



HU000026619T2

(19) **HU****MAGYARORSZÁG**
Szellemi Tulajdon Nemzeti Hivatala(11) Lajstromszám: **E 026 619**(13) **T2**

EURÓPAI SZABADALOM

SZÖVEGÉNEK FORDÍTÁSA

(21) Magyar ügyszám: **E 11 730199**(22) A bejelentés napja: **2011. 06. 14.**(51) Int. Cl.: **A61K 311/37** (2006.01)**A61K 45/06** (2006.01)**A61P 29/00** (2006.01)

(96) Az európai bejelentés bejelentési száma:

EP 20110730199

(97) Az európai bejelentés közzétételi adatai:

EP 2582366 A1 **2011. 12. 22.**

(86) A nemzetközi (PCT) bejelentési szám:

PCT/EP 11/002905

(97) Az európai szabadalom megadásának meghirdetési adatai:

EP 2582366 B1 **2015. 10. 28.**

(87) A nemzetközi közzétételi szám:

WO 11157391

(30) Elsőbbségi adatok:

10006202 **2010. 06. 15.** **EP**

(73) Jogosult(ak):

Grünenthal GmbH, 52078 Aachen (DE)

(72) Feltaláló(k):

SCHIENE, Klaus, 41363 Jüchen (DE)
BLOMS-FUNKE, Petra, 52146 Würselen (DE)

(74) Képviselő:

Dr. Miskolczi Mária, Dr. Jalsovszky Györgyné,
Budapest

(54)

Gyógyászati kombináció fájdalom kezelésére

Az európai szabadalom ellen, megadásának az Európai Szabadalmi Közlönyben való meghirdetésétől számított kilenc hónapon belül, felszólalást lehet benyújtani az Európai Szabadalmi Hivatalnál. (Európai Szabadalmi Egyezmény 99. cikk(1))

A fordítást a szabadalmas az 1995. évi XXXIII. törvény 84/H. §-a szerint nyújtotta be. A fordítás tartalmi helyességét a Szellemi Tulajdon Nemzeti Hivatala nem vizsgálta.

(19)



(11)

EP 2 582 366 B1

(12)

EUROPEAN PATENT SPECIFICATION

(45) Date of publication and mention
of the grant of the patent:
28.10.2015 Bulletin 2015/44

(51) Int Cl.:
A61K 31/137 ^(2006.01) **A61K 45/06** ^(2006.01)
A61P 29/00 ^(2006.01)

(21) Application number: **11730199.4**

(86) International application number:
PCT/EP2011/002905

(22) Date of filing: **14.06.2011**

(87) International publication number:
WO 2011/157391 (22.12.2011 Gazette 2011/51)

(54) **PHARMACEUTICAL COMBINATION FOR THE TREATMENT OF PAIN**

PHARMAZEUTISCHE ZUSAMMENSETZUNGEN ZUR SCHMERZBEHANDLUNG

COMBINAISON PHARMACEUTIQUE POUR LE TRAITEMENT DE LA DOULEUR

(84) Designated Contracting States:
**AL AT BE BG CH CY CZ DE DK EE ES FI FR GB
GR HR HU IE IS IT LI LT LU LV MC MK MT NL NO
PL PT RO RS SE SI SK SM TR**

(30) Priority: **15.06.2010 EP 10006202**

(43) Date of publication of application:
24.04.2013 Bulletin 2013/17

(73) Proprietor: **Grünenthal GmbH
52078 Aachen (DE)**

(72) Inventors:
• **SCHIENE, Klaus**
41363 Jüchen (DE)
• **BLOMS-FUNKE, Petra**
52146 Würselen (DE)

(56) References cited:
WO-A1-03/053427 WO-A2-2009/094563
WO-A2-2010/025931 DE-A1- 4 426 245

Note: Within nine months of the publication of the mention of the grant of the European patent in the European Patent Bulletin, any person may give notice to the European Patent Office of opposition to that patent, in accordance with the Implementing Regulations. Notice of opposition shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).

EP 2 582 366 B1

Description

[0001] The present invention relates to a combination comprising as components (a) at least one 3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol compound, and (b) at least one NMDA-antagonist, a pharmaceutical formulation and a dosage form comprising said combination as well as a method of treating pain, e.g. Inflammatory pain or neuropathic pain, wherein components (a) and (b) are administered simultaneously or sequentially to a mammal, whereby component (a) may be administered before or after component (b) and whereby components (a) or (b) are administered to the mammal either via the same or a different pathway of administration.

[0002] The treatment of chronic and acute pain conditions is extremely important in medicine. There is currently a worldwide demand for additional, not exclusively opioid-based, but highly effective pain treatment. The urgent need for action for patient-oriented and purposeful treatment of pain conditions, this being taken to mean the successful and satisfactory treatment of pain for the patient, is documented in the large number of scientific papers that have recently appeared in the field of applied analgesics and fundamental research work on nociception.

[0003] Even if the analgesics that are currently used for treating pain, for example opioids, NA- and 5HT-reuptake inhibitors, NSAIDs and COX inhibitors, are analgesically effective, side effects nevertheless sometimes occur. Sometimes substance combinations comprising two different drugs show super-additive therapeutic effects upon administration. Due to such a super-additive effect the overall dose and accordingly the risk of undesired side effects can be reduced. Examples of combinations comprising 3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol are disclosed in WO2007/128412, WO2007/128413 and WO2010/025931.

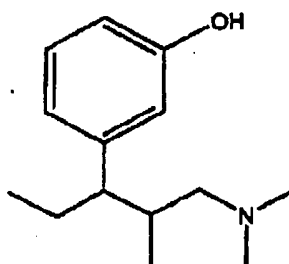
[0004] WO2003053427 describes the use of (+)-(1S,2S)-3-(3-dimethylamino-1-ethyl-2-methyl-propyl)phenol for preparing an anti-emetic agent. DE 44 26 245 relates to 1-phenyl-3-dimethylaminopropane compounds, to a method of preparing them, and to the use of these substances as pharmaceutical active ingredients. WO 2009094563 discloses crystalline form of a (3S)-aminomethyl-5-hexanoic acid prodrug.

[0005] Thus, it was an object of the present invention to find further combinations having improved properties. It was also an object of the present invention to find further combinations that are suitable for the treatment of pain and that preferably exhibit fewer undesired side effects compared to its individual components, if administered in effective doses.

[0006] It has been found that a combination comprising (a) at least one 3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol compound, and (b) at least one NMDA-antagonist selected from the group consisting of (R,S)-ketamine, (S)-ketamine and memantine exhibits an analgesic effect. If these components are present in the composition in such a weight ratio that a super-additive or synergistic therapeutic effect is observed upon administration to the patients, the overall administered dose may be lowered, so that fewer undesired side-effects will occur.

[0007] Accordingly, the present invention relates to a pharmaceutical combination comprising as components

(a) 3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol of formula (I)



(I),

optionally in form of one of its pure stereoisomers, in particular an enantiomer or a diastereomer, a racemate or in form of a mixture of its stereoisomers, in particular enantiomers and/or diastereomers in any mixing ratio, or any corresponding acid addition salt thereof, and

(b) at least one NMDA-antagonist selected from the group consisting of (R,S)-ketamine, (S)-ketamine and memantine.

[0008] In one embodiment of the inventive combination the compound of formula (I) is selected from

(1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol,
(1S,2S)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol,

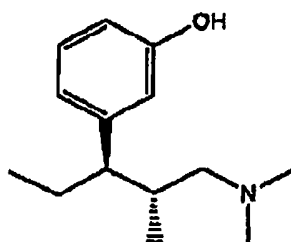
(1R,2S)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol,
(1S,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, and any mixture thereof.

[0009] In another embodiment of the inventive combination the compound of formula (I) is selected from

(1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, and
(1S,2S)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, and any mixture thereof.

[0010] In yet another embodiment the inventive combination comprises

(a) the compound (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol of formula (I'),



(I')

or an acid addition salt thereof, and

(b) at least one NMDA-antagonist selected from the group consisting of (R,S)-ketamine, (S)-ketamine and memantine.

[0011] The compound 3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol of formula (I), its stereoisomers and corresponding salts thereof as well as methods for their preparation are well known, for example, from US 8,248,737 B1.

[0012] The definition of component (a) as used herein includes 3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, derivatives thereof and stereoisomers thereof in any possible form, thereby particularly including solvates and polymorphs, salts, in particular acid addition salts and corresponding solvates and polymorphs.

[0013] The term derivative as used herein particularly includes prodrugs such as ethers and esters of the active substance. Suitable methods for selecting and preparing a prodrug of a given substance are for example described in "Textbook of Drug Design and Discovery", 3rd edition, 2002, chapter 14, pages 410-458, Editors: Krogsgaard-Larsen et al., Taylor and Francis.

[0014] If component (a) is present as mixture of enantiomers, such a mixture may contain the enantiomers in racemic or non-racemic form. A non-racemic form could, for example, contain the enantiomers in a ratio of 60±5:40±5, 70±6:30±5, 80±5:20±5 or 90±5:10±5.

[0015] The compound 3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol and its stereoisomers according to component (a) may be present in the inventive pharmaceutical composition in form of an acid addition salt, whereby any suitable acid capable of forming such an addition salt may be used.

[0016] The conversion of 3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol into a corresponding addition salt, for example, via reaction with a suitable acid may be effected in a manner well known to those skilled in the art. Suitable acids include but are not limited to hydrochloric acid, hydrobromic acid, sulfuric acid, methanesulfonic acid, formic acid, acetic acid, oxalic acid, succinic acid, tartaric acid, mandelic acid, fumaric acid, lactic acid, citric acid, glutamic acid and/or aspartic acid. Salt formation is preferably effected in a solvent, for example, diethyl ether, diisopropyl ether, alkyl acetates, acetone and/or 2-butanone. Moreover, trimethylchlorosilane in aqueous solution is also suitable for the preparation of hydrochlorides.

[0017] The N-methyl-D-aspartate (NMDA) receptor is part of a glutamatergic neurotransmitter-receptor system, the so-called NMDA-receptor/ion-channel complex. It comprises different binding sites that are located inside and outside of the ion-channel. An NMDA receptor antagonist or simply NMDA-antagonist is a substance that interacts with such a binding site and exerts at least partially inhibiting properties related to said binding site.

[0018] NMDA-antagonists are well known to those skilled in the art and particularly include, without being limited thereto, N-containing phosphonic acids, such as norvaline (AP5), D-norvaline (D-AP5), 4-(3-phosphono-propyl)-piper-

azine-2- carboxylic acid (CPP), D-(E)-4-(3-phosphonoprop-2-enyl)piperazine-2-carboxylic acid (D-CPPene), cis-4-(phosphonomethyl)-2-piperidine carboxylic acid (Selfotel, CGS 19755), SDZ-220581, PD-134705, LY-274614 and WAY-126090; quinolinic acids, such as kynurenic acid, 7-chloro-kynurenic acid, 7-chloro-thiokynurenic acid and 5,7-dichloro-kynurenic acid, prodrugs thereof, such as 4-chlorokynurenine and 3-hydroxy-kynurenine; 4-aminotetrahydroquinolin-carboxylates, such as L-689,560; 4-hydroxyquinolin-2(1H)-ones, such as L-701,324; quinoxalinediones, such as licostinel (ACEA-1021) and CGP-68,730A; 4,6-dichloro-indole-2-carboxylate derivatives such as MDL-105,519, gavestinel (GV-150,526) and GV-196,771A; tricyclic compounds, such as ZD-9,379 and MRZ-2/576, (+)-HA-966, morphinan derivatives such as dextromethorphan and dextrophan; benzomorphans, such as BIII-277CL; other opioids, such as dextropropoxyphene, ketobemidone, dextromethadone and D-morphine; amino-adamantanes, such as amantadine and memantine; amino-alkyl-cyclohexanes, such as MRZ-2/579; Ifenprodil and Ifenprodil-like compounds such as eliprodil and PD-196.860; Iminopyrimidines; or other NMDA-antagonists such as nitroprusside, D-cycloserine, 1-aminocyclopropane-carboxylic acid, dizocilpine (MK 801) and its analogs, phencyclidine (PCP), ketamine ((R,S)-2-(2-Chlorophenyl)-2-(methylamino)cyclohexan-1-on), (R)-ketamine, (3)-kelamine, remacemide and its des-glycyl-metabolite FPL-12,495, AR-R-15,896, methadone, sulfazocine, AN19/AVex-144, AN2/AVex-73, Besonprodil, CGx-1007, EAB-318, Felbamate and NPS-1407. NMDA-Antagonists are, for example, disclosed in "Analgesics. Edited by H. Buschmann. T. Christoph, E. Friderichs, C. Maul, B. Sundermann, 2002, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim. Germany, In particular pages 389-428.

[0019] Some NMDA-antagonists such as ketamine and memantine are known to be useful in the treatment of neuropathic pain. In one embodiment of the present invention one or more of these NMDA-antagonists is used as component (b).

[0020] Also included are steroisomers, salts, solvates, polymorphs and derivatives of the NMDA-antagonist component as well as mixtures of any of the foregoing.

[0021] In another embodiment of the inventive combination the NMDA-antagonist according to component (b) is memantine or an acid addition salt thereof such as the hydrochloride addition salt.

[0022] In a further embodiment of the inventive combination the NMDA-antagonist according to component (b) is ketamine or an acid addition salt thereof such as the hydrochloride addition salt.

[0023] In yet another embodiment of the inventive combination the NMDA-antagonist according to component (b) is (R,S)-ketamine or an acid addition salt thereof such as the hydrochloride addition salt.

[0024] In still another embodiment of the inventive combination the NMDA-antagonist according to component (b) is (S)-ketamine or an acid addition salt thereof such as the hydrochloride addition salt.

[0025] Another specific embodiment of the present invention is a combination comprising (a) (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, or an acid addition salt thereof such as the hydrochloride addition salt, and (b) memantine or an acid addition salt thereof such as the hydrochloride addition salt.

[0026] Yet another specific embodiment of the present invention is a combination comprising (a) (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, or an acid addition salt thereof such as the hydrochloride addition salt, and (b) ketamine or an acid addition salt thereof such as the hydrochloride addition salt.

[0027] Yet a further specific embodiment of the present invention is a combination comprising (a) (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, or an acid addition salt thereof such as the hydrochloride addition salt, and (b) (R,S)-ketamine or an acid addition salt thereof such as the hydrochloride addition salt.

[0028] Yet still another specific embodiment of the present invention is a combination comprising (a) (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, or an acid addition salt thereof such as the hydrochloride addition salt, and (b) (S)-ketamine or an acid addition salt thereof such as the hydrochloride addition salt.

[0029] Some NMDA-antagonists comprise functional groups, for example, acidic groups such as carboxy groups which are capable of forming salts with the 3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol component of formula (I), thereby incorporating both components (a) and (b) in one and the same salt.

[0030] Thus, in another embodiment of the present invention the inventive combination comprises components (a) and (b) in form of a salt formed from these two components. Such a salt formation may be partial, i.e. the inventive composition comprises one or both of these components also in their non-salt form, or the salt formation may essentially be complete.

[0031] Both components (a) and (b) as part of the inventive combination may be administered in amounts up to their maximum daily dosage, which is known to those skilled in the art.

[0032] Memantine may preferably be administered to a patient in a daily dosage of 1 to 20 mg.

[0033] Racemic ketamine may preferably be administered to a patient in a daily dosage of 0.1 to 10 mg/kg, the S-enantiomer of ketamine may preferably be administered to a patient in a daily dosage of 1 to 10 mg/kg.

[0034] The compound (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol may preferably be administered to a patient in a daily dosage of 25 to 1000 mg, particularly preferably in a dosage of 50 to 800 mg, more particularly preferably in a dosage of 100 to 600 mg.

[0035] When administered as part of the inventive combination the administered amount per day of component (a) and/or component (b) may be less than the respective maximum daily dosage and be, for example, 75 ± 15 wt.-%, 75 ± 10

wt.-%, 75±5 wt.-%, 50±15 wt.-%, 50±10 wt.-%, 50±5 wt.-%, 25±15 wt.-%, 25±10 wt.-% and 25±5 wt.-% for each of the components.

[0036] In another embodiment of the present invention the inventive combination may contain components (a) and (b) essentially in an equieffective ratio.

[0037] In yet a further embodiment of the inventive combination components (a) and (b) are present in such a weight ratio that the resulting composition will exert a supra-additive or synergistic effect upon administration to a patient. Suitable weight ratios can be determined by methods well known to those skilled in the art.

[0038] Both components (a) and (b) may also be present in the inventive combination in ratios deviating from the equieffective ratio. For, example, each of the components could be present in a range from 1/50 of the equieffective amount to 50 times the equieffective amount, from 1/20 of the equieffective amount to 20 times the equieffective amount, from 1/10 of the equieffective amount to 10 times the equieffective amount, from 1/5 of the equieffective amount to 6 times the equieffective amount, from 1/4 of the equieffective amount to 4 times the equieffective amount, from 1/3 of the equieffective amount to 3 times the equieffective amount, or from 1/2 of the equieffective amount to 2 times the equieffective amount.

[0039] In another embodiment of the present invention the components (a) and (b) can be : administered in a specific dosage regimen to treat pain, for example, neuropathic pain. Components (a) and (b) may be administered simultaneously or sequentially to one another. In each case via the same or different administration pathways.

[0040] The term pain as used herein includes but is not limited to inflammatory pain, neuropathic pain, acute pain, chronic pain, visceral pain, migraine pain and cancer pain.

[0041] Suitable pathways of administration include but are not limited to oral. Intravenous, intraarterial, intraperitoneal, untradermal, transdermal, intrathecal, intramuscular, intranasal, transmucosal, subcutaneous, and rectal administration.

[0042] The inventive combinations are toxicologically safe and are therefore suitable for the treatment of mammals, particularly humans including infants, children and grown-ups.

[0043] Thus, in a further aspect the present invention relates to a pharmaceutical composition comprising an inventive combination as described herein and one or more auxiliary agents.

[0044] In a further aspect the present invention relates to a pharmaceutical dosage form comprising an inventive combination as described herein and one or more auxiliary agents.

[0045] In one embodiment, the inventive pharmaceutical dosage form is suitable for being administered orally, intravenously, intraarterially, intraperitoneally, intradermally, transdermally, intrathecally, intramuscularly, intranasally, transmucosally, subcutaneously, or rectally.

[0046] The inventive formulations and dosage forms may contain auxiliary agents, for example, carriers, fillers, solvents, diluents, colorants and/or binders. The selection of auxiliary agents and of the amounts of the same to be used depends, for example, on how the drug is to be administered, e.g. orally, intravenously, intraarterially, intraperitoneally, intradermally, transdermally, intramuscularly, intranasally or locally, for example for infections of the skin, of the mucous membranes or of the eye.

[0047] Suitable auxiliary agents in the context of this invention are in particular any substances known to a person skilled in the art useful for the preparation of galenical formulations. Examples of suitable auxiliary agents include but are not limited to: water, ethanol, 2-propanol, glycerol, ethylene glycol, propylene glycol, polyethylene glycol, polypropylene glycol, glucose, fructose, lactose, saccharose, dextrose, molasses, starch, modified starch, gelatine, sorbitol, inositol, mannitol, microcrystalline cellulose, methyl cellulose, carboxymethyl cellulose, cellulose acetate, shellac, cetyl alcohol, polyvinyl pyrrolidone, paraffins, waxes, natural and synthetic gums, acacia gum, alginates, dextran, saturated and unsaturated fatty acids, stearic acid, magnesium stearate, zinc stearate, glycerol stearate, sodium lauryl sulphate, edible oils, sesame oil, coconut oil, peanut oil, soybean oil, lecithin, sodium lactate, polyoxyethylene and polypropylene fatty acid ester, sorbitan fatty acid ester, sorbic acid, benzoic acid, citric acid, ascorbic acid, tannic acid, sodium chloride, potassium chloride, magnesium chloride, calcium chloride, magnesium oxide, zinc oxide, silicon dioxide, titanium oxide, titanium dioxide, magnesium sulphate, zinc sulphate, calcium sulphate, potash, calcium phosphate, dicalcium phosphate, potassium bromide, potassium iodide, talcum, kaolin, pectin, crosspovidone, agar and bentonite.

[0048] Pharmaceutical formulations (dosage forms) in the form of tablets, effervescent tablets, chewing tablets, dragees, capsules, drops, juices or syrups are, for example, suitable for oral administration. Oral pharmaceutical formulations may also be in the form of multiparticulates such as granules, pellets, spheres, crystals and the like, optionally compressed into a tablet, filled into a capsule, filled into a sachet or suspended in a suitable liquid medium. Oral pharmaceutical formulations may also be equipped with an enteric coating.

[0049] Pharmaceutical formulations that are suitable for parenteral, topical and inhalative administration include but are not limited to solutions, suspensions, easily reconstitutable dry preparations and sprays.

[0050] Suppositories are a suitable pharmaceutical formulation for rectal administration. Formulations in a deposit, in dissolved form, for example, in a patch optionally with the addition of agents to promote skin penetration, are examples of suitable formulations for percutaneous administration.

[0051] One or both of the components (a) and (b) may be present in the inventive pharmaceutical formulation at least

partially in controlled-release form. Moreover, any controlled release/immediate release combination of said components may also be present in the inventive pharmaceutical formulation. For example, one or both of the components may be released from the inventive formulations with a certain delay, e.g. if administered orally, rectally or percutaneously. Such formulations are particularly useful for "once-daily" or "twice-daily" preparations, which only have to be taken once a day, respectively, twice a day. Suitable controlled-release materials are well known to those skilled in the art.

[0052] The inventive pharmaceutical formulations may be produced using materials, means, devices and processes that are well known in the prior art of pharmaceutical formulations, as described for example in "Remington's Pharmaceutical Sciences", A.R. Gennaro (ed.), 17th edition, Mack Publishing Company, Easton, Pa. (1985), in particular in part 8, chapters 76 to 93.

[0053] In order to obtain a solid pharmaceutical formulation such as a tablet, for example, the components of the pharmaceutical composition may be granulated with a pharmaceutical carrier, for example conventional tablet ingredients such as corn starch, lactose, saccharose, sorbitol, talcum, magnesium stearate, dicalcium phosphate or pharmaceutically acceptable gums, and pharmaceutical diluents, for example water, in order to form a solid composition that contains the components in homogeneous distribution. The term "homogeneous distribution" is taken to mean that the components are distributed uniformly over the entire composition, so that said composition may easily be divided into equally effective unit dose forms, such as tablets, pills or capsules and the like. The solid composition is then divided into unit dose forms. The tablets or pills of the pharmaceutical composition according to the invention may also be coated or compounded in a different manner, in order to provide a dose form with a controlled release.

[0054] If one of the components is to be released prior to the other component, for example at least 30 minutes or 1 hour beforehand, pharmaceutical formulations having a corresponding release profile may be prepared. An example of such a formulation is an osmotically driven release system for achieving a delayed release of one component via a coating that itself contains the other component which is accordingly released earlier. In a release system of this kind, which is particularly suitable for oral administration, at least part, and preferably all, of the surface of the release system, preferably those parts that will come into contact with the release medium, is/are semipermeable, preferably equipped with a semipermeable coating, so the surface(s) is/are permeable to the release medium, but substantially, preferably entirely, impermeable to the active ingredient, the surface(s) and/or optionally the coating comprising at least one opening for releasing the active ingredient. Moreover, precisely that/those surface(s) that is/are in contact with the release medium is/are provided with a coating containing and releasing the other component. This is preferably taken to mean a system in tablet form comprising a release opening, an osmotic pharmaceutical composition core, a semipermeable membrane and a polymer portion that exerts pressure upon swelling. A suitable example of this kind of system is the system distributed by ALZA Corporation, USA under the tradenames OROS®, in particular, the OROS® Push-Pull™ System, the OROS® Delayed Push-Pull™ System, the OROS® Multi-Layer Push-Pull™ system, the OROS® Push-Stick System and also, in specific cases, the L-OROS™.

[0055] Embodiments and examples of osmotically driven release systems are, for example; disclosed in US patents 4,765,989, 4,783,337 and 4,612,008.

[0056] A further example of a suitable pharmaceutical formulation is a gel-matrix tablet, such as the products developed by Penwest Pharmaceuticals (for example, under TimeRX). Suitable examples are provided in US patents 5,330,761, 5,399,362, 5,472,711 and 5,455,046. Particularly suitable is a retarding matrix formulation, with an inhomogeneous distribution of the pharmaceutically active composition, whereby, for example, one component can be distributed in the outer region (the portion that comes into contact with the release medium most quickly) of the matrix and the other component is distributed inside the matrix. On contact with the release medium, the outer matrix layer initially (and rapidly) swells and firstly releases the first component, followed by the significantly (more) retarded release of the other component. Examples of a suitable matrix include matrices with 1 to 80 % by weight of one or more hydrophilic or hydrophobic polymers as pharmaceutically acceptable matrix formers. A further example of a suitable matrix may be inferred from US 4,389,393.

[0057] The amount of the inventive pharmaceutically active combination to be administered to the patient may vary depending on different factors well known to those skilled in the art, for example, the weight of the patient, the route of administration, or the severity of the illness.

[0058] In another one of its aspects the present invention relates to a combination as described herein for the treatment of pain, wherein the pain is preferably selected from inflammatory pain, neuropathic pain, acute pain, chronic pain, visceral pain, migraine pain and cancer pain.

[0059] In a further aspect the present invention relates to the use of an inventive combination as described herein for the treatment of pain, said pain preferably including but not being limited to inflammatory pain, neuropathic pain, acute pain, chronic pain, visceral pain, migraine pain and cancer pain.

[0060] In another aspect the present invention relates to the use of an inventive combination as described herein for the preparation of a medicament for the treatment of pain, said pain preferably including but not being limited to inflammatory pain, neuropathic pain, acute pain, chronic pain, visceral pain, migraine pain and cancer pain.

[0061] The present invention also relates to a kit comprising a combination or dosage forms comprising a combination

or individual components of the combination.

Pharmacological methods:

5 Randall-Selitto test in rats

[0062] The weight ratios of the components (a) and (b) that will lead to a supra-additive effect (synergistic effect) of the inventive pharmaceutical composition may be determined via the test of Randall and Selitto as described in Arch. Int. Pharmacodyn., 1957,111:409 to 419, which is a test of mechanical nociception that may be adapted to evaluate mechanical hyperalgesia in models of inflammatory or neuropathic pain.

[0063] Acute inflammation is induced by an intraplantar injection of 0.1 ml of a carrageenan-solution (0.5 % in distilled water) into a hind paw of a rat. The mechanical nociceptive threshold is measured 4 hours later using an Algesiometer (Ugo Basile, Italy). The device generates a mechanical force with a linear increase over time. The force is applied to the dorsal surface of the inflamed rat hind paw via a cone-shaped stylus with a rounded tip (2 mm tip diameter). The nociceptive threshold is defined as the force (in grams) at which the rat vocalises (cut-off force 260 g). The mechanical nociceptive threshold is measured at different timepoints after the drug or vehicle administration. The antinociceptive and/or antihyperalgesic activity of the tested substance is expressed as percentages of maximum possible effect (%MPE). The group size is n = 12.

[0064] The analysis of the results with respect to a supra-additive effect of the inventive pharmaceutical composition comprising the components (a) and (b) is carried out via statistical comparison of the theoretical additive ED₆₀-value with the experimentally determined ED₅₀-value of a so-called fixed ratio combination (isobolographic analysis according to Tallarida JT, Porreca F, and Cowan A. Statistical analysis of drug-drug and site-site Interactions with isobolograms. Life Sci 1989; 45: 947 - 961).

The interaction studies presented herein were performed using equieffective doses of the two components, calculated from the ratio of the respective ED₅₀ values of the components if administered alone.

[0065] The application route was intravenous (i.v.) for (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol hydrochloride (hereinafter referred to as tapentadol hydrochloride) and intraperitoneal (i.p.) for ketamine ((R,S)-ketamine hydrochloride) and memantine (memantine hydrochloride). When tapentadol hydrochloride was applied alone, the peak effect was reached 15 min p. appl. (timepoint of first - measurement) and an ED₅₀-value of 1.75 (1.69-1.81) mg/kg i.v. was calculated. Ketamine hydrochloride induced a dose-dependent analgesic effect with an ED₅₀-value of 20.2 (19.0 - 21.3) mg/kg i. p. respectively, reaching the peak effect 15 min p. appl. According to their respective timepoint of peak effect, (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol hydrochloride was applied 15 min and ketamine hydrochloride 15 min before timepoint of measurement of the interaction-experiments (both components were applied simultaneously). Memantine hydrochloride induced a dose-dependent analgesic effect with an ED₅₀-value of 19.1 (17.0 - 22.0) mg/kg i. p. respectively, reaching the peak effect 15 min p. appl.. According to their respective timepoint of peak effect, (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol hydrochloride was applied 15 min and memantine hydrochloride 15 min before timepoint of measurement of the interaction-experiments (both components were applied simultaneously).

[0066] Thus, the time point of ED₅₀ calculation of both combinations corresponded to the timepoint of the peak effect of the respective compound. The isobolographic analysis revealed that the experimental ED₅₀-values of the combinations were significantly lower than the respective theoretical ED₅₀-values. Thus, the combination studies demonstrated significant synergistic interaction of (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol hydrochloride with the NMDA-antagonists, ketamine hydrochloride and memantine hydrochloride.

[0067] The results of the isobolographic analysis are summarized in the following table.

[0068] Experimental ED₅₀ values of tapentadol hydrochloride (A) and ketamine hydrochloride or A and memantine hydrochloride, respectively, and isobolographic analysis of the interaction between A and these NMDA-antagonists:

5
10
15
20
25
30
35
40
45
50
55

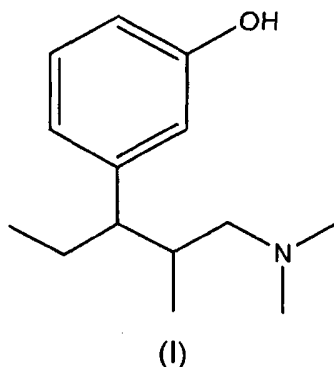
Substance / ED ₅₀ [mg/kg]	tapentadol hydrochloride (A)	ketamine hydrochloride	memantine hydrochloride	Theoretical ED ₅₀ of the combination	Experimental ED ₅₀ of the combination	Inter-action
tapentadol hydrochloride (A) + ketamine hydrochloride	1.75 (1.69 - 1.81)	20.2 (19.0 - 21.3)	-	11.0 (10.6 - 11.3)	9.2 (8.6 - 9.7)	supra-additive (p < 0.001)
tapentadol hydrochloride (A) + memantine hydrochloride	1.75(1.69 -1.81)	-	19.1 (17.0 -22.0)	10.4 (9.7 - 11.1)	9.6 (9.0 - 1 0.2)	supra-additive (p< 0.05)
p: Level of statistical significance						

[0069] The dose ratio of A to ketamine hydrochloride can be calculated to be 1:11.5 and the dose ratio of A to memantine hydrochloride can be calculated to be 1:10.9.

Claims

1. A combination comprising as component(s):

(a) at least one 3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol of formula (I),



optionally in form of one of its stereoisomers, in particular an enantiomer or a diastereomer, a racemate or in form of a mixture of its stereoisomers, in particular enantiomers and/or diastereomers in any mixing ratio, or any corresponding acid addition salt thereof, and

(b) at least one NMDA-antagonist selected from the group consisting of (R,S)-ketamine or an acid addition salt thereof such as the hydrochloride addition salt, (S)-ketamine or an acid addition salt thereof such as the hydrochloride addition salt, and memantine or an acid addition salt thereof such as the hydrochloride addition salt thereof.

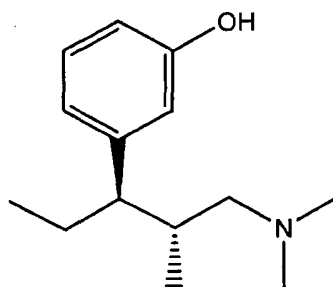
2. The combination according to claim 1, **characterized in that** the compound of formula (I) is selected from

(1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol,
(1S,2S)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol,
(1R,2S)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol,
(1S,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, and any mixture thereof.

3. The combination according to claim 2, **characterized in that** the compound of formula (I) is selected from

(1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol,
(1S,2S)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, and any mixture thereof.

4. The combination according to claim 2 or 3, **characterized in that** the compound of formula (I) is (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol of formula (I'),



(I')

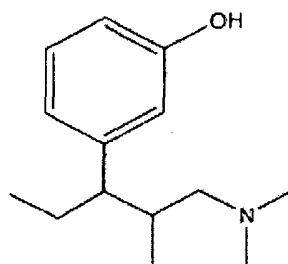
or an acid addition salt thereof, whereby the acid addition salt of hydrochloride is preferred.

5. A pharmaceutical composition comprising a combination according to any one of claims 1-4 and optionally one or more auxiliary agents:
6. A dosage form comprising a combination according to any one of claims 1-4 and optionally one or more auxiliary agents.
7. The dosage form according to claim 6, **characterized in that** it is suitable for oral, intravenous, intraarterial, intraperitoneal, intradermal, transdermal, intrathecal, intramuscular, intranasal, transmucosal, subcutaneous, or rectal administration.
8. The dosage form according to claim 6 or 7, **characterized in that** one or both of the components (a) and (b) is/are present in controlled-release form.
9. The combination according to any one of claims 1-4 for use in the treatment of pain.
10. The combination according to claim 9, wherein the pain is selected from inflammatory pain, neuropathic pain, acute pain, chronic pain, visceral pain, migraine pain and cancer pain.
11. The combination according to claim 9, wherein the pain is inflammatory pain.
12. Use of the combination according to any one of claims 9 to 11 for the manufacture of a medicament for the treatment of pain in a mammal, which comprises administering an effective amount of the combination according to any one of claims 1 to 4 to the mammal.
13. The use according to claim 12, **characterized in that** component (a) and component (b) of the combination are administered simultaneously or sequentially to the mammal wherein compound (a) may be administered before or after compound (b) and wherein compounds (a) and (b) are administered to the mammal either by the same or different pathway of administration.
14. The use according to claims 12 or 13, **characterized in that** the pain is selected from inflammatory pain, neuropathic pain, acute pain, chronic pain, visceral pain, migraine pain, and cancer pain.
15. The use according to any one of claims 12 to 14, **characterized in that** the pain is inflammatory pain.

Patentansprüche

1. Kombination, die als Komponente(n) umfasst:

(a) mindestens ein 3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol der Formel (I)



(I)

optional in der Form eines seiner Stereoisomere, insbesondere eines Enantiomers oder Diastereomers, eines Racemats oder in der Form einer Mischung seiner Stereoisomere, insbesondere Enantiomere und/oder Diastereomere in jedem Mischungsverhältnis, oder jedes entsprechenden Säureadditionssalzes davon, und (b) mindestens einen NMDA-Antagonisten, der ausgewählt wird aus der Gruppe bestehend aus (R,S)-Ketamin oder einem Säureadditionssalz davon, wie dem Hydrochloridadditionssalz, (S)-Ketamin oder einem Säureadditionssalz davon, wie dem Hydrochloridadditionssalz, und Memantin oder einem Säureadditionssalz davon, wie dem Hydrochloridadditionssalz.

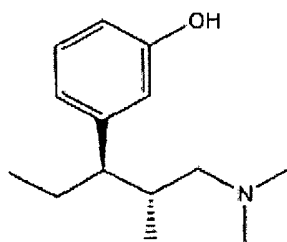
2. Kombination gemäß Anspruch 1, **gekennzeichnet dadurch, dass** die Verbindung der Formel (I) ausgewählt wird aus

(1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol,
 (1S,2S)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol,
 (1R,2S)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol,
 (1S,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, und jeder Mischung daraus.

3. Kombination gemäß Anspruch 2, **gekennzeichnet dadurch, dass** die Verbindung der Formel (I) ausgewählt wird aus

(1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol,
 (1S,2S)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol, und jeder Mischung daraus.

4. Kombination gemäß Anspruch 2 oder 3, **gekennzeichnet dadurch, dass** die Verbindung der Formel (I) (1R,2R)-3-(3-Dimethylamino-1-ethyl-2-methyl-propyl)-phenol der Formel (I')



(I')

oder ein Säureadditionssalz davon ist, wobei das Säureadditionssalz von Hydrochlorid bevorzugt ist.

5. Pharmazeutische Zusammensetzung bestehend aus einer Kombination gemäß einem der Ansprüche 1-4 und optional einem oder mehreren Hilfsmitteln.

6. Dosierungsform bestehend aus einer Kombination gemäß einem der Ansprüche 1-4 und optional einem oder mehreren Hilfsmitteln.

7. Dosierungsform gemäß Anspruch 6, **gekennzeichnet dadurch, dass** sie sich für die orale, intravenöse, intraarte-

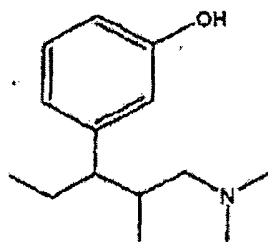
rielle, intraperitoneale, intradermale, transdermale, intrathecale, intramusculaire, intranasale, transmukosale, subkutane und rektale Verabreichung eignet.

8. Dosierungsform gemäß Anspruch 6 oder 7, **gekennzeichnet dadurch, dass** eine oder beide der Komponenten (a) und (b) in der Form der kontrollierten Freisetzung vorliegt/vorliegen.
9. Kombination gemäß einem der Ansprüche 1-4 zur Verwendung in der Behandlung von Schmerzen.
10. Kombination gemäß Anspruch 9, wobei die Schmerzen ausgewählt werden aus Entzündungsschmerzen, neuropathischen Schmerzen, akuten Schmerzen, chronischen Schmerzen, viszeralen Schmerzen, Migräneschmerzen und Krebschmerzen.
11. Kombination gemäß Anspruch 9, wobei die Schmerzen Entzündungsschmerzen sind.
12. Verwendung der Kombination gemäß einem der Ansprüche 9 bis 11 zur Herstellung eines Medikaments zur Behandlung von Schmerzen bei einem Säugetier, welche die Verabreichung einer wirksamen Menge der Kombination gemäß einem der Ansprüche 1-4 bei dem Säugetier umfasst.
13. Verwendung gemäß Anspruch 12, **gekennzeichnet dadurch dass** Komponente (a) und Komponente (b) dem Säugetier gleichzeitig oder aufeinander folgend verabreicht werden, wobei Verbindung (a) vor oder nach Verbindung (b) verabreicht werden kann, und wobei die Verbindungen (a) oder (b) dem Säugetier entweder über denselben oder unterschiedliche Verabreichungspfade verabreicht werden.
14. Verwendung gemäß Anspruch 12 oder 13, **gekennzeichnet dadurch, dass** die Schmerzen ausgewählt werden aus Entzündungsschmerzen, neuropathischen Schmerzen, akuten Schmerzen, chronischen Schmerzen, viszeralen Schmerzen, Migräneschmerzen und Krebschmerzen.
15. Verwendung gemäß einem der Ansprüche 12 bis 14, **gekennzeichnet dadurch, dass** die Schmerzen Entzündungsschmerzen sind.

Revendications

1. Association comprenant comme constituant(s) :

(a) au moins un 3-(3-diméthylamino-1-éthyl-2-méthyl-propyl)-phénol de formule (I)



(I),

éventuellement sous forme d'un de ses stéréoisomères, en particulier d'un énantiomère ou d'un diastéréoisomère, d'un racémate ou sous forme d'un mélange de ces stéréoisomères, en particulier d'énantiomères et/ou de diastéréoisomères en n'importe quel rapport de mélange, ou n'importe lequel de ces sels d'additions d'acides correspondants, et

(b) au moins un antagoniste de NMDA choisi dans le groupe consistant en la (R,S)-kétamine ou un de ses sels d'addition d'acide tels que le chlorhydrate, la (S)-kétamine ou un de ses sels d'addition d'acide tels que le chlorhydrate et la mémantine ou un de ses sels d'addition d'acide tels que le chlorhydrate.

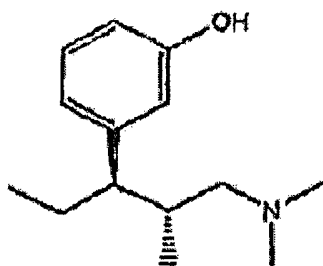
2. Association suivant la revendication 1, **caractérisée en ce que** le composé de formule (I) est choisi entre

le (1R,2R)-3-(3-diméthylamino-1-éthyl-2-méthyl-propyl)-phénol,
le (1S,2S)-3-(3-diméthylamino-1-éthyl-2-méthyl-propyl)-phénol,
le (1R,2S)-3-(3-D1_méthylamino-1-éthyl-2-méthyl-propyl)-phénol,
le (1S,2R)-3-(3-diméthylamino-1-éthyl-2-méthyl-propyl)-phénol et n'importe lequel de leurs mélanges.

3. Association suivant la revendication 2, **caractérisée en ce que** le composé de formule (I) est choisi entre

(1R,2R)-3-(3-diméthylamino-1-éthyl-2-méthyl-propyl)-phénol,
le (1S,2S)-3-(3-diméthylamino-1-éthyl-2-méthyl-propyl)-phénol et n'importe lequel de leurs mélanges.

4. Association suivant la revendication 2 ou 3, **caractérisée en ce que** le composé de formule (I) est le (1R,2R)-3-(3-diméthylamino-1-éthyl-2-méthyl-propyl)-phénol de formule (I'),



(I')

ou un de ses sels d'addition d'acide, le sel d'addition d'acide consistant en chlorhydrate étant préféré.

5. Composition pharmaceutique comprenant une association suivant l'une quelconque des revendications 1 à 4 et éventuellement un ou plusieurs agents auxiliaires.

6. Forme posologique comprenant une association suivant l'une quelconque des revendications 1 à 4 et éventuellement un ou plusieurs agents auxiliaires.

7. Forme posologique suivant la revendication 6, **caractérisée en ce qu'elle** est apte à l'administration orale, intraveineuse, intra-artérielle, intrapéritonéale, intradermique, transdermique, intrathécale, intramusculaire, intranasale, transmuqueuse, sous-cutanée ou rectale.

8. Forme posologique suivant la revendication 6 ou 7, **caractérisée en ce que** l'un des ou les deux constituants (a) et (b) est/sont présent(s) sous une forme à libération contrôlée.

9. Association suivant l'une quelconque des revendications 1 à 4, pour une utilisation dans le traitement de la douleur.

10. Association suivant la revendication 9, la douleur étant choisie entre la douleur inflammatoire, la douleur neuropathique, la douleur aiguë, la douleur chronique, la douleur viscérale, la douleur due à la migraine et la douleur due au cancer.

11. Association suivant la revendication 9, dans laquelle la douleur est la douleur inflammatoire.

12. Utilisation de l'association suivant l'une quelconque des revendications 9 à 11 pour la production d'un médicament pour le traitement de la douleur chez un mammifère, qui comprend l'administration d'une quantité efficace de l'association suivant l'une quelconque des revendications 1 à 4 au mammifère.

13. Utilisation suivant la revendication 12, **caractérisée en ce que** le constituant (a) et le constituant (b) de l'association sont administrés simultanément ou séquentiellement au mammifère, le constituant (a) pouvant être administré avant ou après le constituant (b), et les constituant (a) et (b) étant administrés au mammifère par la même voie d'administration ou bien par des voies d'administration différentes.

EP 2 582 366 B1

14. Utilisation suivant la revendication 12 ou 13, **caractérisée en ce que** la douleur est choisie entre la douleur inflammatoire, la douleur neuropathique, la douleur aiguë, la douleur chronique, la douleur viscérale, la douleur due à la migraine et la douleur due au cancer.

5 **15.** Utilisation suivant l'une quelconque des revendications 12 à 14, **caractérisée en ce que** la douleur est la douleur inflammatoire.

10

15

20

25

30

35

40

45

50

55

REFERENCES CITED IN THE DESCRIPTION

This list of references cited by the applicant is for the reader's convenience only. It does not form part of the European patent document. Even though great care has been taken in compiling the references, errors or omissions cannot be excluded and the EPO disclaims all liability in this regard.

Patent documents cited in the description

- WO 2007128412 A [0003]
- WO 2007128413 A [0003]
- WO 2010025931 A [0003]
- WO 2003053427 A [0004]
- DE 4426245 [0004]
- WO 2009094563 A [0004]
- US 8248737 B1 [0011]
- US 4765989 A [0055]
- US 4783337 A [0055]
- US 4612008 A [0055]
- US 5330761 A [0056]
- US 5399362 A [0056]
- US 5472711 A [0056]
- US 5455046 A [0056]
- US 4389393 A [0056]

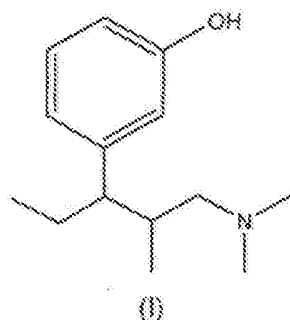
Non-patent literature cited in the description

- Textbook of Drug Design and Discovery. Taylor and Francis, 2002, 410-458 [0013]
- Analgesics. Wiley-VCH Verlag GmbH & Co. KGaA, 2002, 389-428 [0018]
- Remington's Pharmaceutical Sciences. Mack Publishing Company, 1985 [0052]
- *Arch. Int. Pharmacodyn.*, 1957, vol. 111, 409-419 [0062]
- **TALLARIDA JT ; PORRECA F ; COWAN A.** Statistical analysis of drug-drug and site-site Interactions with isobolograms. *Life Sci*, 1989, vol. 45, 947-961 [0064]

Az EP 2 582 366 lajstromszámú európai szabadalom igénypontjainak magyar fordítása:

1. Kombináció, amely komponens(ek)ként tartalmaz:

(a) legalább egy (I) képletű 3-(3-dimetilamino-1-etil-2-metil-propil)-fenolt:



adott esetben sztereoizomerjei egyikének, különösen enantiomer vagy diasztereomer, racemát formájában vagy sztereoizomerjei, különösen enantiomerjei és/vagy diasztereomerjei bármilyen keverési arányú keveréke, vagy bármilyen megfelelő savaddíciós sója formájában, és

(b) legalább egy NMDA-antagonistát, amelyet az (R,S)-ketamin vagy savaddíciós sója, így például hidroklorid addíciós sója, (S)-ketamin vagy savaddíciós sója, így például hidroklorid addíciós sója, és memantin vagy savaddíciós sója, így például hidroklorid addíciós sója által alkotott csoportból választunk.

2. Az 1. igénypont szerinti kombináció, azzal jellemezve, hogy az (I) képletű vegyületet az

(1R,2R)-3-(3-dimetilamino-1-etil-2-metil-propil)-fenol,

(1S,2S)-3-(3-dimetilamino-1-etil-2-metil-propil)-fenol,

(1R,2S)-3-(3-dimetilamino-1-etil-2-metil-propil)-fenol,

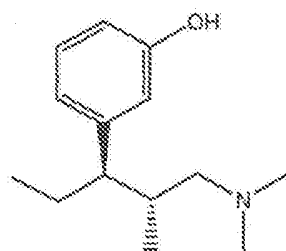
(1S,2R)-3-(3-dimetilamino-1-etil-2-metil-propil)-fenol és ezek bármilyen keveréke közül választjuk.

3. A 2. igénypont szerinti kombináció, azzal jellemezve, hogy az (I) képletű vegyületet az

(1R,2R)-3-(3-dimetilamino-1-etil-2-metil-propil)-fenol,

(1S,2S)-3-(3-dimetilamino-1-etil-2-metil-propil)-fenol és ezek bármilyen keveréke közül választjuk.

4. A 2. vagy 3. igénypont szerinti kombináció, azzal jellemezve, hogy az (I) képletű vegyület (I') képletű (1R,2R)-3-(3-dimetilamino-1-etil-2-metil-propil)-fenol:



(I')

vagy savaddíciós sója, ahol a hidroklorid savaddíciós sóét részesítjük előnyben.

5. Gyógyászati készítmény, amely az 1-4. igénypontok bármelyike szerinti kombinációt és adott esetben egy vagy több segédanyagot tartalmaz.

6. Adagolási forma, amely az 1-4. igénypontok bármelyike szerinti kombinációt és adott esetben egy vagy több segédanyagot tartalmaz.

7. A 6. igénypont szerinti adagolási forma, azzal jellemezve, hogy alkalmas orális, intravénás, intraarteriális, intraperitoneális, intradermális, transzdermális, intratekális, intramuszkuláris, intranazális, transzmukozális, szubkután vagy rektális adagolásra.

8. A 6. vagy 7. igénypont szerinti adagolási forma, azzal jellemezve, hogy az (a) és (b) komponensek közül egy vagy mindkettő szabályozott hatóanyag-leadású formában van.

9. Az 1-4. igénypontok bármelyike szerinti kombináció fájdalom kezelésében való alkalmazásra.

10. A 9. igénypont szerinti kombináció, ahol a fájdalmat a gyulladásos fájdalom, neuropátiás fájdalom, akut fájdalom, krónikus fájdalom, zsigeri fájdalom, migrénes fájdalom és a rákos fájdalom közül választjuk.

11. A 9. igénypont szerinti kombináció, ahol a fájdalom gyulladásos fájdalom.

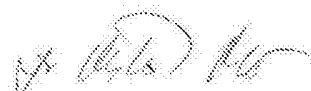
12. A 9-11. igénypontok bármelyike szerinti kombináció alkalmazása gyógyszer előállítására emulzióban fájdalom kezelésére, amely tartalmazza az 1-4. igénypontok bármelyike szerinti kombináció hatékony mennyiségének emulzióknak történő adagolását.

13. A 12. igénypont szerinti alkalmazás, azzal jellemezve, hogy a kombináció (a) komponensét és (b) komponensét egy időben vagy egymást követően adagoljuk az emulzióknak, ahol az (a) vegyületet adagolhatjuk a (b) vegyület előtt vagy után, és ahol az (a) és (b) vegyületeket az emulzióknak ugyanolyan vagy különböző úton adagoljuk.

14. A 12. vagy 13. igénypont szerinti alkalmazás, azzal jellemezve, hogy a fájdalmat a gyulladásos fájdalom, neuropátiás fájdalom, akut fájdalom, krónikus fájdalom, zsigeri fájdalom, migrénes fájdalom és a rákos fájdalom közül választjuk.

15. A 12-14. igénypontok bármelyike szerinti alkalmazás, azzal jellemezve, hogy a fájdalom gyulladásos fájdalom.

A meghatalmazott:



Dr. Miskolczi Mária
Ügyvéd
111 Budapest, Egy József u. 40.
Tel: 215-4148 Fax: 215-4508