition salt form or in quaternary ammonium salt form. The nasal mode of administration creates a simple method of administration which rapidly	
25	gives results and can be easily carried out by the patient himself, e.g. by administering a liquid form for nasal administration, for example a nasal spray or drop solution using a nasal applicator, or by inserting a gelatinous sponge or lyophilisate soaked in the active substance, or by blowing the galenic form in powder form into the nostrils.	25
30	The compounds of the invention may be present in the liquid form for nasal administration in a proportion of 1 to 30%, preferably 5 to 20%, especially 10 to 15% (weight/volume). The present invention accordingly provides also to a liquid form for nasal administration,	30
	containing (1) a compound of the invention	
35	 (2) a preservative, especially benzalkonium chloride, and (3) a liquid diluent or a carrier, suitable for application to the nasal mucous membrane. The proportion of benzalkonium chloride in the compositions according to the invention is preferably ca. 0.002 to ca. 0.02, especially ca. 0.01% (weight/volume) of the total composition. 	35
40	In accordance with the invention, the above-mentioned forms of administration may be administered to the nasal mucous membrane, e.g. as drops or as a spray. As described hereinafter, however, they are preferably administered as a spray, i.e. as finely dispersed droplets. One further possible way of bringing the above-mentioned liquid form for nasal administration into contact with the nasal mucous membrane is to soak a gelatinous sponge (SPONGOSTAN) or	40
45	lyophilistate with the substance and then to insert the sponge into the nostrils. The liquid diluent or carrier employed in conveniently water (pharmaceutical grade). An aqueous salt solution is preferred in particular. The liquid forms for nasal administration according to the invention are formulated such that they allow administration to be effected nasally. With this	45
50	in mind, they can e.g. also contain minimal amounts of further desired components or excipients, e.g. additional preservatives, or e.g. ciliary stimulants such as caffeine. The liquid forms for nasal administration according to the invention preferably have a pH value of 5.5 to 6.	50
	The liquid forms for nasal administration should also have an appropriate isotonicity and viscosity. They preferably have an osmotic pressure of ca. 260 to ca. 380 m0sm/litre. The desired viscosity of the compositions according to the invention depends on the relevant form of administration, e.g. whether nasal drops or a nasal spray are administered. For nasal drops, a	
55	viscosity of ca. 2 to ca. 40×10^{-3} Pa.S is suitable. For nasal sprays, a viscosity of less than 2×10^{-3} Pa.S is suitable. If desired, the liquid forms for nasal administration may also contain further components,	55
60	especially conventional pharmaceutically available surface-active agents. In this connection and as a further aspect of the present invention, it was found that the use of surface-active compounds in the pasal administration of the compounds of the invention increases their respection through	60

60 in the nasal administration of the compounds of the invention increases their resorption through

e.g. of the general formula,

the nasal mucous membrane and improves the initial bio-availability. In this case, preference is given to non-ionic surface-active agents, for example polyoxyalkylene ethers or higher alcohols,

RO $\left\{ (CH_2)_{n} - O \right\}_{x} H$

5 5 wherein RO signifies the radical of a higher alkanol, especially a higher alkanol such as lauryl or cetyl alcohol, or of an alkylphenol, or a sterol, especially lanosterol, dihydrocholesterol or cholesterol, as well as mixtures of two or several such ethers. Preferred polyoxyalkylene ethers which can be used for the present invention are polyoxyethylene- and polyoxypropylene-ethers (i.e. 10 wherein n is the above-mentioned formula is 2 or 3), especially lauryl-, cetyl- and cholesterylpo-10 lyoxyethylene- and -polyoxypropylene-ethers, as well as mixtures of two or several such ethers. Especially suitable polyethers for use according to the invention are those in which the average value of the recurring units in the polyoxyalkylene component (x in the above formula) lies between 4 and 75, especially between 8 and 30 and particularly between 16 and 26. The 15 polyethers may be obtained in accordance with known methods. A large choice of such pro-15 ducts is available commercially and is sold e.g. by Amerchol under the trade name Solulan (R), by KAO Soap, ICI and Atlas under the trade names Emalex (R), Brij (R) and Laureth (R), and by Croda under the trade name Cetomacrogol (R). Examples of polyoxyalkylene ethers which are suitable for use according to the invention, e.g. 20 20 (POE=polyoxyethylene ethers: POP=polyoxypropylene ether; x=average value of the recurring units in the POE/POP component) are listed in the following:-1. Cholesterylethers: Solulan(R) C-24-POE, x=241.1 25 25 Ethers of lanolin alcohols: 2. Solulan(R) 16-POE, x=162.1 Solulan(R) 26-POE, x=25Solulan(R) 75-POE, x=752.3 30 30 2.4 Solulan(R) PB-10-PPE, x=10Solulan(R) 98-POE, x=10-partly acetylated 2.5 Solulan(R) 97-POE, x=9-wholly acetylated 3. Laurylethers: 35 35 3.1 Emalex(R) 709/Laureth(R)-POE, x=93.2 Laureth(R) 4/Brii(R) 30-POE, x=4Laureth(R) 23/Brij(R) 35-POE, x=233.3 Cetylethers: 4. 40 40 4.1 Cetomacrogol(R)-POE, x=20 to 24 Lanolin alcohols are also known as wool fat alcohols and are a mixture of cholesterol, dihydrocholesterol and lanosterol. Preferred polyethers for use according to the invention are cholesteryl polyoxyethylene ethers, 45 e.g. polyethers of the above formula, wherein n=2 and RO is a cholesteryl radical, in particular 45 polyethers wherein the number of recurring units in the polyoxyethylene component is 16 to 26, especially about 24. These polyethers are preferably free from impurities, and especially from other polyoxyalkylene ethers. They preferably contain at least 75%, most particularly at least 85% and especially at 50 50 least 90% (weight) of the pure cholesteryl polyoxyethylene ether. If a surface-active agent, e.g. a polyoxyalkylene ether, is used, the amount present in the compositions according to the invention will depend on the surface-active agent used in particular, the form of administration (e.g. drops or spray) and the desired effect. In general, the amount of surface-active agent employed is between ca. 2.0 and ca. 200 55 55 (preferably up to ca. 100, especially up to ca. 20), especially between ca. 5 and ca. 30 (preferably up to ca. 15) and in particular ca. 10 mg/ml. For nasal administration, the liquid forms for nasal administration are preferably placed in an applicator, which is equipped with a device that enables the composition to be applied to the nasal mucous membrane, e.g. a nasal spray applicator. Such applicators are known per se and include those which are suitable for the administration 60 of liquid preparations as drops or as a spray to the nasal mucous membrane. Since the dosingof the compounds of the invention should be as exact as possible, the use of spray applicators in which an exact control over the quantity administered is possible is generally preferred. Suitable appliance for administration are e.g. atomizers such as pump dispensers or aerosol

65 cans. In the latter case, the applicator contains a composition according to the invention as well

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	as a propellant which is suitable for use in a nasal spray applicator. The atomizer appliance is provided with an appropriate spray device which enables the composition to be applied to the nasal mucous membrane. Such devices are known in general. The container, e.g. a nasal spray applicator, may contain a quantity of the composition which is sufficient for a single nasal dose or for administration several doses, e.g. over a period of several days or weeks. The amounts of the individual doses will preferably correspond to the	5
1	above-mentioned doses. Applicators as defined above are preferably spray applicators for nasal usage. They preferably enable the composition contained therein to be administered in single doses of ca. 0.05 to ca. 0.15 ml, e.g. ca. 0.1 ml.	10
	Suitable compositions, as well as the individual components 1,2 and 3 for use in an applicator, are those which have been previously described. The dosages which are suitable for use similarly correspond to the dosages given previously.	
1	Furthermore, the invention relates to a process for the production of a liquid form for nasal administration, containing (1) compounds according to the invention (2) a preservative, especially benzalkonium chloride, and	15
2	(3) a liquid diluent or a carrier, which are suitable for administering to the nasal mucous membrane, as well as optionally a surface-active agent which is suitable for administering to the nasal mucous membrane, characterised in that the components are intimately mixed together, and if desired, the composition obtained is placed in an applicator which is provided with a spray device which enables the composition thus obtained to be administered to the nasal mucous membrane. Furthermore, a sponge (SPONGO-STAN) may be soaked with the compo-	20
2	sition obtained and the soaked sponge can be inserted into the nostrils. The stability of the composition according to the invention can be determined in the usual	25
•	way. The compositions according to the invention containing benzalkonium chloride are stable towards contamination by germs, e.g. as in standard tests such as those described by S. Urban et al. in Zbl.Bakt.Hyg.I Abt.Orig.B.1972,478–484 (1981) and S.Urban,Acta Pharm.Technol.22,	30
3	0 247–253 (1976). For example, the cell count of the standard bacteria, namely E.coli ATCC 8739, Pseud.aeruginosa ATCC 9027, Staph.aureus ATCC 6538, Strept. pyogenes ATCC 8668 and standard fungi Cand.albicans ATCC 10231, Sacch.cerevisae ATCC 9763. Aspergillus niger ATCC 16404 and Pen.steckil ATCC 10499, is reduced to 0.1% or less within 24 hours after	30
3	injecting them with the composition, as can be shown in standard tests. In a stability test, the nasal spray composition of the following example 1 was kept for 3 months at 30°C under a nitrogen atmosphere in a glass container. Pseud.aeruginosa ATCC 9027, Staph.aureus ATCC 6538, Strept. pyogenes ATCC 8668 and the fungi cand.albicans ATCC 10231, Sacch.cerevisae ATCC 9763, Aspergillus niger ATCC 16404 and Pen.stechii	35
4	ATCC 10499 were added until a cell count of ca. 2×10 ⁵ organisms was reached in the injected liquid. Within 2 hours, the germ count had reduced to less than 0.1%. Within 4 weeks, the germs could no longer be detected. Equally favourable results are obtained if the compounds of the invention are administered in a	40
4	galenic form, which is in the form of a powder and is introduced by blowing into the nostrils. The compounds of the invention other than compounds of formula I wherein A is 3,5- dichlorophenyl, B is CO, C is O and D is tropanyl antagonise the pulmonary depressor reflex in animals.	45
5	The action of the compounds may be observed in spontaneously breathing rabbits which are anesthetized by a continuous infusion of sodium pentobarbital. Both vagi are intact and the systemie, arterial blood pressure, heart beat, breathing rate and platelet count are normal. Pulmonary embolism is produced by injecting 1 mg Sephadex G-25 beads suspended in 0.2 ml dextran (6%) in 1 minute intervals in 6 animals into the right atrium. Pretreatment with the compounds of the invention i.v. at a dose of from 0.1 to 1 mg/kg	50
5	produces a reduction in mortality and an improvement in the cardiovascular and breathing reflex parameters resulting during the developing lung embolism. For this indication, the appropriate dosage will, of course, vary depending upon, for example,	55
	the compound of the invention employed, the host, the mode of administration and the nature and severity of the condition being treated. However, in general, satisfactory results in animals are indicated to be obtained at daily dosages from about 0.1 mg/kg to about 5 mg/kg animal body weight. In larger mammals, for example humans, an indicated daily dosage is in the range	
. 6	O from about 5 mg to about 50 mg of a compound of the invention conveniently administered, for example, in divided doses up to four times a day, preferably parenterally. The compounds are preferably administered in the form of a pharmaceutical composition e.g. as described above.	60
6	The compounds of the invention may be administered for the lung embolism, by any conventional route, in particular enterally (if appropriate), preferably orally, e.g., in the form of tablets or	65

	capsules, or parenterally (if appropriate), e.g., in the to or by the nasal route.	form of injectable solutions or suspensions,	
E	The compounds of the invention also inhibit cancel indicated by standard tests, e.g. an inhibition of cis-perrets at a dose of from about 0.005 to about 0.5	platinin (10 mg/kg i.v.) induced emesis in	5
5	The compounds of the invention furthermore are u HT ₃ -induced gastro-intestinal disorders, e.g. as indica 189002 at the same order of activity.	seful in the treatment of other serotonin	
10	The compounds are useful in the treatment of disc movements in the intestines and intestinal disorders tors, including diarrhea, e.g. sercretory diarrhea, bact	arising or from activation of 5-HT ₃ recep-	10
15	traveller's diarrhea and psychogenic diarrhea, Crohn's syndrome. The compounds are also indicated to be hypersecretion in the intestines, e.g. as a result of in peptic ulcer, biliary dyskinesia, appendicitis, ulcerative	disease, spastic colon and irritable bowel useful in the treatment of disorders due to flammation such as arising out of gastritis,	15
	leading to increased 5-HT secretion. Furthermore, the compounds are useful in the treat peristaltic movements in the stomach and/or stomac receptors, including those arising from decreased gas	h disorders arising from activation of 5–HT ₃ stric emptying, including treatment of oeso-	
20	phageal motility disturbances, achalasia, hiatus hernia gastroduodeinal reflux, stomach hypotonia and pyloru. The compounds are moreover useful in treatment For all these indications, the compounds may be a	, cardia insufficiency, gastrooesophageal and us hyperplasia. of schizophrenia and mania and anxiety.	20
25	rhinitis indications and in the same manner as described Toxicity and Tolerability: Toxicity and Tolerability studies may be effected in	in European Patent Publication No. 189002.	25
	of the invention to determine the upper dosage. Toxicity studies may be effected for example in the weeks.		30
30	For compound E over 26 weeks the no toxic effect p.o For the rat it was 16 to 45 mg/kg per day p.o. have the same order of tolerability. In healthy human were well tolerated without relevant side effects.	Other compounds of the invention may	00
35	The following examples illustrate the invention.		35
	EXAMPLE 1: Tablets for oral administration Tablets containing the constituents as specified be and are used in the indications specified above.	elow were produced in conventional manner	
40	Compound E in form of hydrochloride (corresponding to 15 mg free base) Hydroxy-propyl-cellulose 1.2 mg		40
	Corn Starch 12.0 mg Lactose 92.8 mg		45
45	Silica 0.6 mg Magnesium stearate 1.5 mg	_	45
	Tablet weight 125.0 mg		
50	EXAMPLE 2: Capsules for oral administration Capsules containing the constituents as specified and are used in the indications specified above.	below are produced in conventional manner	50
55		16.9 mg	55
•	Silica	28.7 mg 1.5 mg	
60		3 mg 	60
	EXAMPLE 3: Injection solution for i.v. administration	on	
65	A composition for injection is made up in conven- mg a day.	tional manner and is used at a dose of 10	65

65 I wherein:

hydrochloride Acetic acid (99 to 100%)* 1.2 0.6 0.6 0.6 Sodium acetate 3. H ₂ O* 1.8 3.18 3.18 3.18 Sodium holoride 8 8 7.5 6.5 10 Water for injection to 1.0 ml 10 mg free base, 0=2 mg free base, 0=3 mg free base, 0=4 mg free base, 0=4 mg free base pH value 4.3; 18 mg free base, 0=4 mg free base, 0=4 mg free base, 0=4 mg free base, 0=4 mg free base pH value 4.3; 18 mg free base, 0=4 mg free base, 0=4 mg free base pH value 4.3; 18 mg free base, 0=4 mg free base pH value 4.3; 18 mg free base, 0=4 mg free base pH value 4.3; 18 mg free base, 0=4 mg free base pH value 4.3; 18 mg free base, 0=4 mg free base pH value 4.3; 18 mg free ba								
hydrochloride Acetic acid (99 to 100%)* 1.2 0.6 0.6 0.6 Sodium acetate 3. H ₂ 0* 1.8 3.18 3.18 3.18 Sodium holoride 8 8 7.5 6.5 10 Water for injection to 1.0 ml 10 ml gree base, \(\text{D} = \text{D} = \text{D} \) mg free base, \(\text{D} = \text{D} = \text{D} \) mg free base pH value 4.3; *Buffer used 1/30 molar* 15 \(EXAMPLE 4: \text{ Capsules for oral administration} \) 5 mg and 15 mg capsules (A and B respectively) containing the constituents as specified below were produced in conventional manner and are used in the indications specified above 2-4 times a day in the case of A and once a day in the case of B. 20 \(Amg \) B mg Compound E in form of hydrochloride 20 ms 44.92 79.29 Lactose 200 mesh 84.929 79.29 Lactose 200 mesh 84.929 79.29 Lactose 200 mesh 84.93 79.29 Correstarch 120.00 120.00 Silica 3.0 mg 300 mg Capsules containing other weights can be formulated in conventional manner. 31 \(EXAMPLE 5: \text{Nasal liquid Composition} \) Capsules containing other weights can be formulated in conventional manner. 32 \(EXAMPLE 5: \text{Nasal liquid Composition} \) The solution obtained is filtered (e.g. through a 0.2 \(\mu\) mfilter) and filled into a nasal canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered e.g. for the treatment of rhinitis, unger embolism or to improve the absorption of other active agents. 50 \(EXAMPLE 6: \text{Nasal liquid Composition} \) Components Components 51 \(\text{The solution obtained is filtered} \) (e.g. through a 0.2 \(\mu\) mfilter) and filled into a nasal canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered e.g. for the treatment of rhinitis, unger embolism or to improve the absorption of other active agents. 52 \(\text{Lamble E: Nasal liquid Composition} \) The solution obtained is filtered (e.g. through a 0.2 \(\mu\) mfilter) and filled into a nasal spray canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered in analogous manner to th			-	A	В	С		
Sodium chloride	5	hydrochloride Acetic acid (99 to 100%)*					5
11	10	Sodium chloride Water for injection (1)=1 mg free base, (2)=2 pH value 4.3;		8 1.0 ml	7.5	6.5		10
A mg B mg Compound E in form of hydrochloride 5.641 16.92 12. Lactose 200 mesh 84.929 79.29 Lactose 100 mesh 84.43 79.29 Corn starch 120.00 120.00 Silica 1.5 1.5 Magnesium stearate 3.0 3.0 Capsules containing other weights can be formulated in conventional manner. 35 EXAMPLE 5: Nasal liquid Composition Components Cuantity of components Cuantity of components 40 indol-3-yl-carboxylic acid-endo-8-methyl-8-aza-bicyclo[3,2,1]bct-3-yl-ester. HCl 100 mg benzalkonium chloride 0.1 mg NaCl (0.9% aqueous solution) 0.6 ml distilled water The solution obtained is filtered (e.g. through a 0.2 µm filter) and filled into a nasal canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered e.g. for the treatment of rhinitis, lung embolism or to improve the absorption of other active agents. EXAMPLE 6: Nasal liquid Composition Components Cuantity of components Cuantity of components 55 EXAMPLE 6: Nasal liquid Composition Components Cuantity of components Components Components Ouantity of components 56 EXAMPLE 6: Nasal liquid Composition Components Ouantity of components The solution obtained is filtered (e.g. through a 0.2 µm filter) and filled into a nasal spray canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered in analogous manner to that disclosed in example 5. The active agents in Examples 1 to 6 may be replaced by the following compounds of formula	15	EXAMPLE 4: Capsules 5 mg and 15 mg caps below were produced in	ules (A and conventiona	B respe I manne	ctively) contair r and are used	I in the indications s	as specified pecified above	15
Compound E in form of hydrochloride	20							20
hydrochloride 5.641 16.92 Lactose 200 mesh 84.929 79.29 Lactose 100 mesh 84.43 79.29 Corn starch 120.00 120.00 Silica 1.5 1.5 Magnesium stearate 3.0 3.0 Topical scontaining other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules containing other weights can be formulated in conventional manner. Capsules capsules capsules capsules capsule			A mg		B mg			
Magnesium stearate 3.0 3.0 300 mg 300	25	hydrochloride Lactose 200 mesh Lactose 100 mesh Corn starch	84.929 84.43 120.00		79.29 79.29 120.00			25
Capsules containing other weights can be formulated in conventional manner. 35	•							
Components Quantity of Components Quant	30		300 mg	-	300 mg			30
Components Components Quantity of components 40 indol-3-yl-carboxylic acid-endo-8-methyl- 8-aza-bicyclo[3,2,1]oct-3-yl-ester. HCl 100 mg benzalkonium chloride 0.1 mg NaCl (0.9% aqueous solution) 0.6 ml distilled water 0.4 ml 45 The solution obtained is filtered (e.g. through a 0.2 μm filter) and filled into a nasal canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered e.g. for the treatment of rhinitis, lung embolism or to improve the absorption of other active agents. 50 EXAMPLE 6: Nasal liquid Composition Quantity of components 51 -methyl-N-endo-9-methyl-9-azabicyclo- [3,3,1]indol-3-yl-carboxylic acid amide 50 mg benzalkonium chloride 0.1 mg NaCl (0.9% aqueous solution) 0.83 ml distilled water 0.17 ml The solution obtained is filtered (e.g. through a 0.2 μm filter) and filled into a nasal spray canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered in analogous manner to that disclosed in example 5. The active agents in Examples 1 to 6 may be replaced by the following compounds of formula		Capsules containing otl	her weights	can be f	formulated in c	conventional manner.		
Components components components 40 indol-3-yl-carboxylic acid-endo-8-methyl- 8-aza-bicyclo[3,2,1]oct-3-yl-ester. HCl 100 mg benzalkonium chloride 0.1 mg NaCl (0.9% aqueous solution) 0.6 ml distilled water 0.4 ml The solution obtained is filtered (e.g. through a 0.2 μm filter) and filled into a nasal canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered e.g. for the treatment of rhinitis, lung embolism or to improve the absorption of other active agents. EXAMPLE 6: Nasal liquid Composition Components Components Cuantity of components 50 mg benzalkonium chloride 0.1 mg NaCl (0.9% aqueous solution) 0.83 ml distilled water 0.17 ml The solution obtained is filtered (e.g. through a 0.2 μm filter) and filled into a nasal spray canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered in analogous manner to that disclosed in example 5. The active agents in Examples 1 to 6 may be replaced by the following compounds of formula	35	EXAMPLE 5: Nasal liqui	id Compositi	on				35
8-aza-bicyclo[3,2,1]oct-3-yl-ester. HCl		Components			•			
The solution obtained is filtered (e.g. through a 0.2 μm filter) and filled into a nasal canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered e.g. for the treatment of rhinitis, lung embolism or to improve the absorption of other active agents. 50 EXAMPLE 6: Nasal liquid Composition Components Components 1-methyl-N-endo-9-methyl-9-azabicyclo-[3,3,1]indol-3-yl-carboxylic acid amide benzalkonium chloride 0.1 mg NaCl (0.9% aqueous solution) 0.83 ml distilled water Components The solution obtained is filtered (e.g. through a 0.2 μm filter) and filled into a nasal spray canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered in analogous manner to that disclosed in example 5. The active agents in Examples 1 to 6 may be replaced by the following compounds of formula	40	8-aza-bicyclo[3,2,1]oct-3- benzalkonium chloride NaCl (0.9% aqueous solu	yl-ester. HC		0.1 mg 0.6 ml			40
Components Components Components 1-methyl-N-endo-9-methyl-9-azabicyclo- [3,3,1]indol-3-yl-carboxylic acid amide 50 mg benzalkonium chloride 0.1 mg NaCl (0.9% aqueous solution) 0.83 ml distilled water 0.17 ml Components 158 1-methyl-N-endo-9-methyl-9-azabicyclo- [3,3,1]indol-3-yl-carboxylic acid amide 50 mg benzalkonium chloride 0.1 mg NaCl (0.9% aqueous solution) 0.83 ml distilled water 0.17 ml Components 158 158 159 159 150 150 150 150 150 150	45	a gelatinous foam (SPON	GOSTAN) is	soaked	gh a 0.2 μm fi with the solut	ion. It is administere	ed e.g. for the	45
Components Components Components	50	EXAMPLE 6: Nasal liqui	d Compositi	on				50
[3,3,1]indol-3-yl-carboxylic acid amide 50 mg benzalkonium chloride 0.1 mg NaCl (0.9% aqueous solution) 0.83 ml distilled water 0.17 ml 60 The solution obtained is filtered (e.g. through a 0.2 μm filter) and filled into a nasal spray canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered in analogous manner to that disclosed in example 5. The active agents in Examples 1 to 6 may be replaced by the following compounds of formula		Components		-				
The solution obtained is filtered (e.g. through a 0.2 μ m filter) and filled into a nasal spray canister, or a gelatinous foam (SPONGOSTAN) is soaked with the solution. It is administered in analogous manner to that disclosed in example 5. The active agents in Examples 1 to 6 may be replaced by the following compounds of formula	55	[3,3,1]indol-3-yl-carboxyli benzalkonium chloride NaCl (0.9% aqueous solu	c acid amide		0.1 mg 0.83 ml	• .	•	55
		The solution obtained i canister, or a gelatinous analogous manner to that The active agents in Ex	foam (SPON t disclosed i	GOSTAN n examp	gh a 0.2 μ m fi N) is soaked w Ne 5.	ith the solution. It is	administered in	60

						÷		n(VI)		
	No.	A=11 R ₁	B= -C0- R ₂	Z .	CO-Position	С	Conf.	D = VIII (pos.)	R ₈	
5										
	1	Н	Н	NH	3	NH	endo	3 (VI)	CH3	
10	2	5-F	Н	NCH ₃	3	0	endo	3 (VI)	Н	
10	3	н	2-01	NH	3	0	endo	2 (VI)	CH ₃	
	4	Н	2-0CH ₃	NH	3	0	endo	2 (VI)	CH ₃	
15	5	Н	3-J	NH	4	0	endo	2 (VI)	CH ₃	
	6	Н	Н	NH	4	0	endo	2 (VI)	CH ₃	
	7	Н	Н	NH	4	0	endo	3 (VI)	CH ₃	
20	8	5-C1	H	NH	3	0	endo	2 (VI)	CH ₃	
	9	4-0CH ₃	Н	NH	3	0	endo	2 (VI)	CH ₃	
	10	5-0CH ₃	Н	NH	3	0	endo	2 (VI)	CH ₃	
25	11	Н	Н	NCH ₃	3	0	endo	2 (VI)	CH ₃	
	12	Н	Н .	NH	3	0	exo	2 (VI)	CH ₃	
	13	5-F	Н	NH	3	NH	endo	2 (VI)	CH ₃	
30	14	Н	Н	NCH ₃	3	NH	endo	2 (VI)	CH ₃	
	15	Н	2-CH ₃	NH	3	NH	endo	2 (VI)	CH ₃	
	16	Н	Н	NH	3	NH	exo	2 (VI)	CH ₃	
35	17	Н	Н	NH	3	NH	endo	2 (VI)	CH ₃	
	18	5 - C1	Н	NH	3	Н	endo	2 (VI)	CH ₃	
40	19	Н	Н	NH	3	0	endo	3 (VI)	Bz	
40	20	н	Н	NCH ₃	3	0	endo	3 (VI)	Bz	
	21	5-F	Н	NH	3	0	endo	3 (VI)	Bz	
45	22	Н	Н	S.	3	0	endo	3 (VI)	CH ₃	
	23	Н	Н	S	3	NH	endo	3 (VI)	CH ₃	
	24	Н	Н	0	3	NH	endo	3 (VI)	CH ₃	
50	25	Н	н	0	3	0	endo	3 (VI)	CH ₃	
	26	Н	Н	CH ₂	3	NH	endo	3 (VI)	CH ₃	

J

					· .			n(VI)		
-	No.	A=II, R ₁	B= -CO- R ₂	Z	Carboxyl- Position	С	Conf.	D = VIII(pos.)	R ₈	5
5										3
	27	Н	Н	NH	3	NH	exo	4 (VI)	CH3	
10	28	Н	H	NH	3	0	exo	4 (VI)	CH ₃	10
	29	Н	Н	NH	3	0	endo	3 (VI)	CH ₃	
	30	Н	Н	NH	3	0	endo	2 (VI)	n-C ₃ H ₇	
15	31	Н	Н	NH	3	0	exo	2 (VI)	Bz	15
	32	Н	Н	NH	3	0	endo	2 (VI)	Bz	
00	33	Н	Н	NH	3	0	endo	2 (VI)	Н	20
20	34	5-F	Н	NH	3	0	endo	3 (VI)	Н	20
-	35	Н	Н	NCH ₃	3	. 0	endo	3 (VI)	Н	-
25	36	Н	Н	NH	3	0	endo	3 (VI)	Н	25
25	37	5-CH ₃	Н	NH	3	0	endo	3 (VI)	сн3	20
	38	Н	2-CH ₃	NH	3	0	endo	3 (VI)	CH ₃	
30	39	5-F	Н	NCH ₃	. 3	0	endo	3 (VI)	CH ₃	30
	40	5-F	Н	NH	3	0	endo	3 (VI)	CH ₃	
	41	5-F	Н	NCH ₃	3	0	.endo	3 (VI)	Bz	
35	42	Н	Н	NCH ₃	3	0	endo	3 (VI)	CH ₃	35
	43	5-CH ₃	Н	NH	3	NH	endo	3 (VI)	CH ₃	
	44	Н	- Н	NH	5	0	endo	2 (VI)	CH ₃	
40	45	Н	H	NH	5	0	endo	3 (VI)	CH ₃	40
	46	Н	3-J	NH	· 5	0.	endo	3 (VI)	CH ₃	
	47	Н	Н	NH	4	NH	exo	2 (VI)	CH ₃	
45	48	Н	Н	NH	4	NH	endo	2 (VI)	CH ₃	45
	49	Н	H	NH	5	Н	endo	2 (VI)	CH ₃	
	50	Н	Н	NH	3	0	-	VIII (3)	-	

C

								n(IV)	
5	No.	A=III, R ₄	B= R ₅	-co- R ₆	R ₇	С	Conf.	D = VIII(pos.)	R ₈
J		· · · · · ·							
	51	0CH ₃	Н	NHCH ₃	C1	0		VIII (3)	
10	52	0CH ₃	Н	NH ₂	C1	0		2 (VI)	Bz
	53	0CH ₃	Н	NH ₂	C1	0	exo	2 (VI)	Н
	54	0CH ₃	Н	NHCH ₃	C1	0	endo	2 (VI)	CH ₃
15	55	0CH ₃	Н	N(CH ₃) ₂	Н	0	exo	2 (VI)	Bz
	56	осн ₃	Н	NH ₂	C1	0.	endo	2 (VI)	CH ₃
20	57	OCH ₃	Н	NH ₂	C1	0	endo	2 (VI)	Н
20	58	OCH ₃	Н	NH ₂	Н	0	endo	2 (VI)	Н
	59	0CH ₃	Н	NH ₂	Н	0	exo	2 (VI)	Н
25	60	OCH ₃	Н	NH ₂	Н	0	endo	2 (VI)	CH ₃
25	61	OCH ₃	Н	N(CH ₃) ₂	Н	0	endo	2 (VI)	CH ₃
	62	C1 .	Н	NH ₂	Н	0	endo	2 (VI)	CH ₃
30	63	0CH ₃	I	NH ₂	Н	0	endo	2 (VI)	CH3
	64	0CH ₃	I	NHCH ₃	H	0	endo	3 (VI)	. CH ³
	65	0CH ₃	Н	NHCH ₃	H	0	endo	3 (VI)	CH ₃
35	66	C1	Н	NO ₂	Н	0	endo	2 (VI)	CH ₃
	67	0CH ₃	Н	Br	Н	0	endo	2 (VI)	CH ₃
	68	Н	C1	Н	C1	0	endo	3 (VI)	CH ₃
40	69	0CH ₃	Н	1-Pyrrolyl	C1	0	endo	2 (VI)	CH3
	70	0CH ₃	Н	1-Pyrrolyl	Н	0	endo	2 (VI)	CH3
	71	0CH ₃	Н	NHCH ₃	C1	NH		VIII (3)	
45	72	H	c1	H	C1	C1		VIII (3)	
	73	OCH ₂	Н	NH2	c1	NH		VIII (3)	

	No.	Formula R _l	II, B=	= -CO- Z	Carboxyl- Position	С	Conf.	D = Group	R ₈	
5					-				· · · · · · · · · · · · · · · · · · ·	5
	74	H :	Н	NH	3	0	exo	(X)		
10	75	H	Н	NH	3	0	endo	(XVII) (Z=OCH ₃)*	CH3	1.0
ΙŲ	76	Н	Н	NH	3	0	(endo)	(XVI) (r=3)	CH ₃	10
	77	Н	H į	NH .	3	0	endo	(XI)	CH ₃	
15	78	Н	Н	NH	3	0	(endo)	(XVI) (r=2)	CH ₃	15
	79	Н	Н	NH	3 .	0	(exo)	(XVI) (r=3)	CH ₃	
_	80	Н	Н	NH	3	. 0	endo	(X)		
20	81	Н	Н	NH	3	0	exo	(XIII)(t=1)	CH ₃	20
	82 (-)	Н	Н	NH	3	0	endo	(XI)	CH ₃	
	83 (+)	Н	Н	NH	3	0	endo	(XI)	CH ₃	
25	84	Н	Н .	NH	· 3	0	endo	(XII)	CH ₃	25
	85	Н	Н .	NH	5	0	endo	(XVII) (Z=OCH ₃)*	CH ₃	-
30		······································						J		30
	() = R	ting is i	n Chair	form						
35	* = (1s*, 3r*	, 5R*,	5R*)						35
00	Bz in F	R ₈ = Benz	yl		·				-	50
40			-							40
.5	No.	A=I R 4	II, B= R ₅	-co- ^R 6	R ₇	С	Conf.	n(VI) D = VIII(pos.)	R ₈	***
45		00:		1 11011		AU.		/v\		45
	86	OCH	_	NHCH ₃	C1	NH	exo	(X)		
	87	.OCH	a H	NHCH	C1	NH	endo	(X)		

No.	۳ م م	rmula I = -CO-	T,	Carboxvl	-		L1_	Formula	XVIII						
	2 & T	R ₁ R ₂	2	Position	ی	Conf.	R ₈	Rg		R ₁₁	R ₁₂	E .	5	0	a.
										-					
88 (-)	Ŧ	Ŧ	-NH-	က	10-	S	CH,	I	I	1	ı	2	0	0	
89	I	±	-NH-	က	-0-	~	Ğ.	I	I	I	Ξ	2	0	0	_
06	×	=	-N-	က	-0-	RS	ິ ສິ	I	=	=	I		0		0
16	Ξ	Ξ	-N-	က	-0-	RS	, E	I	I	I	=	0	_		_
95	=	Ŧ	-HH-	3	-0-	RS	Ğ.	Ξ	I	=	=	2	0	_	0
93	I	=	-N-	က	-0-	RS	GF,	I	Ŧ	I	Ξ	_	_		0
94	工	=	-N-	က	-0-	RS	, =	СH	£.	CH,	CH,	_	_		0
95	≖	=	-N-	က	-0-	RS	CH ₃) =	, =	· =) =	2	_		0
96	I	I	-NH-	က	-NH-	RS	· =	Ŧ	=	Ŧ	I	_	0	_	0
26	I	±	-HN-	က	- NH-	RS	CH ₃	ェ	I	I	Ξ	_	0	_	0

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.0		- .	- poor
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E			0
R ₁₂		Ξ	=
R		Ŧ	I
R10		=	Ξ.
Rg		x	<u> </u>
8 8		Ŧ	CH ₃
Conf.		RS	RS
ပ		· H	¥
R ₇		C1	C
ж 6		NH,	NHCH ₃
R 5			=
R 4		0СН,	0CH ₃ H
No.		86	66
	R ₄ R ₅ R ₆ R ₇ C Conf. R ₈ R ₉ R ₁₀ R ₁₁ R ₁₂ m q o	R ₄ R ₅ R ₆ R ₇ C Conf. R ₈ R ₉ R ₁₀ R ₁₁ R ₁₂ m q o	R ₄ R ₅ R ₆ R ₇ C Conf. R ₈ R ₉ R ₁₀ R ₁₁ R ₁₂ m q o o och, H NH, C1 NH RS H H H H H I O I

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R ₈	CH ₃	CH ₃	CH ₃	CH ₃	CH ₃	CH ₃	НЭ	СНЗ		;	ł	;		!	:
c	ຕີ	က	က	က	က	က	1	w.		ı	ı	i		ŧ	1 .
Position Conf. of g	anti	syn	አ	४	४	४	(15*, 5R*, 6R*)	anti		RS	RS	RS		RS	RS
Posi of g										က	3	က		က	က
D = Group of Formula	(VII)	(\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	(VI)	(NI)	(IA)	(VI)	(IX)	(\(\) \(\)		(VIII)	(VIII)	(VIII)		(VIII)	(1111)
ပ	0	0	0	0	0	0	0	Ħ		0	Ħ	볼		¥	¥
Position of B	(3)	(3)	(3)	(3)	(2)	(3)	(3)	(3)	·	1	1	· ·			,
æ	00	00	00	00	20	200	8	8		S0,	50,	່ວວ		SO2NH2 CO	SO2NH2 CO
2	¥	H	1	!	s t	;	I N	H	R ₇	I	=	工		SON	50^{2}_{2} N
χ	1	;	N=CH	-	HD=HD	0-сн,	7	;	R ₆	CH,	CH,		₁ 2	ェ	Ξ
R ₂	I	I	工	1	I	Ξ	=	I	R ₅	Ŧ	Ŧ	0=	≥ -0-	Ŧ,	Ξ
٣ ا	×	=	×	Ŧ	±	×	Ξ	: =	R ₄	Ξ	Ŧ	Ξ		ен э	I
A = Group of Formula	(11)	(11)	(11)	(v)	(1V)	(1V)	(11)	(11)		· (III)	(111)	(111)		(111)	(111)
No.	100	101	102	103	104	105	106	107		108	109	110		111	112

mp. (° C)	133-134	109-110	143-144	104-105	78,5-79,5	178-179 (Dec.) (Oxalate)	131-132	126-127	106-107	139-140
R14 1						62 ^H 5				C ₂ H ₅
R ₁₃	C2H5	C ₂ H ₅	$c_2^H_5$	C ₂ H ₅	C2H5	c ₂ H ₅	C ₂ H ₅	C ₂ H ₅	$c_2^H_5$	$c_2^H_5$
of q	2	2	2	2	7	2	7	2	2	2
D Group of Formula	XIX	XIX	XIX	XIX	XIX	XIX	XIX	XIX	XIX	XIX
Conf.	1	1	ı		. 1	1		ı	ı	ı
ပ	H.	H	HN	Ŧ	H	0	¥	HZ.	HN	H
Carboxyl- Position	2	е	4	9	7	S	ស	2	2	ស
. 2	HN	H	H	H	H	Ŧ	HN	Ĭ	Ħ	NCH ₃
No. R ₁ R ₂	×	I	I	· =	I	±	3-Br	3-1	3-CH ₃	· =
۾ آ	I	±	×	Ŧ	I	I	Ŧ	=	· =	I
No.	113	114	115	116	1117	118	119	120	121	122

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No.	A = 111, R4 R5	111, R ₅	B = C0 R6	R7	U	Conf.	n (VI) D = (VIII)(pos.)	15.) R ₈	mp. (° C)
123	Ŧ	C1	±	C1	0	I	3 (VI)	CH ₃	170-171 (Malonate)
124	±	Cl	Ŧ	CJ	0	1	(VIII) 3	ı	159-160 (Malonate)
				A					
No.	R .	A = II, R ₁ R ₂	B = C0 Z	Carboxyl- Position	cyl- ion	U	Conf.	D = (XX)(pos.)) mp. (° C)
124	エエ	= =	H H	ю r		0 0	2R*, 3S* 2S*, 3S*	(XX) 3 (XX) 3	230-232 (Dec.) 270-272 (Dec.) (Hydrochloride
									,

mp. (° C)	271-272 279-281	248-250 (Dec.) (Hydrochloride)	112-113 Described Europ. Pat. App. No. 200 444
. 8 8	=	CH ₃	
(VIII)(pos.) D = n (VI)	(VIII) 3 2 (VI)	D = n (VI) 2 (VI)	D = n (VI) 2 (VI) 3 (VI)
ပ	NH CH ₂	0	O H
Carboxyl- Position	າ ວ	က	с т ст
2	H H	1	1 1
B = C0 R ₂	н н	11b, B = CO H	IIc, B = CO H H
A = IIa, R ₁	6-0CH ₃	A H	ч н Н
No.	127	129	130

mр. (°С)	242-243 (Hydrochloride)	233-234 (Hydrochloride)		247-248 (Hydrogenoxalate)
R ₈	CH ₃	cH ₃		CH ₃
(VIII)(pos.) D = n(VI)	3 (VI)	3 (VI)		NCH ₃ 3 (VI)
J	0	0		NCH ₃
Carboxyl- Position	2	က		ന
Z 0	! !		00	H
$B = C$ R_2	ı		B = C0	ェ
$A = IId, B = C0$ $R_1 \qquad R_2$	t		A = II,	=
No.	132	133		134

135 н н 136 н н	ν ν	ო ო	0 0	9s 9r	(XXI) (XXI)	176
No. A = III, R4 F	III, B = CO R ₅ R ₆	R _Ž C	Conf.	٥	88 8	mp. (° C)

No.	A = II, R ₁	B = C0 R ₂	2	Carboxyl- Position	ں	Conf.	D = n (VI)	88 8	mp. (° C)
138	138 6-0сн ₃	≖	H	m ·	0	MESO	2 (VI)	CH ₃	243-244 (Hydrochloride)
139	Н0-9	Ŧ	¥	Ю	0	MESO	2 (VI)	CH ₃	290 (Dec.) (Hydrochloride)
140	5-CH ₃	±	I	က	0	MESO	2 (VI)	CH ₃	284-286 (Dec.) (Hydrochloride)
141	ェ	±	Ħ	က	ı	MESO	2 (VI)	CH ₃	278-280 (Dec.) (Hydrochloride)
	A = 11	, B = C0					. = 0		
142	=	±	Ä	ო	0	ENDO	(XXIII)	сн3	274-276 (Dec.) (Hydrochloride)
	A = 11	, B = C0					= 0		
143	Ξ	Ξ	ĭ	Ю	0	MESO	(XXII)	CH ₃	262-263 (Dec.) (Hydrochloride)

No.	$A = II$ R_1	IIe, B R ₂	B = C0	Carboxyl- Position	J	Conf.	D = n(VI)	R ₈	mp. (° C)
144	1	1	HN.	m	0	1	2 (VI)	CH ₃	181-184 (Malonate)
	# 4	11,	B = C0				" Q		
145	I	x	H	m	•	ı	(XXIVa)	CH ₃	221-223
							= Q		
146	=	±	H	က	1	í	(XXIVb)	CH ₃	230
							D = n(VI)		
147	五	×	0=0-1 V	က	0	ENDO	2 (VI)	CH ₃	170-171
148	H0-9	エ	NH NH	့က	CH ₂	MESO	2 (VI)	=	> 280

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D = XXI	XXV known from Europ. Pa-tent appl. 94 742	mp. 229-230° С
Conf.	t	described in Brit. Patent appl.
ن	Ŧ.	described in Brit. Patent
Carboxyl- Position		nerein CH_3 and R_{18} = H
R7	C1	ein ₁₃ and R
. c0 . R	NH 2	Compound of formula Ia wherein 150 R ₁₅ = CH ₃ , R ₁₆ = CH ₃ an
II, B =	=	formula = CH ₃ ,
No. $A = III$, $B = CO$ $R_4 \qquad R_5 \qquad R_6$	149 0CH ₃	Compound of formula Ia wh 150 R ₁₅ = CH ₃ , R ₁₆ =
No.	149	Сотро 150

The above compounds charaterised by the melting point are new (except where otherwise stated). They can be produced according to the processes described in Belgian Patents 897 117 and 903 984 as well as the following reference examples:

3

Reference example for the preparation of the compound No. 147 1-Acetyl-1H-indol-3-carboxylic acid 8-methyl-8-azabicyclo[3.2.1]oct-3lpha-yl-ester 2.84 g 1H-indol-3-carboxylic acid 8-methyl-8-azabicyclo[3.2.1]oct-3 α -yl-ester are dissolved at 30° ml tetrahydrofuran. The solution is cooled to 0° and treated at this temperature dropwise 5 5 with 5.9 ml butyl-lithium within 15 minutes. A slightly exothermic reaction takes place. The mixture is stirred at 0° for 1 hour, then cooled to -10° and treated dropwise with a solution of 0.75 ml acetyl chloride in 4 ml tetrahydrofuran. The mixture is stirred overnight at room temperature and then partitioned between 2 N aqueous sodium carbonate solution and CH2 Cl2. The organic phase is evaporated to give the title compound, which cristallizes from CH2 10 10 Cl₂/C₂H₅OH, m.p. 170–171°. Reference example for the preparation of the compound No. 113 N-[2-(N,N-Diethylamino)ethyl]indol-2-carboxamide To a suspension of 4.83 g indol-2-carboxylic acid and 3.8 g N-hydroxy-succinimide in 60 ml 15 abs. acetonitrile is added at room temperature a solution of 6.8 g dicyclohexyl-carbodiimide in 15 30 ml abs. diethyl ether, whereby the temperature arises rapidly to 33°. The suspension goes into solution and urea precipitates. The mixture is stirred at room temperature for 3 hours, filtered and the filtrate washed with acetonitrile. The filtrate is treated dropwise with 8.5 ml (60 mM) diethylaminoethyl amine, whereby the temperature rises from 20° to 28°. The mixture is left 20 overnight and then is partitioned between 1 N aqueous sodium carbonate solution and CH₂ Cl₂. 20 The organic phase is evaporated to give the title compound, recrystallised from CH2 Cl2 / hexane m.p. 133-134°. Reference example for the preparation of the compound No. 141. 25 25 Indol-3-carbonyl-8-methyl-8-azabicyclo[3.2.1]oct-3β-yl-ane (a) 3-Chloro-8-methyl-8-azabicyclo[3.2.1]octane To a solution of 95 g pseudotropine in 420 ml abs. CHCl₃ are added dropwise at 0° 195 ml thionyl chloride within 20 minutes. The reaction mixture is refluxed for 4 hours, then left overnight at room temperature and then heated to 60-65° for 3 hours. The mixture is then 30 diluted with CH₂Cl₂ to a volume of ca. 100 to 150 ml and poured on ice-water. A 35% aqueous 30 NaOH solution is added to a pH 11 and then dry ice to a pH 10. After extraction with CH₂Cl₂ and distillation of the extracts (second fraction \sim 97-98°) the title compound is obtained. (b) 3-Cyano-8-methyl-8-azabicyclo[3.2.1]octane To a solution of 18.16 g KCN in 28 ml H_2O is added a solution of 42 g of the step (a) 35 35 compound in 90 ml ethanol and the mixture heated to 80°. Thereafter the mixture is refluxed for 22 hours. The mixture is evaporated to about 1/4 of its volume in a rotatory evaporator, then rendered alkaline with potassium carbonate and extracted with ether. Distillation of the residue yields at about 0.4 mm Hg and 85° the title compound. 40 40 (c) 3-Methoxycarbonyl-8-methyl-8-azabicyclo[3.2.1]octane To 30 g of the step (b) compound in 300 ml methanol and 3.7 ml water is introduced within 1 hour gaseous HCl, whereby the temperature rises to 60° (cooling). The mixture is left at room temperature for 18 hours. The resulting white suspension is filtered, the filtrate is concentrated, 45 rendered alkaline with potassium carbonate to a pH 10 and extracted with ether (3 times). The ether phase is washed with water and evaporated to give the title compound, as an oil, b.p. 72-74° / 0.13-0.15 mm Hg. (d) 3-Carboxy-8-methyl-8-azabicyclo[3.2.1]octane A solution of 25.2 g of the step (c) compound in 20 ml methanol is treated portionwise with 50 70 ml 2N aqueous NaOH solution within 30 minutes (pH at the end 13.2). The mixture is left at room temperature for 3 hours and then treated with the same amount of 2N HCl to a pH 5.8. The reaction mixture as chromatographed on about 300 ml amberlite TR 120 (H+form) using 10% NH₃ as eluant to give the title compound (recrystallised from ethanol/hexane), m.p. 55 222-224° (decomp.). 55 (e) 3-Chloro-carbonyl-8-methyl-8-azabicyclo[3.2.1]octane To a solution of 4.22 g. of the step (d) compound in 50 ml CH₂Cl₂ are added dropwise at about 15° 2.8 ml oxalyl chloride diluted with 5 ml CH2 Cl2. The resulting white suspension is 60 stirred 30 minutes at room temperature, diluted with 50 ml hexane, filtered and washed with 60 CH₂Cl₂ / hexane (1:2), to yield the hydrochloride of the title compound, decomp. from 205°. (f) Indol -3-carbonyl-8-methyl-8-azabicyclo[3.2.1]oct-3β-yl-ane To a Grignard reagent prepared from 1.44 g magnesium, 3.75 ml methyl iodide and 55 ml

65 abs. ether is added dropwise at boiling temperature a solution of 3.51g indole in 20 ml abs.

5	ether. The resulting silvergrey mixture is refluxed for 1 hour, then cooled to 0° and treated portionwise with 6.72 g of the hydrochloride of the step (e) compound, under slight exothermic reaction. A resin precipitates. The mixture is allowed to come to room temperature, whereby the resin solidifies. After leaving overnight water and CH_2Cl_2 are added. Stirring is effected until a white suspension results, which is extracted with CH_2Cl_2 (3 times). The aqueous phase is extracted with CH_2Cl_2 and 10 to 15% $\text{C}_2\text{H}_5\text{OH}$ (5 times). The evaporated extracts (about 5 g) are dissolved in $\text{CH}_2\text{Cl}_2+10\%$ CH $_3\text{OH}$ and filtered (residue about 2 g). The solution is chromato-	5
10	graphed on 250 g silicagel KG 004 using $CH_2CI_2+10\%$ C_2H_5OH as an eluant whereby the title compound is obtained (800 g). The residue and 800 mg of the compound obtained by chromatography are together recrystalized from H_2O/C_2H_5OH to give the hydrochloride of the title compound, m.p. 278–280° (decomp.). Reference example for the preparation of the compound No. 148	10
15	3 -(6-Hydroxyindolyl)-8-azabicyclo[3.2.1]-3β-methyl-ketone (a) 8-Benzyl-8-azabicyclo[3.2.1]octane-3β-acetic acid ethyl ester To 14 g 3-carbethoxy-methylen-8-benzyl-8-azabicyclo[3.2.1]octane in 300 ml aqueous NH $_3$ and 100 ml toluene at -40° are added 2.5 g sodium. The resulting blue mixture is decomposed after 5 minutes with solid NH $_4$ CL and the NH $_3$ is distilled off. After addition of water the mixture is extracted with CH $_2$ Cl $_2$ and chromatographed on silicium dioxide with ethyl acetate/hexane (1:8)	15
20	to yield the title compound as a colourless oil.	20
25	(b) 8-Azabicyclo[3.2.1]octane-3 β -acetic acid ethyl ester To a solution of 8.4 g of the step (a) compound in 350 ml C_2H_5OH are added 1 g Pd/C and the mixture is hydrogenated 4 hours. The mixture is filtered and evaporaacted to give the title compound as a colourless oil.	25
30	(c) 8-Benzyloxycarbonyl-8-azabicyclo[3.2.1]-3β-acetic acid ethyl ester 5.7 g of the step (b) compound, 100 ml toluene and 7.5 ml triethylamine are treated with 12.3 ml chloroformic acid benzyl ester. The mixture is heated 3 hours to 50°, then poured into 200 ml 0.1 N HCl and extracted 3 times with CH ₂ Cl ₂ . The organic phase is dried (Na ₂ SO ₄) and evaporated to give the title compound as a colourless oil.	30
35	(d) 8-Benzyloxycarbonyl-8-azabicyclo[3.2.1]-3β-acetic acid 9.1 g of the step (c) compound are dissolved in 60 ml ethanol, treated with 60 ml 2 N aqueous NaOH solution and refluxed 1 hour. After removal of ethanol by distillation, the remaining aqueous phase is acidified by addition of 10% tartaric acid and extracted with CH ₂ Cl ₂ . The combined organic phases are dried and evaporated to give the title compound as a yellowish foam.	35
40	(e) 8-Benzyloxycarbonyl-8-azabicyclo[3.2.1]-3β-acetic acid chloride A solution of 7.3 g of the step (d) compound in 60 ml CHCl ₃ is treated with 4.4 ml thionyl chloride and refluxed 2 hours. The solution is then treated several times with toluene and evaporated to give the title compound.	40
45	(f) 3-(6-Methoxyindolyl)-8-benzyloxycarbonyl-8-azabicyclo[3.2.1]-3β-methyl ketone Methyl magnesium iodide, prepared from 1.6 ml methyl iodide and 630 mg magnesium in 80 ml ether, is treated at room temperature with 1.5 g 6-methoxyindole in 70 ml ether. The mixture is heated 2 hours under reflux, then cooled to 0° and treated with 3.2 g of the step (e)	45
50	compound in 50 ml toluene. The reaction mixture is stirred 2 hours, poured into 2 N hydrochloric acid and extracted 3 times with CH_2CI_2 . The CH_2CI_2 -phases are washed with aqueous sodium bicarbonate solution and chromatographed on silicium dioxide eluting with ethyl acetate / hexane (1:10>1:1) to give the title compound as a colourless foam.	50
55	(g) 3 -(6 -Hydroxyindolyl)-8-azabicyclo[$3.2.1$]- 3β -methyl-ketone 300 mg of the step (f) compound in 30 ml CH $_2$ Cl $_2$ are treated at -78° with a solution of 0.8 ml boron tribromide in 10 ml CH $_2$ Cl $_2$. After stirring 1 hour at -78° and $2\frac{1}{2}$ hours at 0° aqueous sodium bicarbonate solution is added and the mixture extracted with n-butanol (3 times). The organic phases are evaporated and the residue is chromatographed on silica gel with CH $_2$ Cl $_2$ /	55
60	CH_3 OH / aq. NH ₃ (95:5:1>85:15:1) to yield the compound as colourless crystalls, m.p.>280°.	60
	CLAIMS 1. Use of a mono or bicyclic carbocyclic, or heterocyclic carboxylic, acid ester or amide, or an imidazolyl carbazol, in the manufacture of a medicament suitable for the treatment of stress-related psychiatric disorders, for increasing vigilance, for the treatment of rhinitis or serotonin-induced disorders or coadministration with another active agent to increase the bioavailability	65

	thereof, or for nasal administration.	
. 5	2. Use of a mono or bicyclic carbocyclic, or heterocyclic carboxylic, acid ester or amide, or an imidazolyl carbazol, for the treatment of stress-related psychiatric disorders, for increasing vigilance, for the treatment of rhinitis or serotonin-induced nasal disorders or coadministration with another active agent to increase the bioavailability thereof, or for nasal administration.	5
	3. Use of a serotonin 5-HT ₃ antagonist in the manufacture of a medicament suitable for the treatment of stress-related psychiatric disorders for increasing vigilance, for the treatment of rhinitis or serotonin-induced nasal disorders or coadministration with another active agent to increase the bioavailability thereof, or for nasal administration.	-
10	4. Use of a serotonin 5–HT ₃ antagonist for the treatment of stress-related psychiatric disorders, for increasing vigilance, for the treatment of rhinitis or serotinin-induced nasal disorders or coadministration with another active agent to increase the bioavailability thereof, or for nasal administration.	10
15	 5. Use according to any one of claims 1 to 4 for stress-related psychiatric disorders. 6. Use according to any one of claims 1 to 4 for serotonin-induced nasal disorders. 7. Use according to any one of claims 1 to 4 for increasing vigilance. 8. Use according to any one of claims 1 to 4 for the treatment of rhinitis. 	15
20	 9. Use according to any one of claims 1 to 4 for the treatment of impaired approach oriented behaviour in stressful situations. 10. Use according to any one of claims 1 to 4 for co-administration with another active 	20
	agent to increase bioavailability thereof. 11. Use according to any one of claims 1 to 4 for nasal administration. 12. Use according to any one of claims 1 to 4 for nasal coadministration with another active	
25	agent. 13. A nasal composition comprising a mono or bicyclic carbocyclic, or heterocyclic carboxylic, acid ester or amide or an imidazolyl carbazol.	25
30	 14. A nasal composition comprising a serotonin 5–HT₃ antagonist. 15. A nasal composition according to claim 13 or 14 containing another active agent. 16. Use of a mono or bicyclic carbocylic or heterocyclic carbocylic acid ester or amide other than a compound of formula I wherein A is 3,5-dichloro-phenyl, B is CO, C is O, D is tropanyl in the manufacture of a medicament suitable for the treatment of lung embolism. 17. Use of a mono or bicyclic carbocyclic or heterocyclic carbocylic acid ester or amide other than a compound of formula I wherein A is 3,5-dichloro-phenyl, B is CO, C is O, D is tropanyl 	30
35	for the treatment of lung embolism. 18. Use or composition as defined in any one preceding claim wherein the compound is a mono or bicyclic carbocylic, or a heterocyclic carboxylic acid, ester, or amide, of a cyclic alcohol, or amine, containing nitrogen as a ring atom.	35
	19. Use or composition according to claim 18 wherein the compound is a heterocyclic compound.	
40	20. Use of composition according to claim 18 wherein the compound is a bicyclic heterocyclic compound.	40
	21. Use or composition according to any one preceding claim wherein the ester or amide or serotonin $5-HT_3$ antagonist is a compound of formula I.	-
45	A-B-C-D (I) wherein A is a group of formula	45

X-Y is -CH=CH-, $-O=CH_2-$ or -N=CH-,

Z is $-CH_2-$, $-NR_3-$, -O- or -S-,

R₁ and R₂ are independently hydrogen, halogen, (C_{1-4}) alkyl, (C_{1-4}) alkoxy, hydroxy, amino, (C_{1-4}) alkylamino, $di(C_{1-4})$ alkylamino, mercapto or (C_{1-4}) alkylthio,

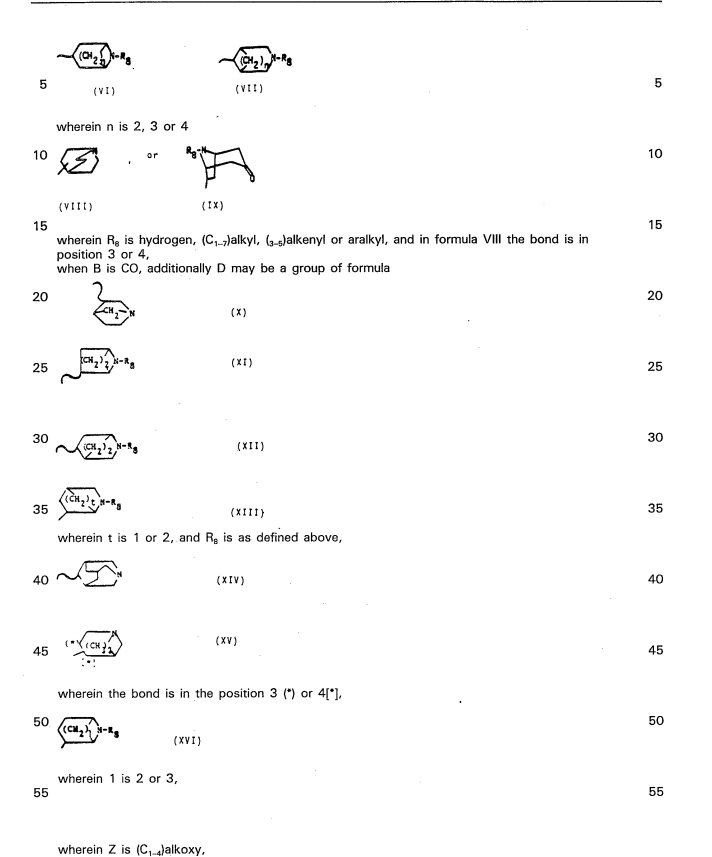
R₃ is hydrogen, (C_{1-4}) alkyl, acyl (C_{3-5}) alkenyl, aryl or arylalkyl, and

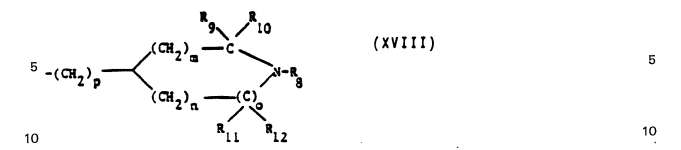
R₄ to R₇ are, independently, hydrogen, amino, nitro, (C_{1-4}) alkylamino, $di(C_{1-4})$ alkylamino, halo
55 gen, (C_{1-4}) alkoxy, (C_{1-4}) alkyl, (C_{1-4}) alkanoylamino, pyrrolyl, sulfamoyl, or carbamoyl

55 B is -CO- or $-SO_2-$,

C is -O- or -NH- or a bond

D is a group of formula





wherein $R_{\rm 9}$ to $R_{\rm 12}$ are independently hydrogen or (C $_{\rm 1-4}$)alkyl, m is 0, 1 or 2 and

n, o, p independently are 0 or 1,

15

$$_{20}$$
 -(CH₂)_q -N $_{R_{14}}$ (XIX)

wherein q is 2 or 3,

 R_{13} and R_{14} independently are (C_{1-4})alkyl,

25



30 wherein the bond is in position 3 or 4,

H....

(XXI)

5

5

10 H₃C CH₂ N-R₃

(XXII)

10

15

15 H₃C N-R₈

(XXIII)

20

20

$$\sim$$
N $(CH_2)_3$ N-R₈

(XXIVa)

25

25

(XXIVb)

30

35



(XXY)

30

40

and R₈ is as defind above in free base form, acid addition salt form or quaternary ammonium salt form, or a compound of formula la

5 (Ia)

10

15

5

wherein R_{15} is hydrogen, (C_{1-10}) alkyl, (C_{3-9}) cycloalkyl, (C_{3-6}) alkenyl, phenyl or phenyl (C_{1-3}) alkyl and one of the groups R_{16} , R_{17} and R_{18} is hydrogen, (C_{1-6}) alkyl, (C_{3-7}) cycloalkyl, (C_{2-6}) alkenyl or phenyl(C_{1-3})alkyl, and the others independently are hydrogen, or (C_{1-4})alkyl.

22. Use or composition according to claim 21 wherein the compound is of formula I wherein 15 A is chosen from formula II, III, IV and V, R_3 is other than acyl C is -O- or -NH- and D is chosen from formula VI to XVIII, with the proviso that, when A is formula III, B is CO and C is NH, D is not a group of formula VI, in free base, in acid addition salt form or in quaternary ammonium salt form.

23. Use or composition according to claim 21 for the promotion of approach-oriented social 20 behaviour in stressful situations or increasing vigilance.

20 24. Use or composition according to any one of claims 1 to 18 wherein the ester or amide or serotonin 5-HT3 antagonist is a dicarbocyclic or heterocyclic carboxylic acid ester, or carboxylic acid amide, of a piperidinol containing an alkylene bridge or of piperidylamine containing an

alkylene bridge or of an ester or amide of a substituted benzoic acid and of a piperdidinol, or 25 piperidylamine, containing an alkylene bridge, with the proviso that in each benzoic acid amide the alkylene bridge of the piperidyl ring is bonded to the nitrogen atom and to a cyclic carbon

25. Use or composition according to claim 21 wherein the compound is of formula I has a group of formula II'

30

25

35 wherein the free bond may be situated in any of the rings,

35

X' is $-CH_2-$, $-NR_3-$, -O-, -S-, R'_1 and R'_2 independently of one another are hydrogen, halogen, (C_{1-4}) alkyl, (C_{1-4}) alkoxy, hydroxy, amino, (C_{1-4}) alkylamino, di (C_{1-4}) alkylamino, mercapto or (C_{1-4}) alkylthio, and R_3' is hydrogen, (C_{1-4}) alkyl, (C_{3-5}) alkenyl, aryl or aralkyl, or a group of formula III'

40

45

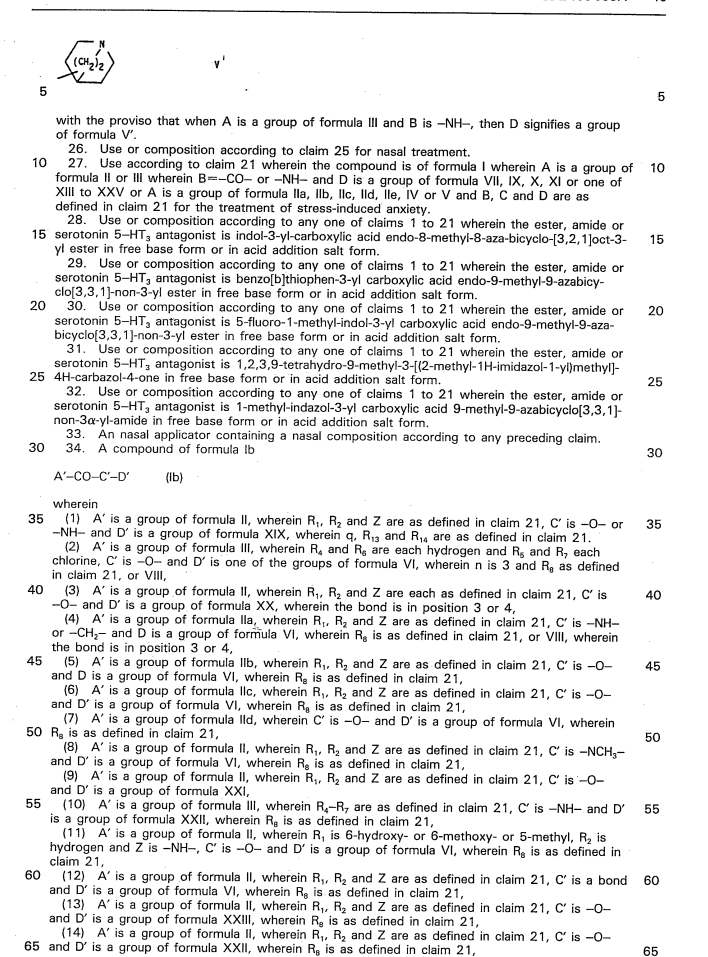
 R'_4 to $R'_{7'}$ independently of one another, are hydrogen, amino, nitro, (C_{1-4}) alkylamino, di (C_{1-4}) lamino, halogen, (C₁₋₄alkoxy, (C₁₋₄alkyl, (C₁₋₄)alkanoylamino or pyrrolyl,

B signifies -O- or -NH-, 50 D signifies a group of formula IV', 50

55

wherein

n is 2, 3 or 4 and R_8' is hydrogen, (C_{1-7}) alkyl, (C_{3-5}) alkenyl or aralkyl, or a group of formula V',



	(15) A' is a group of formula IIe, wherein Z is as defined in claim 21, C' is $-0-$ and D' is a group of formula VI, wherein R_8 is as defined in claim 21,	
	(16) A' is a group of formula II, wherein R_1 , R_2 and Z are as defined in claim 21, C' is a bond and D' is a group of formula XXIVa, wherein R_0 is as defined in claim 21,	5
5	(17) A' is a group of formula II, wherein R_1 , R_2 and Z are as defined in claim 21, C' is a bond and D' is a group of formula (XXIVb), wherein R_8 is as defined in claim 21. (18) A' is a group of formula II, wherein R_1 and R_2 are as defined in claim 21, Z is N-acyl, C'	J
	is $-O-$ and D' is a group of formula VI, wherein R_8 is as defined in claim 21, (19) A' is a group of formula II, wherein R_1 hydroxy, R_2 is hydrogen and Z is $-NH-$, C' is	
10	$-CH_2-$ and D' is a group of formula VI, wherein R ₈ is as defined in claim 21. in free base form, in acid addition salt form or quaternary ammonium salt form.	10
	35. A compound of claim 34 which is:— N-[2-(N,N-diethylamino)ethyl]indolyl-2-carboxamide N-[2-(N,N-diethylamino)ethyl]indolyl-3-carboxamide	
15	N-[2-(N,N-diethylamino)ethyl]indolyl-4-carboxamide N-[2-(N,N-diethylamino)ethyl]indolyl-6-carboxamide	15
	N-[2-(N,N-diethylamino)ethyl]indolyl-6-carboxamide 5-indolylcarboxylic acid-[2-(N,N-diethylamino)ethyl]ester N-[2-(N,N-diethylamino)ethyl]-3-bromoindolyl-5-carboxamide	
20	N-[2-(N,N-diethylamino)ethyl]-3-iodoindolyl-5-carboxamide N-[2-(N,N-diethylamino)ethyl]-3-methylindolyl-5-carboxamide	20
	N-[2-(N,N-diethylamino)ethyl]-1-methylindolyl-5-carboxamide 3,5-dichloro-benzoic acid-9-methyl-9-aza-bicyclo[3,3,1]non-3-yl ester 3,5-dichloro-benzoic acid-azabicyclo[2,2,2]oct-3-yl ester	
25	(2R*,3S*)-1H-indolyl-3-carboxylic acid 2-methyl-1-aza-bicyclo[2,2,2]oct-3-yl ester (2S*,3S*)-1-indolyl-3-carboxylic acid 2-methyl-1-azabicyclo[2,2,2]oct-3-yl ester	25
	3RS-(2,3-dihydro-1H-indol-5-yl carboxylic acid) 1-azabicyclo[2,2,2]oct-3-yl amide 2-methyl-indazol-3-carboxylic acid 8-methyl-8-azabicyclo[3,2,1]oct-3α-yl ester 1-methyl-indazol-3-carboxylic acid 8-methyl-8-azabicyclo[3,2,1]oct-3α-yl ester	
30	Thiophenyl-2-carboxylic acid 9-methyl-9-azabicyclo[3,3,1]non-3 α -yl ester Thiophenyl-3-carboxylic acid 9-methyl-9-azabicyclo[3,3,1]non-3 α -yl ester	.30
	1-acetyl-1H-indolyl-3-carboxylic acid-8-methyl-8-azabicyclo[3,2,1]oct-3α-yl ester 1H-indolyl-3-carboxylic acid 9-methyl-9-azabicyclo[3,3,1]nonan-3-yl methyl amide 9r-benzthiophenyl-3-carboxylic acid-3-aza-adamanthan-9-yl ester	
35	9s-benzthiophenyl-3-carboxylic acid-3-aza-adamantan-9-yl ester 4-amino-5-chloro-2-methoxy benzoic acid 7-dimethyl-(1α H,3 α ,5H)-aza-bicyclo[3,3,1]non-3-yl am-	35
	ide 6-methoxy-1H-indolyl-3-carboxylic acid-(1αH,5αH)-8-methyl-8-azabicyclo[3,2,1]oct-3-yl ester	
40	6-hydroxy-1H-indolyl-3-carboxylic acid-(1αH,5αH)-8-methyl-8-azabicyclo[3,2,1]oct-3-yl ester Indolyl-3-carbonyl-8-methyl-8-azabicyclo[3,2,1]oct-3β-yl-ane	40
	1H-indolyl-3-carboxylic acid 6,6,9-trimethyl-9-azabicyclo[3,2,1]non-3-yl ester 1H-indolyl-3-carboxylic acid 7,7,9-trimethyl-9-azabicyclo[3,3,1]non-3-yl ester 5-methoxy-6-fluor-indolyl-3-carboxylic acid 8-methyl-8-azabicyclo[3,2,1]oct-3-yl ester	
45	1H-indolyl-3-carboxylic acid 3-methyl-3,9-diazabicyclo[3,3,1]non-9-yl ester Indolyl-3-carboxylic acid 9-methyl-3,9-diazabicyclo[3,3,1]non-3-yl amide 5-methyl-indolyl-3-carboxylic acid $(1\alpha H,5\alpha H)$ -8-methyl-8-azabicyclo[3,2,1]oct-3 α -yl ester	45
	3-(6-hydrooxyindolyl)-8-azabicyclo[3,2,1]-3 β -methylketone, or 3-(6-methoxy-2,3-dihydroindolyl)-8-azabicyclo[3,2,1]-3 β -methylketone	
50	in free base form, in acid addition salt form or in a quaternary ammonium salt form. 36. A process for the production of a compound of formula lb A'-B'-C'-D' as defined in claim 34 in free base form or in acid addition salt form or in quaternary ammonium salt form,	50
	which includes the step of (a) for the production of a compound of formula lb wherein C' is -O- or -NH-, reacting a compound of formula XXX	-
55	·	55
	wherein A' is as defined above, or a reactive derivative thereof,	_
60		60
	H–C'–D' (XXXI)	

or a precursor of this compound, or (b) for the production of a compound of formula lb wherein C' is -CH₂-, reacting a compound of formula XXXII 5 A'-Mg-Hal (XXXII) 5 wherein A' is as defined above, and Hal is chlorine, bromine or iodine, with an appropriate compound of formula XXXIII 10 10 CIOC-CH2-D' (XXXIII) wherein D' is as defined above, in free base form or protected form, under conditions of a Grignard reaction, and removing any protected group present, (c) for the production of a compound of formula lb wherein C' is a direct bond reacting a 15 compound of formula XXXII as defined above, with an appropriate compound of formula XXXIV CI-OC-D' (XXXIV 20 wherein D' is as defined above, 20 under conditions of a Grignard reaction, (d) for the production of a compound of formula lb wherein C' is a -NCH₃- group, reacting a compound of formula XXX as defined above, with an appropriate compound of formula XXXI wherein C' is -NCH3-, 25 25 (e) for the production of a compound of formula lb wherein Z is N-acyl- wherein an appropriate compound of formula I wherein A is a compound of formula II and Z is -NH-, B is CO, C is -O- or -NH- and D is as defined in claim 3 is acylated and recovering the compound of formula lb in free base form, acid addition salt form or quaternary ammonium salt form. 37. Use of a compound of claim 21 wherein (i) A is a group of formula II or III and B is 30 -CO-, C is -O- or -NH- and D is a group of formula VII, IX, X, XI and one of XIII to XXV and (ii) A is a group of formula IIa, IIb, IIc, IId, IIe, IV or V in the manufacture of a medicament suitable for use as an anxiolytic. 38. Use of a compound of formula I wherein (i) A is a group of formula II or III and B is 35 35 -CO-, C is -O- or -NH- and D is a group of formula VII, IX, X, XI and one of XIII to XXV, and (ii) A is a group of formula IIa, IIb, IIc, IId, IIe, IV or V as an anxiolytic. 39. A process for the production of a pharmaceutical composition adapted for the treatment of psychiatric disorders for increasing vigilance, for the treatment of rhinitis or for serotonininduced nasal disorders or lung embolism which comprises working up a mono or bicyclic 40 40 carboxylic, or heterocyclic carboxylic acid or ester or amide or an imidazolyl carbazol, with pharmaceutical carriers and diluents in the manufacture of unit dosage formulations of said indications. 40. A process for the production of a pharmaceutical composition having improved bioavailability which comprises working up a mono or bicyclic carboxylic or heterocyclic carboxylic acid 45 ester or amide or an imidazolyl carbazol, e.g. a compound of formula I or la as defined hereinbefore with another active agent, e.g. a peptide and if desired formulating as a unit dosage form. 41. A process for the production of a nasal composition which comprises working up a mono or bicyclic carboxylic or heterocyclic acid ester or amide or an imidazolyl carbazol, e.g. a 50 50 compound of formula I or la as defined hereinbefore, with an appropriate nasal carrier, and optionally incorporating a surface active agent and optionally filling the resultant composition into a nasal applicator. 42. A composition or use substantially as hereinbefore described with reference to any one of the examples. 55 55 43. Use of mono or bicyclic carboxylic or heterocyclic carboxylic acid esters or amides of a cyclic alcohol, or cyclic amine, containing nitrogen as a ring atom, in free base form or in acid addition or quaternary ammonium salt form as a 5-HT₃ antagonists in the manufacture of a medicament suitable for the treatment of serotonin-induced psychiatric disorders. 44. Use according to claim 43 wherein the psychiatric disorders are chosen from one of the 60 60 following: anxiety, social withdrawal, affective disorders, psychoses and other stress-related illnesses, disorders, of vigilance, e.g. geriatric illnesses.