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(54) **NOVEL PHOSPHODIESTERASE INHIBITORS**
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Description

FIELD OF THE INVENTION

- 5 **[0001]** The present invention relates to novel compounds with phosphodiesterase inhibitory activity, as well as to their use as therapeutic agents in the treatment of inflammatory diseases and conditions.

BACKGROUND OF THE INVENTION

- 10 **[0002]** Phosphodiesterases are enzymes that catalyse the hydrolysis of cyclic AMP and/or cyclic GMP in cells to 5-AMP and 5-GMP, respectively, and as such they are critical to cellular regulation of cAMP or cGMP levels. Of the 11 phosphodiesterases identified so far, phosphodiesterase (PDE) 4, PDE7 and PDE8 are selective for cAMP. PDE4 is the most important modulator of cAMP expressed in immune and inflammatory cells such as neutrophils, macrophages and T-lymphocytes (Z. Huang and J.A. Mancini, *Current Med. Chem.* 13, 2006, pp. 3253-3262). As cAMP is a key
 15 second messenger in the modulation of inflammatory responses, PDE4 has been found to regulate inflammatory responses of inflammatory cells by modulating proinflammatory cytokines such as TNF α , IL-2, IFN- γ , GM-CSF and LTB₄. Inhibition of PDE4 has therefore become an attractive target for the therapy of inflammatory diseases such as asthma, chronic obstructive pulmonary disease (COPD), rheumatoid arthritis, atopic dermatitis, Crohn's disease etc. (M.D. Houslay et al., *Drug Discovery Today* 10 (22), 2005, pp. 1503-1519). As atopic dermatitis (AD) patients have increased
 20 PDE-activity, PDE4-inhibition would also appear to be a viable treatment of AD (*Journal of Investigative Dermatology* (1986), 87(3), 372-6).

- [0003]** The PDE4 gene family consists at least of four genes, A, B, C and D, which have a high degree of homology (V. Boswell Smith and D. Spina, *Curr. Opin. Investig. Drugs* 6(11), 2006, pp. 1136-1141). The four PDE4 isoforms are differentially expressed in different tissues and cell types. Thus, PDE4B is predominantly expressed in monocytes
 25 and neutrophils, but not in cortex and epithelial cells, while PDE4D is expressed in lung, cortex, cerebellum and T-cells (C. Kroegel and M. Foerster, *Exp. Opin. Investig. Drugs* 16(1), 2007, pp. 109-124). It has been speculated that inhibition of PDE4D in the brain is associated with the adverse effects found when administering PDE4 inhibitors clinically, primarily nausea and emesis, whereas inhibition of PDE4B is associated with anti-inflammatory effects (B. Lipworth, *Lancet* 365, 2005, pp. 167-175). However, the PDE inhibitors developed so far are not believed to be specific for any of the four
 30 PDE4 isoforms.

- [0004]** Numerous PDE4 inhibitors have been studied for their therapeutic effect on inflammatory diseases, primarily asthma, inflammatory bowel disease and COPD. The first of these, theophylline, is a weak, non-selective phosphodiesterase inhibitor used in the treatment of respiratory diseases such as asthma and COPD. Treatment with theophylline may, however, give rise to both mild and severe adverse effects, e.g. arrhythmia and convulsions, restricting the clinical
 35 utility of theophylline (Kroegel and Foerster, *supra*). As phosphodiesterase has remained an attractive target for anti-inflammatory therapy, several other, more selective PDE4 inhibitors have been developed and investigated in a clinical setting. The clinical development of many of the first-generation PDE4 inhibitors such as rolipram was discontinued due to dose-limiting side effects, primarily nausea and emesis. Second-generation PDE4 inhibitors with apparently less pronounced adverse effects are currently in clinical trials (Houslay, *supra*).

- 40 **[0005]** Recently developed PDE-4 inhibitors are for example disclosed in EP 0771794 and EP 0943613. WO 96/31476 discloses structurally different 4-substituted-3,5-dichloropyridines which are inhibitors of cyclic AMP phosphodiesterase.

[0006] US 2002/0128290 discloses oxygen containing heterocyclic compounds which have PDE4 inhibitory activity.

- [0007]** There is a continued need for developing novel PDE4 inhibitors which have a more favourable therapeutic window, i.e. fewer adverse effects, while retaining their therapeutic anti-inflammatory effect. An overview of preclinical
 45 and clinical trials with selective PDE4 inhibitors, including such inhibitors aimed for the treatment of atopic dermatitis and psoriasis, was recently given in *Inflammation & Allergy: Drug Targets*, 2007, 6(1), 17-26.

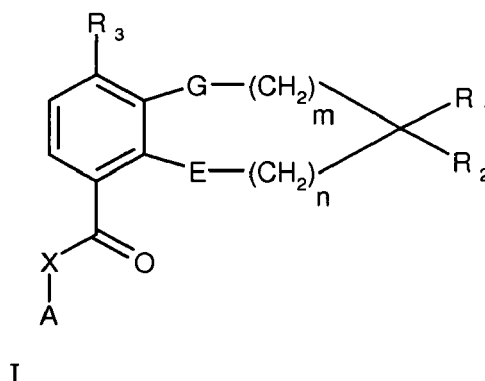
SUMMARY OF THE INVENTION

- 50 **[0008]** The inventors have surprisingly found that novel compounds of the present invention exhibit PDE4 inhibitory activity and may be useful as therapeutic agents for inflammatory allergic diseases such as bronchial asthma, allergic rhinitis, and nephritis; autoimmune diseases such as rheumatoid arthritis, multiple sclerosis, Crohn's disease, and systemic lupus erythematosus; diseases of the central nervous system such as depression, amnesia, and dementia; organopathy associated with ischemic reflux caused by cardiac failure, shock, and cerebrovascular diseases, and the like;
 55 insulin-resistant diabetes; wounds; AIDS, and the like.

[0009] Compounds of the present invention may also be beneficial in preventing, treating or ameliorating a variety of diseases, such as dermal diseases or conditions, such as proliferative and inflammatory skin disorders and in particular psoriasis, epidermal inflammation, alopecia, skin atrophy, steroid induced skin atrophy, skin ageing, photo skin ageing,

acne, dermatitis, atopic dermatitis, seborrheic dermatitis, contact dermatitis, urticaria, pruritis, and eczema.

[0010] Accordingly, the present invention relates to a compound according to formula I,



wherein m and n independently represent 0, 1, 2, 3, 4, 5, 6, or 7;

wherein G and E independently represent sulphur, oxygen, -N=, -N(R₅)-, or -N(R₅)C(O)-, and

R₁ and R₂, together with the carbon atom to which they are attached, form a heterocyclic ring comprising one or two heteroatoms selected from oxygen, sulphur, -S(O)-, -S(O)₂-, -N(R₅)-, one or more carbon atoms in said heterocyclic ring being optionally substituted with one or more, same or different substituents selected from R₄;

R₃ is halogen, hydroxy, C₁₋₆-alkyl, C₂₋₆-alkenyl, C₂₋₆-alkynyl, halo-C₁₋₆-alkyl, C₁₋₆-alkoxy, halo-C₁₋₆-alkoxy, C₁₋₆-alkylthio, formyl, C₁₋₆-alkoxycarbonyl, C₁₋₆-alkylcarbonyl, or aminocarbonyl;

R₄ is hydrogen, amino, thioxo, C₁₋₆-alkyl, halo-C₁₋₆-alkyl, hydroxy-C₁₋₆-alkyl, C₁₋₆-alkoxy, halogen, oxo, or hydroxy;

R₅ is hydrogen, C₁₋₆-alkyl, halo-C₁₋₆-alkyl, C₁₋₆-alkylcarbonyl, hydroxy-C₁₋₆-alkyl, C₁₋₆-alkoxycarbonyl, C₁₋₆-alkylsulfonyl, C₁₋₆-alkylaminosulfonyl or aminosulfonyl;

X is a bond, -CH₂-, or -NH-;

A is aryl comprising 6-10 carbon atoms, C₃₋₁₀-cycloalkyl, C₃₋₁₀-cycloalkenyl, aryl-C₁₋₆-alkyl, heteroaryl comprising 1-10 carbon atoms and 1-5 hetero atoms, heteroaryl-C₁₋₆-alkyl, the heteroaryl comprising 1-10 carbon atoms and 1-5 hetero atoms, hetero-C₃₋₁₀-cycloalkyl comprising 1-3 hetero atoms or hetero-C₃₋₁₀-cycloalkenyl comprising 1-3 hetero atoms, optionally substituted with one or more, same or different substituents selected from R₄;

and pharmaceutically acceptable salts, hydrates, N-oxides or solvates thereof.

[0011] In another aspect, the invention relates to a pharmaceutical composition comprising a compound of general formula I as defined above together with a pharmaceutically acceptable vehicle or excipient or pharmaceutically acceptable carrier(s), optionally together with one or more other therapeutically active compound(s).

[0012] In yet another aspect, the invention relates to a compound according to formula I as defined above, and pharmaceutically acceptable salts, hydrates, N-oxides or solvates thereof, for use in the prophylaxis, treatment or amelioration of dermal diseases or conditions, or acute or chronic cutaneous wound disorders. In yet another aspect, the invention relates to a method of preventing, treating or ameliorating dermal diseases or conditions, or acute or chronic cutaneous wound disorders, the method comprising administering to a person suffering from at least one of said diseases an effective amount of one or more compounds of formula I as defined above and pharmaceutically acceptable salts, hydrates, N-oxides or solvates thereof; optionally together with a pharmaceutically acceptable carrier or one or more excipients, and optionally in combination with other therapeutically active compounds.

DETAILED DESCRIPTION OF THE INVENTION

[0013] The term "hydrocarbon radical" is intended to indicate a radical containing only hydrogen and carbon atoms, it may contain one or more double and/or triple carbon-carbon bonds, and it may comprise cyclic moieties in combination with branched or linear moieties. Said hydrocarbon comprises 1-20 carbon atoms, and preferably comprises 1-12, e.g. 1-6, e.g. 1-4, e.g. 1-3, e.g. 1-2 carbon atoms. The term includes alkyl, alkenyl, cycloalkyl, cycloalkenyl, alkynyl and aryl, arylalkyl, as indicated below.

[0014] The term "aryl" is intended to indicate a radical of aromatic carbocyclic rings comprising 6-20 carbon atoms, such as 6-14 carbon atoms, preferably 6-10 carbon atoms, in particular 5- or 6-membered rings, optionally fused carbocyclic rings with at least one aromatic ring, such as phenyl, naphthyl, indenyl and indanyl.

[0015] The term "heteroaryl" is intended to indicate radicals of heterocyclic aromatic rings comprising 1-6 heteroatoms (selected from O, S and N) and 1-20 carbon atoms, such as 1-5 heteroatoms and 1-10 carbon atoms, such as 1-5

heteroatoms and 1-6 carbon atoms, such as 1-5 heteroatoms and 1-3 carbon atoms, in particular 5- or 6-membered rings with 1-4 heteroatoms selected from O, S and N, or optionally fused bicyclic rings with 1-4 heteroatoms, and wherein at least one ring is aromatic, e.g. pyridyl, quinolyl, isoquinolyl, indolyl, tetrazolyl, thiazolyl, imidazolyl, pyrazolyl, oxazolyl, isoxazolyl, thienyl, pyrazinyl, isothiazolyl, benzimidazolyl and benzofuranyl.

[0016] In the present context, the term "alkyl" is intended to indicate the radical obtained when one hydrogen atom is removed from a hydrocarbon. Said alkyl comprises 1-20, preferably 1-12, such as 1-6, such as 1-4 carbon atoms. The term includes the subclasses normal alkyl (*n*-alkyl), secondary and tertiary alkyl, such as methyl, ethyl, *n*-propyl, isopropyl, *n*-butyl, isobutyl, *sec.*-butyl, *tert.*-butyl, pentyl, isopentyl, hexyl and isohexyl.

[0017] The term "cycloalkyl" is intended to indicate a saturated cycloalkane radical comprising 3-20 carbon atoms, preferably 3-10 carbon atoms, in particular 3-8 carbon atoms, such as 3-6 carbon atoms, including fused bicyclic rings, e.g. cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, or cycloheptyl.

[0018] The term "heterocycloalkyl" is intended to indicate a cycloalkane radical as described above, wherein one or more carbon atoms are replaced by heteroatoms, comprising 1-19 carbon atoms, e.g. 2-4 carbon atoms, further comprising 1-6 heteroatoms, preferably 1, 2, or 3 heteroatoms, selected from O, N, or S, which may optionally be oxidised once or twice, e.g. [1,3]dioxole, oxetane, [1,3]dioxolane, [1,3]dioxane, tetrahydrothiopyran, tetrahydrothiopyran-1,1-dioxide, tetrahydrothiopyran-1-oxide, piperidine, tetrahydrothiophene, [1,3]-dithiane, thietane, [1,3]-dithiane-1,3-dioxide, or thietane-1-oxide, or including fused bicyclic rings with 1-4 heteroatoms, wherein at least one ring comprises a heteroatom, and wherein the other ring may for example be a carbocyclic ring, e.g. isoindolyl.

[0019] The term "alkenyl" is intended to indicate a mono-, di-, tri-, tetra- or pentaunsaturated hydrocarbon radical comprising 2-10 carbon atoms, in particular 2-6 carbon atoms, such as 2-4 carbon atoms, e.g. ethenyl, propenyl, butenyl, pentenyl or hexenyl.

[0020] The term "cycloalkenyl" is intended to indicate mono-, di- tri- or tetra-unsaturated non-aromatic cyclic hydrocarbonsradicals, comprising 3-20 carbon atoms, including fused bicyclic rings, typically comprising 3-10 carbon atoms, such as 3, 4, or 6 carbon atoms, e.g. cyclopropenyl, cyclobutenyl, cyclopentenyl, cyclohexenyl, cycloheptenyl.

[0021] The term "heterocycloalkenyl" is intended to indicate a cycloalkene radical as described above, wherein one or more carbon atoms are replaced by heteroatoms, comprising 1-19 carbon atoms, e.g. 2-4 carbon atoms, further comprising 1-6 heteroatoms, preferably 1, 2, or 3 heteroatoms, selected from O, N, or S, including fused bicyclic rings with 1-4 heteroatoms, wherein at least one ring comprises a heteroatom and wherein the other ring may for example be a carbocyclic ring, e.g. dihydrofuranyl, or 2,5-dihydro-1H-pyrrolyl.

[0022] The term "arylalkyl" is intended to indicate an aryl radical as defined above covalently joined to an alkyl group, e.g. benzyl.

[0023] The term "heteroarylalkyl" is intended to indicate a heteroaryl radical as defined above covalently joined to an alkyl group.

[0024] The term "alkynyl" is intended to indicate an hydrocarbon radical comprising 1-5 triple C-C bonds and 2-20 carbon atoms, typically comprising 2-10 carbon atoms, in particular 2-6 carbon atoms, such as 2-4 carbon atoms, e.g. ethynyl, propynyl, butynyl, pentynyl or hexynyl.

[0025] The term "halogen" is intended to indicate a substituent from the 7th main group of the periodic table, such as fluoro, chloro, bromo and iodo.

[0026] The term "haloalkyl" is intended to indicate an alkyl group as defined above substituted with one or more halogen atoms as defined above, e.g. difluoromethyl.

[0027] The term "hydroxyalkyl" is intended to indicate an alkyl group as defined above substituted with one or more hydroxy, e.g. hydroxymethyl, hydroxyethyl, hydroxypropyl.

[0028] The term "alkoxy" is intended to indicate a radical of the formula -OR', wherein R' is alkyl as indicated above, e.g. methoxy, ethoxy, *n*-propoxy, isopropoxy, butoxy, etc.

[0029] The term "alkoxycarbonyl" is intended to indicate a radical of the formula -C(O)-O-R', wherein R' is alkyl as indicated above, e.g. methoxycarbonyl, ethoxycarbonyl, *n*-propoxycarbonyl, isopropoxycarbonyl, etc.

[0030] The term "alkylcarbonyl" is intended to indicate a radical of the formula -C(O)-R', wherein R' is alkyl as indicated above, e.g. ethanoyl, acetyl.

[0031] The term "aminosulfonyl" is intended to indicate a radical of the formula -S(O)₂-NR", wherein R' is as indicated above, e.g. -SO₂Me.

[0032] The term "heterocyclic ring" is intended to include the definitions heteroaryl, heterocycloalkyl and heterocycloalkenyl as defined above, further including annelated ring systems with each other or with cyclic hydrocarbons, e.g. 2,5-dihydrobenzo(b)dioxocine, 2,3,5,8-tetrahydro-[1,4]dioxocine, 5,8-dihydro-[1,4]dioxocine.

[0033] The term "pharmaceutically acceptable salt" is intended to indicate salts prepared by reacting a compound of formula I with a suitable inorganic or organic acid, such as hydrochloric, hydrobromic, hydroiodic, sulfuric, nitric, phosphoric, formic, acetic, 2,2-dichloroacetic, adipic, ascorbic, L-aspartic, L-glutamic, galactaric, lactic, maleic, L-malic, phthalic, citric, propionic, benzoic, glutaric, gluconic, D-glucuronic, methanesulfonic, salicylic, succinic, malonic, tartaric, benzenesulfonic, ethane-1,2-disulfonic, 2-hydroxy ethanesulfonic acid, toluenesulfonic, sulfamic or fumaric acid. Pharma-

aceutically acceptable salts of compounds of formula I may also be prepared by reaction with a suitable base such as sodium hydroxide, potassium hydroxide, magnesium hydroxide, calcium hydroxide, silver hydroxide, ammonia or the like, or suitable non-toxic amines, such as lower alkylamines, for example triethylamine, hydroxy-lower alkylamines, for example 2-hydroxyethylamine, bis-(2-hydroxyethyl)-amine, cycloalkylamines, for example dicyclohexylamine, or benzylamines, for example N,N'-dibenzylethylenediamine, and dibenzylamine, or L-arginine or L-lysine. Salts obtained by reaction with a suitable base include, but are not limited to sodium salts, choline salts, 2-(dimethylamino)-ethanol salts, 4-(2-hydroxy-ethyl)-morpholin salts, L-lysine salts, N-(2-hydroxyethyl)-pyrrolidine salts, ethanolamine salts, potassium salts, tetrabutylammonium salts, benzyltrimethylammonium salts, cetyltrimethylammonium salts, tetramethylammonium salts, tetrapropylammonium salts, tris(hydroxymethyl)aminomethane salts, N-methyl-D-glucamine salts, silver salts, ben-zethonium salts, and triethanolamine salts.

[0034] The term "solvate" is intended to indicate a species formed by interaction between a compound, e.g. a compound of formula I, and a solvent, e.g. alcohol, glycerol or water, wherein said species are in a solid form. When water is the solvent, said species is referred to as a hydrate.

Embodiments of the present invention

[0035] In one or more embodiments of the present invention E and G are both oxygen.

[0036] In one or more embodiments of the present invention m and n are both one.

[0037] In one or more embodiments of the present invention m and n are both zero.

[0038] In one or more embodiments of the present invention R_1 and R_2 , together with the carbon atom to which they are attached, form a heterocyclic ring comprising one or two heteroatoms selected from the group consisting of -O-, -S-, -S(O)-, -S(O)₂-, -N=, and -N(R₅)-; one or more carbon atoms in the heterocyclic ring being optionally substituted with one or more substituents, same or different, selected from R₄.

[0039] In one or more embodiments of the present invention R_1 and R_2 , together with the carbon atom to which they are attached, form a 4-, 5- or 6-membered heterocyclic ring, in particular a 6-membered heterocyclic ring.

[0040] In one or more embodiments of the present invention the heterocyclic ring is tetrahydropyran, oxetane, [1,3]dioxolane, [1,3]dioxane, tetrahydrothiopyran, tetrahydrothiopyran-1,1-dioxide, tetrahydrothiopyran-1-oxide, piperidine, tetrahydrothiophene, [1,3]-dithiane, thietane, [1,3]-dithiane-1,3-dioxide, thietane-1-oxide, or thiethane-1,1-dioxide.

[0041] In one or more embodiments of the present invention the heterocyclic ring formed of R_1 and R_2 together with the carbon atom to which they are attached comprises one heteroatom or two heteroatoms in said ring.

[0042] In one or more embodiments of the present invention the heteroatom is located in position 4 of the heterocyclic ring. The heteroatom may, for example, be O.

[0043] In one or more embodiments of the present invention the heteroatom(s) is/are oxygen, sulphur, -S(O)-, or -S(O)₂-.

[0044] In one or more embodiments of the present invention A represents heteroaryl comprising 1-10 carbon atoms and 1-5 hetero atoms or heteroaryl-C₁₋₆-alkyl, the heteroaryl comprising 1-10 carbon atoms and 1-5 hetero atoms.

[0045] In one or more embodiments of the present invention A represents pyridyl, pyrazinyl or quinolyl.

[0046] In other embodiments A may represent phenyl.

[0047] In one or more embodiments of the present invention A is substituted with halogen, in particular chlorine, fluorine, bromine or iodine.

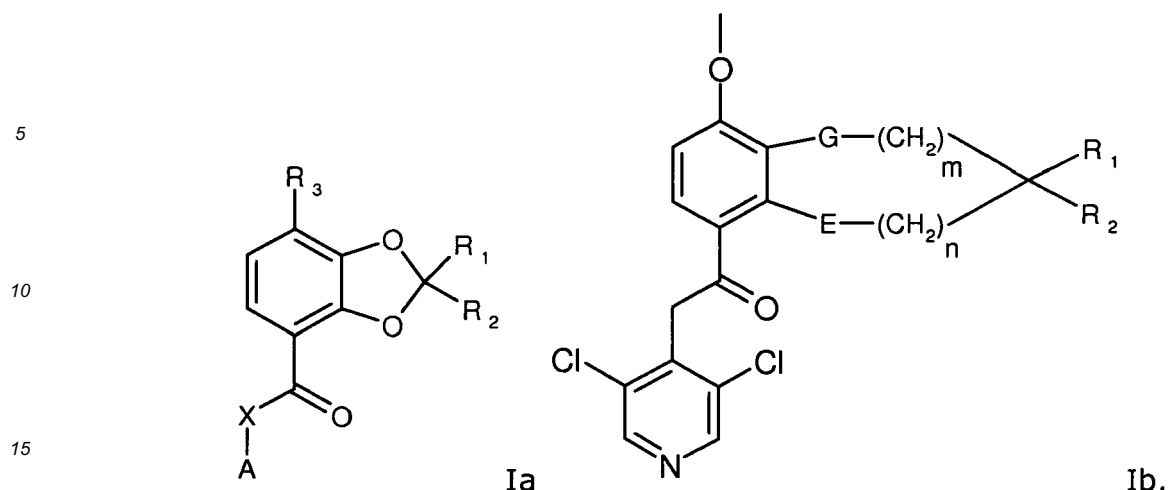
[0048] In one or more embodiments of the present invention R_3 represents C₁₋₆ alkoxy, C₁₋₆ halogenalkyl, or halogen.

[0049] In one or more embodiments of the present invention R_3 represents methoxy or ethoxy.

[0050] In one or more embodiments of the present invention X is -CH₂- or -NH-.

[0051] In one or more embodiments of the present invention A is 4-(3,5-dichloro-pyridyl).

[0052] In one or more embodiments of the present invention, the compound of formula I is represented by formula Ia or Ib



wherein X, A, G, E, R₁, R₂, R₃, R₄, R₅, m and n are as defined above.

[0053] In a particular embodiment of the present invention X = -NH- when R₃ represents C₁₋₆ alkoxy.

[0054] The present invention includes all embodiments wherein X, A, G, E, R₁, R₂, R₃, R₄, R₅ are combined in any manner described herein.

[0055] In particular compounds of formula I may be selected from one of the following compounds:

2-((3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-yl)ethanone (compound 101), N-(3,5-Dichloropyridine-4-yl)-7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-carboxamide (compound 102), 2-(3,5-Dichloro-1-oxido-pyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-yl)ethanone (compound 103), 2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-4',5'-dihydro-spiro[1,3-benzodioxole-2,3'-(2H)-thiophen]-4-yl)ethanone (compound 104), 2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 105), 2-((3,5-Dichloropyridine-4-yl)-1-(7-methoxy-1'-[methoxycarbonyl]-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 106), 2-((3,5-Dichloropyridine-4-yl)-1-(7-methoxy-1'-[methylsulfonyl]-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 107), 2-((3,5-Dichloropyridine-4-yl)-1-(7-methoxy-1'-acetyl-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 108), 2-((3,5-Dichloropyridine-4-yl)-1-(7-methoxy-1'-methyl-spiro[1,5-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 109), 2-((3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran]-4-yl)ethanone (compound 110), 2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran 1'-oxide]-4-yl)ethanone (compound 111), 2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran-1',1'-dioxide]-4-yl)ethanone (compound 112), or 2-((3,5-Dichloro-1-oxido-pyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran-1',1'-dioxide]-4-yl)ethanone (compound 113); 2-((3-bromopyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (compound 114); 2-(3-Bromo-pyrazin-2-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (compound 115); 2-(pyrazin-2-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (compound 116); 2-(pyridin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (compound 117); 2-(quinolin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (compound 118); 2-(2,6-Dichloro-phenyl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (compound 119); 2-(2-Chloro-phenyl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (compound 120); 2-(3,5-Dichloropyridine-4-yl)-1-(9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-oxetane]-6-yl)ethanone (compound 121); 2-((3,5-Dichloro-1-oxido-pyridine-4-yl)-1-(9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-oxetane]-6-yl)ethanone (compound 122); 2-((3,5-Dichloropyridine-4-yl)-1-(9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietane]-6-yl)ethanone (compound 123); 2-((3,5-Dichloropyridine-4-yl)-1-(9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietane-1',1'-dioxide]-6-yl)eth-

anone (compound 124);

2-((3,5-Dichloropyridin-1-oxido-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietane-1',1'-dioxide]-6-yl})ethanone (compound 125); 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),2'-(1,3-dioxolane)]-6-yl}ethanone (compound 126);

2-((3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),4'-tetrahydropyran]-6-yl})ethanone (compound 127);

2-((3,5-Dichloro-1-oxido-pyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),4'-tetrahydropyran]-6-yl})ethanone (compound 128);

2-((3,5-Dichloropyridin-4-yl)-1-{9-methoxy-2',2'-dimethyl-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxane]-6-yl})ethanone (compound 129);

2-((3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxane]-6-yl})ethanone (compound 130);

2-((3,5-Dichloro-1-oxido-pyridine-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxane]-6-yl})ethanone (compound 131); and

2-((3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dithiane]-6-yl})ethanone (compound 132)

and pharmaceutically acceptable salts, hydrates, N-oxides or solvates thereof.

[0056] In one or more embodiments of the present invention, the compounds of formula I as defined above are useful in therapy.

[0057] The compounds of formula I may be obtained in crystalline form either directly by concentration from an organic solvent or by crystallisation or recrystallisation from an organic solvent or mixture of said solvent and a cosolvent that may be organic or inorganic, such as water. The crystals may be isolated in essentially solvent-free form or as a solvate, such as a hydrate. The invention covers all crystalline modifications and forms and also mixtures thereof.

[0058] Compounds of formula I may or may not comprise asymmetrically substituted (chiral) carbon atoms which give rise to the existence of isomeric forms, e.g. enantiomers and possibly diastereomers. The present invention relates to all such isomers, either in pure form or as mixtures thereof (e.g. racemates). Pure stereoisomeric forms of the compounds and the intermediates of this invention may be obtained by the application of procedures known in the art. The various isomeric forms may be separated by physical separation methods such as selective crystallization and chromatographic techniques, e.g. liquid chromatography using chiral stationary phases. Enantiomers may be separated from each other by the selective crystallization of their diastereomeric salts with optically active amines, such as 1-ephedrine. Alternatively, enantiomers may be separated by chromatographic techniques using chiral stationary phases. Said pure stereoisomeric forms may also be derived from the corresponding pure stereoisomeric forms of the appropriate starting materials, provided that the reaction occurs stereoselectively or stereospecifically. Preferably, if a specific stereoisomer is desired, said compound will be synthesized by stereoselective or stereospecific methods of preparation. These methods will advantageously employ chiral pure starting materials.

[0059] Compounds of the invention, optionally in combination with other active compounds, may be useful for the treatment of dermal diseases or conditions, or acute or chronic cutaneous wound disorders, in particular for the treatment of proliferative and inflammatory skin disorders, psoriasis, cancer, epidermal inflammation, alopecia, skin atrophy, steroid induced skin atrophy, skin ageing, photo skin ageing, acne, dermatitis, atopic dermatitis, seborrheic dermatitis, contact dermatitis, urticaria, pruritis, and eczema.

[0060] Besides being useful for human treatment, the compounds of the present invention may also be useful for veterinary treatment of animals including mammals such as horses, cattle, sheep, pigs, dogs, and cats.

[0061] For use in therapy, compounds of the present invention are typically in the form of a pharmaceutical composition. The invention therefore relates to a pharmaceutical composition comprising a compound of formula I, optionally together with one or more other therapeutically active compound(s), together with a pharmaceutically acceptable excipient or vehicle. The excipient must be "acceptable" in the sense of being compatible with the other ingredients of the composition and not deleterious to the recipient thereof.

[0062] Conveniently, the active ingredient comprises from 0.05-99.9% by weight of the formulation.

[0063] In the form of a dosage unit, the compound may be administered one or more times a day at appropriate intervals, always depending, however, on the condition of the patient, and in accordance with the prescription made by the medical practitioner. Conveniently, a dosage unit of a formulation contains between 0.1 mg and 1000 mg, preferably between 1 mg and 100 mg, such as 5-50 mg of a compound of formula I.

[0064] A suitable dosage of the compound of the invention will depend, inter alia, on the age and condition of the patient, the severity of the disease to be treated and other factors well known to the practising physician. The compound may be administered either orally, parenterally or topically according to different dosing schedules, e.g. daily or with weekly intervals. In general a single dose will be in the range from 0.01 to 400 mg/kg body weight. The compound may be administered as a bolus (i.e. the entire daily dose is administered at once) or in divided doses two or more times a day.

[0065] In the context of topical treatment it may be more appropriate to refer to a "usage unit", which denotes a single dose which is capable of being administered to a patient, and which may be readily handled and packed, remaining as a physically and chemically stable unit dose comprising either the active material as such or a mixture of it with solid or liquid pharmaceutical diluents or carriers. The term "usage unit" in connection with topical use means a unitary, i.e. a single dose capable of being administered topically to a patient in an application per square centimetre of the infected area of from 0.1 mg to 10 mg, and preferably from 0.2 mg to 1 mg, of the active ingredient in question.

[0066] It is also envisaged that in certain treatment regimes, administration with longer intervals, e.g. every other day, every week, or even with longer intervals may be beneficial.

[0067] If the treatment involves administration of another therapeutically active compound it is recommended to consult Goodman & Gilman's *The Pharmacological Basis of Therapeutics*, 9th Ed., J.G. Hardman and L.E. Limbird (Eds.), McGraw-Hill 1995, for useful dosages of said compounds.

[0068] The administration of a compound of the present invention with one or more other active compounds may be either concomitantly or sequentially.

[0069] The formulations include e.g. those in a form suitable for oral (including sustained or timed release), rectal, parenteral (including subcutaneous, intraperitoneal, intramuscular, intraarticular and intravenous), transdermal, ophthalmic, topical, dermal, nasal or buccal administration. Topical administration of the claimed formulation is particularly suitable.

[0070] The formulations may conveniently be presented in dosage unit form and may be prepared by any of the methods well known in the art of pharmacy, e.g. as disclosed in Remington, *The Science and Practice of Pharmacy*, 20th ed., 2000. All methods include the step of bringing the active ingredient into association with the carrier, which constitutes one or more accessory ingredients. In general, the formulations are prepared by uniformly and intimately bringing the active ingredient into association with a liquid carrier or a finely divided solid carrier or both, and then, if necessary, shaping the product into the desired formulation.

[0071] Formulations of the present invention suitable for oral administration may be in the form of discrete units as capsules, sachets, tablets or lozenges, each containing a predetermined amount of the active ingredient; in the form of a powder or granules; in the form of a solution or a suspension in an aqueous liquid or non-aqueous liquid, such as ethanol or glycerol; or in the form of an oil-in-water emulsion or a water-in-oil emulsion. Such oils may be edible oils, such as e.g. cottonseed oil, sesame oil, coconut oil or peanut oil. Suitable dispersing or suspending agents for aqueous suspensions include synthetic or natural gums such as tragacanth, alginate, acacia, dextran, sodium carboxymethylcellulose, gelatin, methylcellulose, hydroxypropylmethylcellulose, hydroxypropylcellulose, carbomers and polyvinylpyrrolidone. The active ingredients may also be administered in the form of a bolus, electuary or paste.

[0072] A tablet may be made by compressing or moulding the active ingredient optionally with one or more accessory ingredients. Compressed tablets may be prepared by compressing, in a suitable machine, the active ingredient(s) in a free-flowing form such as a powder or granules, optionally mixed by a binder, such as e.g. lactose, glucose, starch, gelatine, acacia gum, tragacanth gum, sodium alginate, carboxymethylcellulose, methylcellulose, hydroxypropylmethylcellulose, polyethylene glycol, waxes or the like; a lubricant such as e.g. sodium oleate, sodium stearate, magnesium stearate, sodium benzoate, sodium acetate, sodium chloride or the like; a disintegrating agent such as e.g. starch, methylcellulose, agar, bentonite, croscarmellose sodium, sodium starch glycollate, crospovidone or the like or a dispersing agent, such as polysorbate 80. Moulded tablets may be made by moulding, in a suitable machine, a mixture of the powdered active ingredient and suitable carrier moistened with an inert liquid diluent.

[0073] Formulations for rectal administration may be in the form of suppositories in which the compound of the present invention is admixed with low melting water soluble or insoluble solids such as cocoa butter, hydrogenated vegetable oils, polyethylene glycol or fatty acids esters of polyethylene glycols, while elixirs may be prepared using myristyl palmitate.

[0074] Formulations suitable for parenteral administration conveniently comprise a sterile oily or aqueous preparation of the active ingredients, which is preferably isotonic with the blood of the recipient, e.g. isotonic saline, isotonic glucose solution or buffer solution. The formulation may be conveniently sterilised by for instance filtration through a bacteria retaining filter, addition of sterilising agent to the formulation, irradiation of the formulation or heating of the formulation. Liposomal formulations as disclosed in e.g. *Encyclopedia of Pharmaceutical Technology*, vol.9, 1994, are also suitable for parenteral administration.

[0075] Alternatively, the compounds of formula I may be presented as a sterile, solid preparation, e.g. a freeze-dried powder, which is readily dissolved in a sterile solvent immediately prior to use.

[0076] Transdermal formulations may be in the form of a plaster or a patch:

Formulations suitable for ophthalmic administration may be in the form of a sterile aqueous preparation of the active ingredients, which may be in microcrystalline form, for example, in the form of an aqueous microcrystalline suspension. Liposomal formulations or biodegradable polymer systems e.g. as disclosed in *Encyclopedia of Pharmaceutical Technology*, vol.2, 1989, may also be used to present the active ingredient for ophthalmic administration.

Formulations suitable for topical or ophthalmic administration include liquid or semi-liquid preparations such as liniments, lotions, gels, sprays, foams, oil-in-water or water-in-oil emulsions such as creams, ointments or pastes; or solutions or suspensions such as drops. Compositions for ophthalmic treatment may preferably additionally contain a cyclodextrin.

[0077] For topical administration, the compound of formula I may typically be present in an amount of from 0.01 to 20% by weight of the composition, such as 0.1% to about 10 %, but may also be present in an amount of up to about 50% of the composition.

[0078] Formulations suitable for nasal or buccal administration include powder, self-propelling and spray formulations, such as aerosols and atomisers. Such formulations are disclosed in greater detail in e.g. Modern Pharmaceutics, 2nd ed., G.S. Banker and C.T. Rhodes (Eds.), page 427-432, Marcel Dekker, New York; Modern Pharmaceutics, 3th ed., G.S. Banker and C.T. Rhodes (Eds.), page 618-619 and 718-721, Marcel Dekker, New York and Encyclopedia of Pharmaceutical Technology, vol. 10, J. Swarbrick and J.C. Boylan (Eds), page 191-221, Marcel Dekker, New York.

[0079] In addition to the aforementioned ingredients, the formulations of a compound of formula I may include one or more additional ingredients such as diluents, buffers, flavouring agents, colourant, surface active agents, thickeners, preservatives, e.g. methyl hydroxybenzoate (including anti-oxidants), emulsifying agents and the like.

[0080] When the active ingredient is administered in the form of salts with pharmaceutically acceptable non-toxic acids or bases, preferred salts are for instance easily water-soluble or slightly soluble in water, in order to obtain a particular and appropriate rate of absorption.

[0081] The pharmaceutical composition may additionally comprise one or more other active components conventionally used in the treatment of dermal disease or conditions, e.g. selected from the group consisting of glucocorticoids, vitamin D and vitamin D analogues, antihistamines, platelet activating factor (PAF) antagonists, anticholinergic agents, methyl-xanthines, β -adrenergic agents, COX-2 inhibitors, salicylates, indomethacin, flufenamate, naproxen, timegadine, gold salts, penicillamine, serum cholesterol lowering agents, retinoids, zinc salts, salicylazosulfapyridine and calcineurin inhibitors.

[0082] The invention is described in further detail in the following examples which are not in any way intended to limit the scope of the invention as claimed.

METHODS OF PREPARATION

[0083] The compounds of the present invention can be prepared in a number of ways well known to those skilled in the art of synthesis. The compounds of formula I may for example be prepared using the reactions and techniques outlined below together with methods known in the art of synthetic organic chemistry, or variations thereof as appreciated by those skilled in the art. Preferred methods include, but are not limited to, those described below. The reactions are carried out in solvents appropriate to the reagents and materials employed and suitable for the transformations being effected. Also, in the synthetic methods described below, it is to be understood that all proposed reaction conditions, including choice of solvent, reaction atmosphere, reaction temperature, duration of experiment and work-up procedures, are chosen to be conditions of standard for that reaction, which should be readily recognized by one skilled in the art. Not all compounds falling into a given class may be compatible with some of the reaction conditions required in some of the methods described. Such restrictions to the substituents which are compatible with the reaction conditions will be readily apparent to one skilled in the art and alternative methods can be used.

[0084] Starting materials are either known compounds which are commercially available, or they may be prepared by routine synthetic methods well known to a person skilled in the art.

[0085] The compounds of the present invention or any intermediate may be purified if required using standard methods well known to a synthetic organic chemist, e.g. methods described in "purification of Laboratory Chemicals", 5th ed. 2003. Starting materials are either known compounds, commercially available, or they may be prepared by routine synthetic methods well known to a person skilled in the art.

GENERAL PROCEDURES, PREPARATIONS AND EXAMPLES

[0086] ^1H nuclear magnetic resonance (NMR) spectra were recorded at 300 MHz and ^{13}C NMR spectra at 75.6 MHz. Chemical shift values (δ , in ppm) are quoted in the specified solvent relative to internal tetramethylsilane ($\delta = 0.00$) or chloroform ($\delta = 7.25$) or deuteriochloroform ($\delta = 76.81$ for ^{13}C NMR) standards. The value of a multiplet, either defined (doublet (d), triplet (t), quartet (q)) or not (m) at the approximate mid point is given unless a range is quoted. (bs) indicates a broad singlet. The organic solvents used were usually anhydrous. Chromatography was performed on Merck silica gel 60 (0.040 - 0.063 mm). The solvent ratios indicated refer to v:v unless otherwise noted.

[0087] The following abbreviations have been used throughout:

DCM	dichloromethane
DMF	N,N'-Dimethylformamide
DMSO	dimethyl sulfoxide
Et	ethyl
EtOAc	ethyl acetate
h	hour
L	litre
LDA	lithium diisopropylamide
LiHMDS	lithium Hexamethyldisilazide
m	milli
Me	methyl
MeOH	methanol
NMR	nuclear magnetic resonance
ppt	precipitate
rt	room temperature
TsCl	<i>p</i> -toluenesulphonyl chloride
THF	tetrahydrofuran
v	volume

Preparative HPLC/MS

[0088] Preparative HPLC/MS was performed on a Dionex APS-system with two Shimadzu PP150 prep. pumps and a Thermo MSQ Plus mass spectrometer. Column: Waters XTerra C-18, 150 mm x 19 mm, 5 μ m; solvents system: A = water (0.1 % formic acid) and B = acetonitrile (0.1 % formic acid); flow rate = 18 mL/min; method (10 min): Linear gradient method going from 10 % B to 100 % B in 6 minutes and staying at 100 % B for another 2 minutes. The fractions were collected based on ion traces of relevant ions and PDA signal (240-400 nm).

Analytical HPLC/MS

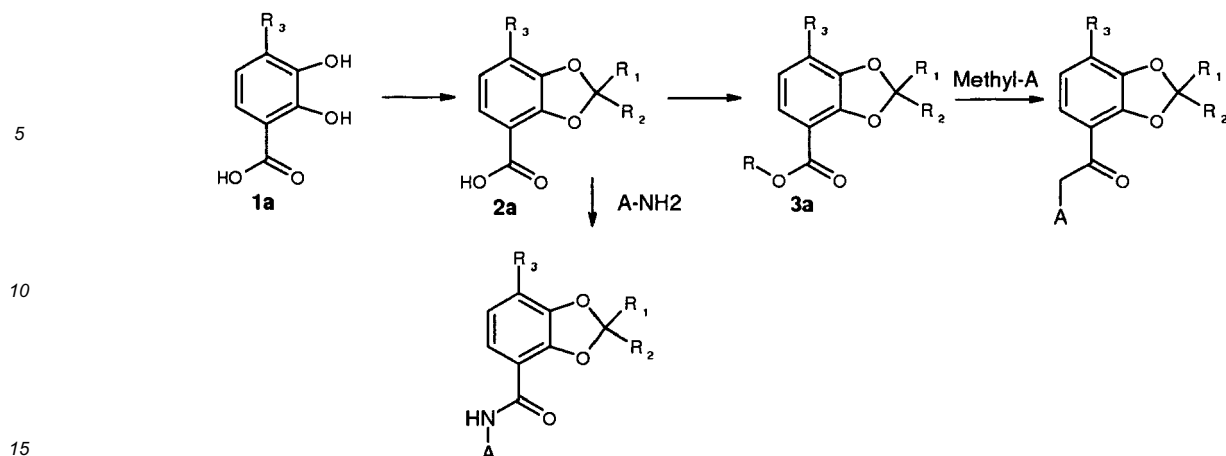
[0089] Method A: Analytical HPLC/MS was performed on a Dionex APS-system with a P680A analytical pump and a Thermo MSQ Plus mass spectrometer. Column: Waters XTerra C-18, 150 mm x 4.6 mm, 5 μ m; solvents system: A = water (0.1 % formic acid) and B = acetonitrile (0.1 % formic acid); flow rate = 1.0 mL/min; method (10 min): Linear gradient method going from 10 % B to 100 % B in 6.6 minutes and staying at 100 % B for another 1.5 minutes.

[0090] Method B: Analytical HPLC/MS was performed on a system consisting of a Waters 2795 HPLC, Micromass ZQ mass spectrometer, Waters 996 PDA. Column: Waters XTerra C-18, 50 mm x 3.0 mm, 5 μ m; solvents system: A = water:acetonitrile 95:5 (0.05 % formic acid) and B = acetonitrile (0.05 % formic acid); flow rate = 1.0 mL/min; method (8 min): Linear gradient method going from 10 % B to 100 % B in 6.0 minutes and staying at 100 % B for 1 minute.

General procedure of preparation:

[0091] The compounds of the invention can for example be prepared by the following general methods:

Compounds of the general formula Ia, wherein R_1 , R_2 , and R_3 are as defined above, can be prepared as follows:



[0092] Starting materials of formula **1a** are prepared according to standard procedures known to chemist skilled in the art. 2,3,4-trimethoxy benzoic acid is selectively di-demethylated at the 2- and 3-position using BCl_3 according to Kaisalo et al. Synth. Commun, (1986), 16, 645-48.

[0093] Subsequent reaction of the deprotected compound in neat ketones, enol ethers, ketals or a mixture of these with or without additional catalysts such as paratoluenesulfonic acid or a lewis acid at temperature from room temperature to 180 °C using either microwave or conventional heating, results in the compounds **2a**.

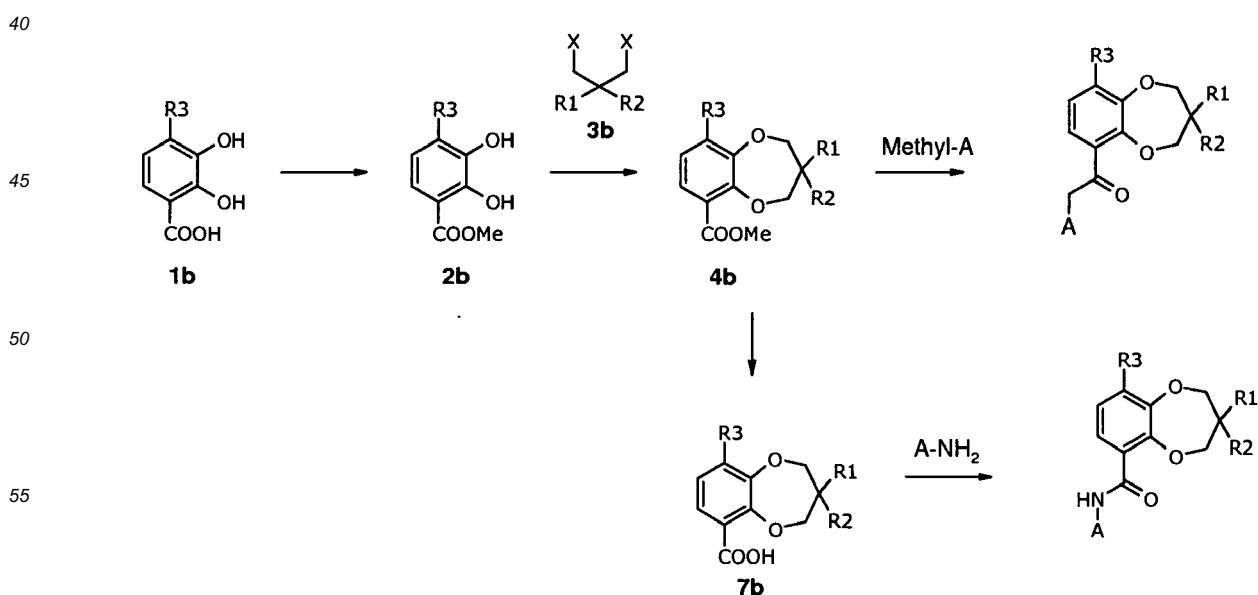
[0094] Reaction of compounds with formula **2a** with MeI (or dimethyl sulphate) in the presence of a suitable base, such as K_2CO_3 $KHCO_3$ or Et_3N in a suitable solvent such as DMF, acetone, THF or DCM at temperatures from room temperature to 100°C give compounds of the formula **3a**.

[0095] The ester **3a** may also be prepared by classical esterification methods using the alcohol and a suitable acid e.g. H_2SO_4 .

[0096] Compounds of the formula Ia ($X=CH_2$) were obtained by condensation of the generated methyl ester with lithio carbanions generated from A-Methyl, wherein A is defined as described above, and a suitable base, such as LDA or LiHMDS in a suitable solvent such as THF at temperatures from minus 78°C to room temperature. Alternatively to the lithio carbanions a grignard reagent may be used.

[0097] Compounds of the formula Ia ($X=NH$) were obtained by reaction of compounds with the formula **2a** with $(COCl)_2$, $SOCl_2$ or PCl_5 in a suitable solvent such as DCM or toluene with or without catalytic amount of DMF at temperatures from 0°C to 70°C to afford the corresponding acid chloride. After evaporation of the solvent *in vacuo* subsequent condensation of the generated acid chloride with nitrogen-anions, generated by addition a suitable base, such as NaH, LDA or LiHMDS in a suitable solvent such as THF at temperatures from minus 78°C to room temperature to $A-NH_2$, wherein A is defined as described above, are performed.

[0098] Compounds of the general formular Ib, wherein R_1 , R_2 , and R_3 are as defined above, can be prepared as follows:



[0099] Esterification of **1b** using standard procedures, e.g. MeOH and H₂SO₄ results in the ester **2b**.

[0100] Alkylation of **2b** using **3b** (X= Br, I, OTs) in the presence of a suitable base, such as K₂CO₃ in a suitable solvent such as DMSO at temperatures from room temperature to 120°C give compounds of the formula **4b**.

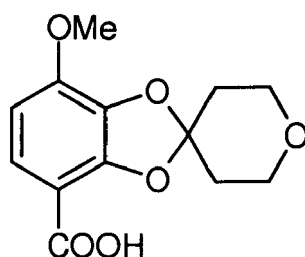
[0101] Compounds of the formula Ib (X=CH₂) were obtained by condensation of the generated methyl ester with lithio carbanions generated from A-Methyl, wherein A is defined as described above, and a suitable base, such as LDA or LiHMDS in a suitable solvent such as THF at temperatures from minus 78°C to room temperature. Alternatively to the lithio carbanions a grignard reagent may be used.

[0102] Esther hydrolysis using standard conditions (acidic or basic) is expected to result in the carboxylic acid **7b**, that can be converted to the carboxylic acid chloride and subsequently reacted with nitrogen-anions (generated from A-NH₂) as described for the synthesis of Ia (X=NH).

Preparation 1:

7-Methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-carboxylic acid (compound 501)

[0103]

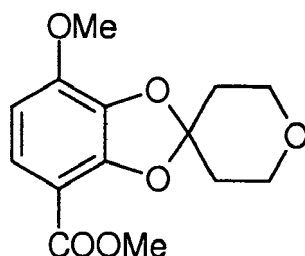


[0104] A suspension of 2,3-dihydroxy-4-methoxybenzoic acid (6.04 g, 32.8 mmol) in 5,6-dihydro-4-methoxy-2H-pyran (20 mL, 152 mmol) was kept at 140 °C for three days. At room temperature ethyl acetate (200 mL) was added and the organic phase was extracted with saturated aqueous NaHCO₃ (2 x 50 mL). The aqueous phase was washed with Et₂O (2 x 40 mL), acidified to pH = 1 with concentrated HCl and extracted with dichloromethane (2 x 50 mL). The organic phase was dried over MgSO₄. Evaporation under reduced pressure afforded traces of 2,3-dihydroxy-4-methoxybenzoic acid along with 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-carboxylic acid (1.23 g, 14%). ¹³C NMR (DMSO) δ 164.9, 148.2, 146.6, 134.5, 123.7, 117.0, 107.1, 106.8, 64.4, 56.0, 35.3.

Preparation 2:

Methyl 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-carboxylate (compound 502)

[0105]



[0106] A suspension of the crude 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-pyran]-4-carboxylic acid (2.17 g, 8.15 mmol), KHCO₃ (2.58 g, 26.0 mmol) and dimethyl sulphate (1.58 mL, 16.7 mmol) in acetone (62 mL) was stirred at room temperature for two days before it was evaporated to dryness under reduced pressure. Ethyl acetate (100 mL) was added. The organic phase was washed with 0.5 M aqueous NaOH (6 x 30 mL) and evaporated to dryness under reduced pressure. The crude product was redissolved in dichloromethane (75 mL), dried over MgSO₄ and evaporated to dryness under reduced pressure. Standard silica gel column chromatography afforded methyl 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-carboxylate (1.87 g, 79%). ¹³C NMR (CDCl₃)

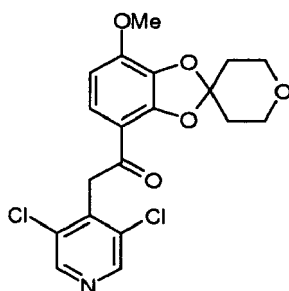
δ 164.9, 149.1, 147.2, 135.2, 124.0, 117.5, 107.1, 106.5, 65.2, 56.4, 51.8, 35.9.

Standard procedure A:

Example 1:

2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-yl)ethanone (compound 101)

[0107]

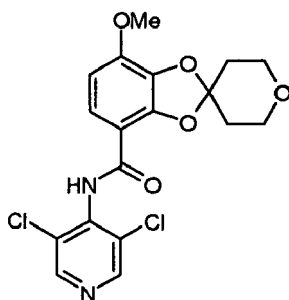


[0108] A solution of methyl 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-carboxylate (1.80 g, 6.42 mmol) and 3,5-dichloro-4-picoline (1.46 g, 8.99 mmol) in tetrahydrofuran (33 mL) was cooled to 0 °C. A 1.0 M solution of lithium bis(trimethylsilyl)amide in tetrahydrofuran (19.3 mL, 19.3 mmol) was added and the reaction mixture was allowed to reach room temperature overnight. Saturated aqueous NH_4Cl (70 mL) was added. The aqueous phase was extracted with dichloromethane (3 x 100 mL). The combined organic phase was washed with water (50 mL), dried over MgSO_4 and evaporated to dryness under reduced pressure. Standard silica gel column chromatography followed by recrystallization from isopropanol afforded 2-(3,5-dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-yl)ethanone (1.90 g, 71%). ^{13}C NMR (DMSO) δ 189.1, 148.2, 147.7, 147.0, 141.2, 134.5, 132.8, 122.0, 118.0, 113.0, 107.8, 64.4, 56.3, 43.5, 35.2.

Example 2:

N-(3,5-Dichloropyridine-4-yl)-7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-carboxamide (compound 102)

[0109]



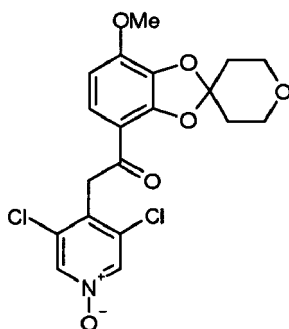
[0110] Oxalyl chloride (92 μL , 1.1 mmol) and a catalytic amount of N,N-dimethylformamide was added to a suspension of 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-carboxylic acid (48 mg, 0.18 mmol) in dichloromethane (2 mL). After stirring for one hour at room temperature, the solvent was removed under reduced pressure and the crude acid chloride was redissolved in tetrahydrofuran (2 mL). A suspension of 3,5-dichloropyridin-4-amine (67 mg, 0.40 mmol) and NaH (a 60% dispersion in mineral oil, 16 mg, 0.40 mmol) in tetrahydrofuran (1 mL) was stirred for three hours at room temperature before it was added dropwise at room temperature to the tetrahydrofuran solution containing the crude acid chloride. After having stirred overnight at room temperature the reaction mixture was diluted with diethyl ether (30 mL) and the organic phase was washed with 0.5 M aqueous NaOH (3 x 10 mL). The organic phase

was dried over MgSO_4 and evaporated to dryness under reduced pressure. Standard HPLC purification afforded N-(3,5-dichloropyridine-4-yl)-7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-carboxamide (14 mg, 19%). ^{13}C NMR (DMSO) δ 160.8, 148.0, 146.5, 146.2, 141.1, 134.1, 130.5, 122.5, 118.2, 108.3, 107.6, 64.2, 56.2, 35.2.

Example 3:

2-(3,5-Dichloro-1-oxido-pyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-yl)ethanone (compound 103)

[0111]

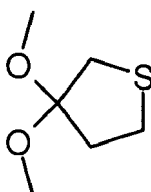


[0112] To a solution of 2-(3,5-dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-yl)ethanone (41 mg, 0.99 mmol) in dichloromethane (0.5 mL) was added 30% H_2O_2 (25 μL) and methyltrioxorhenium(VII) (3 mg). The mixture was stirred at room temperature overnight, added MnO_2 (3 mg) and was stirred for another hour. Water (10 mL) was added and the aqueous phase was extracted with dichloromethane (3 x 10 mL). The combined organic phase was dried over MgSO_4 and evaporated to dryness under reduced pressure. Standard HPLC purification afforded compound **103** (8 mg, 19%). LC/MS (METHOD B): (m/z) 426.1; 428.1 (MH⁺); RT = 2.98 min; purity (UV) = 100%

Preparation 3:

Tetrahydro-3,3-dimethoxythiophen (compound 503).

[0113]

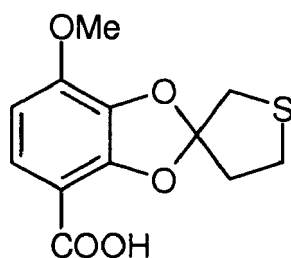


[0114] A solution of tetrahydrothiophen-3-one (10.0 g, 97.9 mmol), methyl orthoformate (21.4 mL, 196 mmol) and para-toluenesulfonic acid monohydrate (50 mg, 0.29 mmol) in dry methanol (25 mL) was refluxed for one hour. Then 1.0 M methanolic NaOMe (0.30 mL, 0.30 mmol) was added and excess of methanol and trimethyl orthoformate was removed by distillation (atmospheric pressure). Further distillation under reduced pressure afforded a mixture of tetrahydrothiophen-3-one (~0.67 g, 7%) and tetrahydro-3,3-dimethoxythiophen (~9.8 g, 67%). ^{13}C NMR (MeOH) δ 113.01, 50.11, 36.90, 36.11, 27.72.

Preparation 4:

7-Methoxy-4',5'-dihydro-spiro[1,3-benzodioxole-2,3'-(2H)-thiophen]-4-carboxylic acid (compound 504)

[0115]

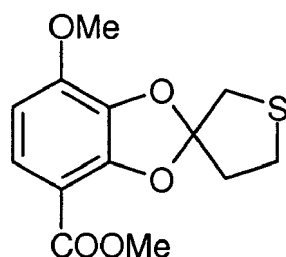


[0116] P-toluenesulfonic acid (54 mg, 0.28 mmol) was added to the mixture of tetrahydro-3,3-dimethoxythiophen (~9.8 g, 66 mmol) and tetrahydrothiophen-3-one (~0.67 g, 6.6 mmol). The oil bath was heated to 145 °C and approximately one equivalent of methanol (2.7 mL, 67 mmol) was distilled off. The temperature was lowered and distillation under reduced pressure afforded 7.04 g of an oil to which 2,3-dihydroxy-4-methoxybenzoic acid (1.00 g, 5.43 mmol) was added. The suspension was exposed to microwave heating (180 °C, one hour) in a sealed reaction vessel. Filtration and subsequent standard HPLC purification afforded compound **504** (164 mg, 11%). LC/MS (METHOD B): (m/z) 267.2(M-1); RT = 2.79 min; purity (UV) = 100%

Preparation 5:

Methyl 7-methoxy-4',5'-dihydro-spiro[1,3-benzodioxole-2,3'-(2H)-thiophen]-4-carboxylate (compound 505)

[0117]

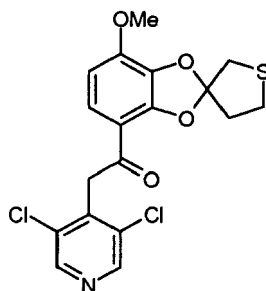


[0118] A suspension of 7-methoxy-4',5'-dihydro-spiro[1,3-benzodioxole-2,3'-(2H)-thiophen]-4-carboxylic acid (161 mg, 0.600 mmol), K₂CO₃ (166 g, 1.20 mmol) and dimethyl sulphate (74 µL, 0.78 mmol) in acetone (1 mL) was kept at 50 °C overnight. At room temperature water (15 mL) was added and the aqueous phase was extracted with ethyl acetate (2 x 20 mL). The organic phase was dried over MgSO₄ and evaporated to dryness under reduced pressure. Standard HPLC purification afforded compound **505** (24 mg, 14%). ¹H NMR (CDCl₃) δ 7.44 (d, 1H), 6.56 (d, 1H), 3.94 (s, 3H), 3.88 (s, 3H), 3.32 (d, 1H), 3.24 (d, 1H), 3.05 (t, 2H), 2.49 (td, 2H).

Example 4:

2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-4',5'-dihydro-spiro[1,3-benzodioxole-2,3'-(2H)-thiophen]-4-yl)ethanone (compound 104)

[0119]

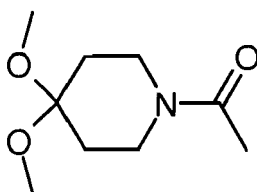


[0120] A solution of methyl 7-methoxy-4',5'-dihydro-spiro[1,3-benzodioxole-2,3'-(2H)-thiophen]-4-carboxylate (24 mg, 85 μ mol) and 3,5-dichloro-4-picoline (21 mg, 0.13 mmol) in tetrahydrofuran (1 mL) was cooled to 0 °C. A 1.0 M solution of lithium bis(trimethylsilyl)amide in tetrahydrofuran (0.26 mL, 0.26 mmol) was added and the reaction mixture was allowed to reach room temperature overnight. Saturated aqueous NH_4Cl (10 mL) was added. The aqueous phase was extracted with dichloromethane (3 x 10 mL). The combined organic phase was washed with water (20 mL), dried over MgSO_4 and evaporated to dryness under reduced pressure. Standard HPLC purification afforded the title compound (12 mg, 34%). ^{13}C NMR (DMSO) δ 189.02, 148.09, 147.51, 147.05, 140.98, 134.48, 132.69, 127.57, 122.31, 112.81, 107.81, 56.35, 43.26, 37.54, 36.54, 25.70.

Preparation 6:

1-Acetyl-4,4-dimethoxy-piperidine (compound 506)

[0121]

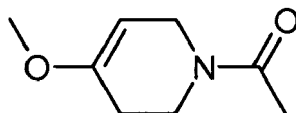


[0122] A solution of 1-acetyl-4-piperidone (17.0 g, 121 mmol), trimethyl orthoformate (26.4 mL, 241 mmol) and para-toluenesulfonic acid monohydrate (80 mg, 0.42 mmol) in dry methanol (34 mL) was refluxed for one hour. Then 1.0 M methanolic NaOMe (0.42 mL, 0.42 mmol) and excess of methanol and trimethyl orthoformate were removed by distillation (atmospheric pressure). Further distillation under reduced pressure afforded 1-acetyl-4,4-dimethoxy-piperidine (20.2 g, 89%). ^1H NMR (DMSO) δ 3.45 - 3.32 (m, 4H), 3.10 (s, 6H), 1.99 (s, 3H), 1.72 - 1.62 (m, 2H), 1.61 - 1.52 (m, 2H).

Preparation 7:

1-Acetyl-1,2,3,6-tetrahydro-4-methoxy-pyridine (compound 507)

[0123]

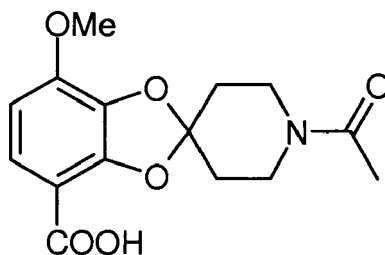


[0124] To 1-acetyl-4,4-dimethoxy-piperidine (20.2 g, 108 mmol) was added para-toluenesulfonic acid monohydrate (80 mg, 0.42 mmol). The mixture was heated to 160 °C and approximately one equivalent of methanol (4.38 mL, 108 mmol) was distilled off. The temperature was lowered and distillation under reduced pressure afforded a mixture of 1-acetyl-4,4-dimethoxy-piperidine (1.4 g, 7%) and 1-acetyl-1,2,3,6-tetrahydro-4-methoxy-pyridine (14.2 g, 85%). ^1H NMR (DMSO) δ 4.68 - 4.62 (m, 1H), 4.00 - 3.88 (m, 2H), 3.59 - 3.49 (m, 2H), 3.49 - 3.45 (m, 3H), 2.19 - 2.12 (m, 1H), 2.09 - 2.03 (m, 1H), 2.03 - 1.96 (m, 3H).

Preparation 8:

7-Methoxy-1'-acetyl-spiro[1,3-benzodioxole-2,4'-piperidine]-4-carboxylic acid (compound 508)

[0125]

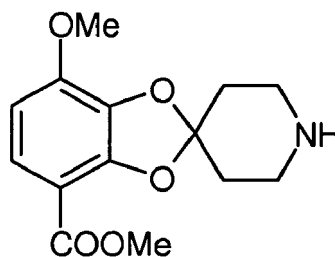


[0126] A mixture of 2,3-dihydroxy-4-methoxybenzoic acid (1.23 g, 6.67 mmol), 1-acetyl-4,4-dimethoxy-piperidine (1.4 g, 7.6 mmol) and 1-acetyl-1,2,3,6-tetrahydro-4-methoxy-pyridine (14.2 g, 91.5 mmol) was exposed to microwave heating (180 °C, one hour) in a sealed reaction vessel. Filtration and subsequent standard HPLC purification afforded compound **508** (0.54 g, 26%). LC/MS (METHOD B): (m/z) 308.2 (MH⁺); RT = 2.27 min; purity (UV) = 95%.

Preparation 9:

Methyl 7-methoxy-spiro[1,3-benzodioxole-2,4'-piperidine]-4-carboxylate (compound 509)

[0127]

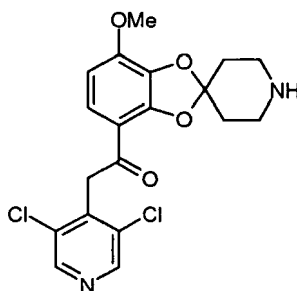


[0128] A solution of 7-methoxy-1'-acetyl-spiro[1,3-benzodioxole-2,4'-piperidine]-4-carboxylic acid (143 mg, 0.467 mmol) and LiOH (224 mg, 9.34 mmol) in water (3 mL) and MeOH (3 mL) was heated to 75 °C for five hours. At room temperature the mixture was neutralized with 2M HCl and evaporated to dryness under reduced pressure. The crude 7-methoxy-spiro[1,3-benzodioxole-2,4'-piperidine]-4-carboxylic acid [LC/MS (METHOD B): (m/z) 266.2 (MH⁺); RT = 1.57 min; purity (UV) = 82%] was refluxed overnight in 1.7 M methanolic HCl (5 mL). At room temperature water (20 mL) was added. The aqueous phase was washed with Et₂O (10 mL), made basic by addition of Na₂CO₃ and extracted with dichloromethane (3 x 10 mL). The organic phase was dried over MgSO₄ and evaporation under reduced pressure afforded methyl 7-methoxy-spiro[1,3-benzodioxole-2,4'-piperidine]-4-carboxylate (75 mg, 57%). ¹H NMR (DMSO) δ 7.31 (d, 1H), 6.72 (d, 1H), 3.87 (s, 3H), 3.78 (s, 3H), 2.96 - 2.77 (m, 4H), 1.94 - 1.83 (m, 4H).

Example 5:

2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 105)

[0129]

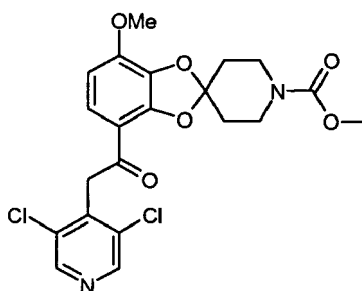


[0130] A solution of methyl 7-methoxy-spiro[1,3-benzodioxole-2,4'-piperidine]-4-carboxylate (75 mg, 0.268 mol) and 3,5-dichloro-4-picoline (65 mg, 0.40 mmol) in tetrahydrofuran (2.5 mL) was cooled to 0 °C. A 1.0 M solution of lithium bis(trimethylsilyl)amide in tetrahydrofuran (0.80 mL, 0.80 mmol) was added and the reaction mixture was allowed to reach room temperature overnight. Saturated aqueous NH₄Cl (10 mL) was added. The aqueous phase was extracted with dichloromethane (3 x 10 mL). The combined organic phase was dried over MgSO₄ and evaporated to dryness under reduced pressure. Standard HPLC purification afforded compound **105** (58 mg, 53%). ¹H NMR (DMSO) δ 8.66 (s, 2H), 7.38 - 7.36 (m, 1H), 6.83 - 6.80 (m, 1H), 4.62 (s, 2H), 3.91 (s, 3H), 3.02 - 2.91 (m, 4H), 2.12 - 1.93 (m, 4H).

Example 6:

2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-1'-[methoxycarbonyl]-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 106)

[0131]

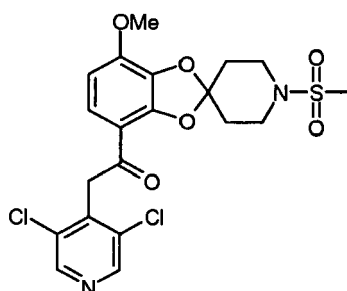


[0132] A solution of 2-(3,5-dichloropyridine-4-yl)-1-(7-methoxy-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (10 mg, 24 μmol), triethylamine (24 μL, 171 μmol) and methyl chloroformate (10 μL, 122 μmol) in dichloromethane (200 μL) was kept at room temperature overnight. Water (500 μL) was added and the aqueous phase extracted with dichloromethane (3 x 500 μL). The organic phase was dried over MgSO₄ and evaporated to dryness under reduced pressure. Standard HPLC purification afforded compound **106** (2.5 mg, 22%). ¹H NMR (DMSO) δ 8.65 (s, 2H), 7.39 (d, 1H), 6.84 (d, 1H), 4.62 (s, 2H), 3.91 (s, 3H), 3.80 - 3.66 (m, 2H), 3.62 (s, 3H), 3.58 - 3.46 (m, 2H), 2.18 - 1.97 (m, 4H).

Example 7:

2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-1'-[methylsulfonyl]-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 107)

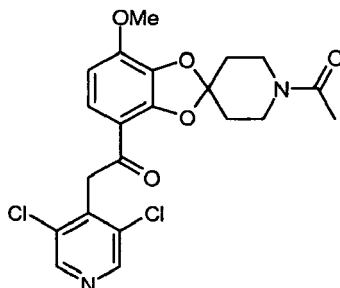
[0133]



[0134] A solution of 2-(3,5-dichloropyridine-4-yl)-1-(7-methoxy-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (10 mg, 24 μmol), triethylamine (24 μL, 171 μmol) and mesyl chloride (10 μL, 122 μmol) in dichloromethane (200 μL) was kept at room temperature overnight. Water (500 μL) was added and the aqueous phase extracted with dichloromethane (3 x 500 μL). The organic phase was dried over MgSO₄ and evaporated to dryness under reduced pressure. Standard HPLC purification afforded compound **107** (1.8 mg, 15%). ¹H NMR (DMSO) δ 8.66 (s, 2H), 7.45 - 7.38 (m, 1H), 6.88 - 6.81 (m, 1H), 4.62 (s, 2H), 3.92 (s, 3H), 3.50 - 3.36 (m, 4H), 2.98 (s, 3H), 2.29 - 2.11 (m, 4H).

Example 8:

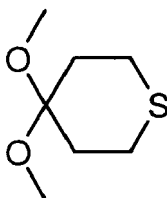
2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-1'-acetyl-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 108)

[0135]

[0136] A solution of 2-(3,5-dichloropyridine-4-yl)-1-(7-methoxy-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (10 mg, 24 μ mol), triethylamine (24 μ L, 171 μ mol) and acetic anhydride (12 μ L, 122 μ mol) in dichloromethane (200 μ L) was kept at room temperature overnight. Water (500 μ L) was added and the aqueous phase extracted with dichloromethane (3 x 500 μ L). The organic phase was dried over $MgSO_4$ and evaporated to dryness under reduced pressure. Standard HPLC purification afforded compound **108** (7.2 mg, 65%). 1H NMR (DMSO) δ 8.66 (s, 2H), 7.40 (d, 1H), 6.84 (d, 1H), 4.63 (s, 2H), 3.92 (s, 3H), 3.90 - 3.84 (m, 1H), 3.75 - 3.68 (m, 1H), 3.64 - 3.58 (m, 1H), 3.55 - 3.49 (m, 1H), 2.21 - 2.15 (m, 1H), 2.15 - 2.05 (m, 5H), 2.01 - 1.94 (m, 1H).

Preparation 10:

4,4-Dimethoxytetrahydro-(4H)-thiopyran (compound 510)

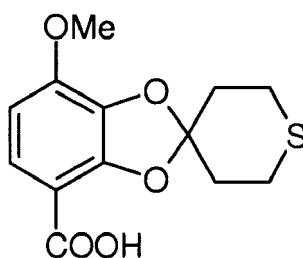
[0137]

[0138] A mixture of tetrahydro-(4H)-thiopyran-4-one (15.0 g, 129 mmol), trimethyl orthoformate (28.3 mL, 258 mmol) and para-toluenesulfonic acid monohydrate (67 mg, 0.35 mmol) in methanol (40 mL) was refluxed for 1 hour. The reaction mixture was cooled to room temperature, 1 M NaOMe (0.35 mL, 0.35 mmol) was added and excess methanol and trimethyl orthoformate was removed by distillation (atmospheric pressure). Further distillation under reduced pressure afforded 4,4-dimethoxytetrahydro-(4H)-thiopyran (20.7 g, 99%). 1H NMR (DMSO) δ 3.07 (s, 6H), 2.56 (m, 4H), 1.84 (m, 4H).

Preparation 11:

7-Methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran]-4-carboxylic acid (compound 511)

[0139]

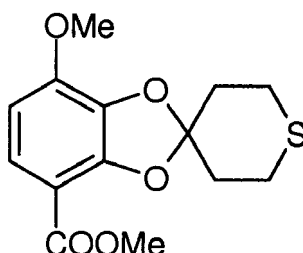


[0140] P-Toluenesulfonic acid (97 mg, 0.51 mmol) was added to 4,4-dimethoxy-tetrahydro-(4H)-thiopyran (20.7 g, 128 mmol) and the mixture was heated to 145 °C and kept at that temperature until approximately one equivalent of methanol (5.17 mL, 128 mmol) was distilled off. The mixture was then cooled to 130 °C and distillation under reduced pressure afforded 10.1 g of a 5:3 mixture of 5,6-dihydro-4-methoxy-(2H)-thiopyran [¹H NMR (DMSO) δ 4.87 (m, 1H), 3.44 (s, 3H), 3.15 (dt, 2H), 2.72 (t, 2H), 2.22 (m, 2H)] and 4,4-dimethoxytetrahydro-(4H)-thiopyran. Without further purification the mixture was added to 2,3-dihydroxy-4-methoxybenzoic acid (2.00 g, 10.9 mmol) and the suspension was exposed to microwave heating (180 °C, one hour) in a sealed reaction vessel. Ethyl acetate (100 mL) was added and the organic phase was first washed with 0.5 M HCl (40 mL) and then extracted with saturated aqueous NaHCO₃ (2 x 30 mL). The aqueous phase was washed with Et₂O (2 x 40 mL), acidified to pH = 1 with concentrated HCl and extracted with dichloromethane (2 x 30 mL). The organic phase was dried over MgSO₄. Evaporation under reduced pressure afforded 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-carboxylic acid (1.86 g, 61%). ¹³C NMR (DMSO) δ 164.9, 148.2, 146.6, 134.5, 123.8, 118.0, 107.2, 106.9, 56.1, 35.9, 25.4.

Preparation 12:

Methyl 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran]-4-carboxylate (compound 512)

[0141]

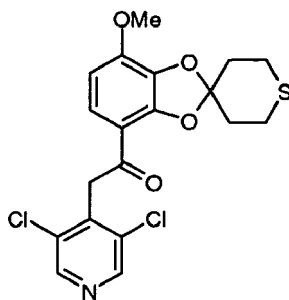


[0142] A suspension of 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-carboxylic acid (570 mg, 2.02 mmol), K₂CO₃ (558 mg, 4.04 mmol) and dimethyl sulphate (0.25 mL, 2.62 mmol) in acetone (14 mL) was stirred at 50 °C overnight. At room temperature water (30 mL) was added. The aqueous phase was extracted with dichloromethane (3 x 15 mL). The combined organic phase was dried over MgSO₄ and evaporated to dryness under reduced pressure. Standard silica gel column chromatography afforded methyl 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran]-4-carboxylate (407 mg, 68%). ¹³C NMR (DMSO) δ 163.8, 148.1, 146.9, 134.6, 123.4, 118.3, 107.1, 105.9, 56.1, 51.6, 35.9, 25.4.

Example 9:

2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (compound 110)

[0143]

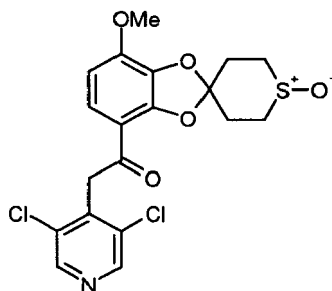


[0144] A solution of methyl 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-carboxylate (40 mg, 0.14 mmol) and 3,5-dichloro-4-picoline (33 mg, 0.20 mmol) in tetrahydrofuran (1.1 mL) was cooled to 0 °C. A 1.0 M solution of lithium bis(trimethylsilyl)amide in tetrahydrofuran (0.41 mL, 0.41 mmol) was added and the reaction mixture was allowed to reach room temperature overnight. Saturated aqueous NH₄Cl (20 mL) was added. The aqueous phase was extracted with dichloromethane (3 x 15 mL). The combined organic phase was dried over MgSO₄ and evaporated to dryness under reduced pressure. Standard HPLC purification afforded 2-(3,5-dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran]-4-yl)ethanone (38 mg, 67%). ¹³C NMR (DM-SO) δ 189.1, 148.2, 147.7, 147.0, 141.3, 134.5, 132.8, 122.0, 119.1, 113.0, 107.9, 56.3, 43.6, 35.9, 25.5.

Example 10:

2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran 1'-oxide]-4-yl)ethanone (compound 111)

[0145]

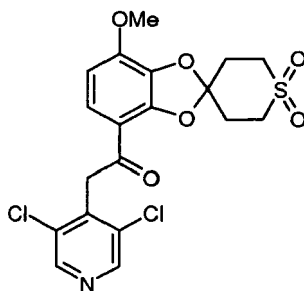


[0146] To a solution of 2-(3,5-dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (17 mg, 40 μmol) in dichloromethane (0.5 mL) was added first 0.25 M H₂O₂ in ethanol (128 μL, 32 μmol) and then secondly methyltrioxorhenium(VII) (1 mg, 4 μmol). The mixture was stirred at room temperature for two days and evaporated to dryness under reduced pressure. Standard HPLC purification afforded 2-(3,5-dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran 1'-oxide]-4-yl)ethanone (7 mg, 40%). ¹H NMR (DMSO) δ 8.66 (s, 2H), 7.42 (d, 1H), 6.85 (d, 1H), 4.63 (s, 2H), 3.93 (s, 3H), 3.17 - 2.94 (m, 4H), 2.69 - 2.55 (m, 2H), 2.36 - 2.24 (m, 2H).

Example 11:

2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran-1',1'-dioxide]-4-yl)ethanone (compound 112)

[0147]

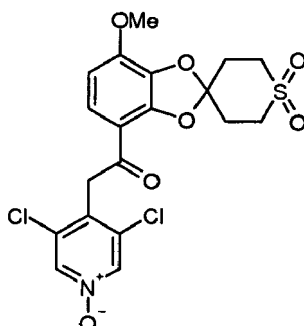


[0148] To a solution of 2-(3,5-dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran]-4-yl)ethanone (11 mg, 26 μ mol) in dichloromethane (0.25 mL) was added meta-chloroperbenzoic acid (10 mg, 58 μ mol) and the reaction was stirred at room temperature overnight. Saturated aqueous NaHCO_3 (1 mL) was added, and the aqueous phase was extracted with dichloromethane (2 x 2 mL). The combined organic phase was dried over MgSO_4 and evaporated to dryness under reduced pressure. Standard HPLC purification afforded 2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran-1',1'-dioxide]-4-yl)ethanone (5 mg, 42%). ^1H NMR (CDCl_3) δ 8.52 (s, 2H), 7.55 (d, 1H), 6.68 (d, 1H), 4.55 (s, 2H), 3.99 (s, 3H), 3.45 - 3.37 (m, 2H), 3.33 - 3.25 (m, 2H), 2.79 - 2.66 (m, 4H).

Example 12:

2-(3,5-Dichloro-1-oxido-pyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran-1',1'-dioxide]-4-yl)ethanone (compound 113)

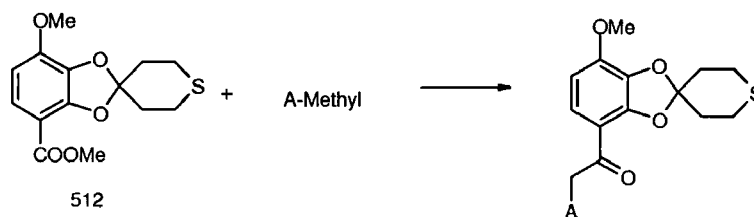
[0149]



[0150] To a solution of 2-(3,5-dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran]-4-yl)ethanone (10 mg, 23 μ mol) in absolute ethanol (2 mL) was added first H_2O_2 (100 μ L, 0.97 mmol) and then secondly methyltrioxorhenium(VII) (2 mg, 8 μ mol). The mixture was stirred at 40 $^\circ\text{C}$ overnight before 5% W/V aqueous NaHSO_3 (10 mL) was added. The aqueous phase was extracted with dichloromethane (3 x 20 mL). The combined organic phase was dried over MgSO_4 and evaporated to dryness under reduced pressure. Standard HPLC purification afforded 2-(3,5-dichloro-1-oxido-pyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran-1',1'-dioxide]-4-yl)ethanone (2.5 mg, 23%). ^1H NMR (DMSO) δ 8.64 (s, 2H), 7.42 (d, 1H), 6.85 (d, 1H), 4.59 (s, 2H), 3.92 (s, 3H), 3.51 (m, 2H), 3.31 (m, 2H), 2.59 (m, 4H).

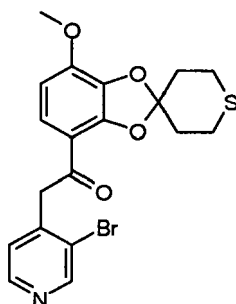
General procedure A:

[0151] LiHMDS (1M in THF, 3.0 eq) was added dropwise to a ice-cold solution of the ester **512** (1 eq) and A-Methyl (1.3 eq) in anhydrous THF. The reaction mixture was stirred at rt for 12 h, H_2O (10 mL) and sat. aq. NH_4Cl (20 mL) were added and then extracted with EtOAc (3 x 50 mL). The combined organic phases were dried (Na_2SO_4), filtered and concentrated under reduced pressure. The residue obtained was purified by flash column chromatography to afford ketone.

**Example 13:**

10 2-(3-bromopyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran]-4-yl)ethanone (compound 114).

15 **[0152]** Prepared according to General Procedure A using 18mg of 4-bromo-3-methylpyridine (yield: 40%)

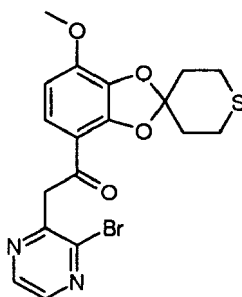


[0153] LC/MS (METHOD B): (m/z) 436.2 (MH⁺); RT = 4.17 min; purity (UV) = 100%

Example 14:

30 2-(3-Bromo-pyrazin-2-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (compound 115).

35 **[0154]** Prepared according to General Procedure A using 17mg of 2-bromo-3-methylpyrazine (yield: 9%)

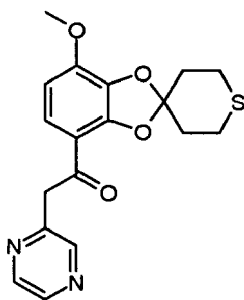


[0155] LC/MS (METHOD B): (m/z) 437.2; 439.22 (MH⁺); RT = 4.22 min; purity (UV) = 100%

Example 15:

50 2-(pyrazin-2-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (compound 116).

55 **[0156]** Prepared according to General Procedure A using 16mg of 2-methylpyrazine (yield: 52%)

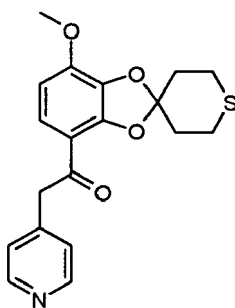


[0157] LC/MS (METHOD B): (m/z) 359.3 (MH⁺); RT = 3.33 min; purity (UV) = 100%

Example 16:

2-(pyridin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran]-4-yl)ethanone (compound 117).

[0158] Prepared according to General Procedure A using 18mg of 4-methylpyridine (yield: 13%)

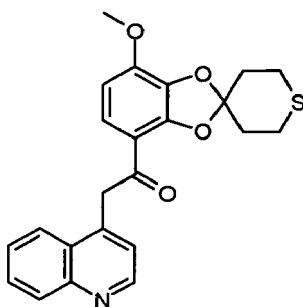


[0159] LC/MS (METHOD B): (m/z) 358.3 (MH⁺); RT = 2.50 min; purity (UV) = 100%

Example 17:

2-(quinolin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran]-4-yl)ethanone (compound 118).

[0160] Prepared according to General Procedure A using 16mg of 4-methylquinoline (yield: 12%)

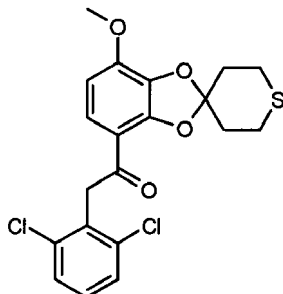


[0161] LC/MS (METHOD B): (m/z) 408.3 (MH⁺); RT = 3.33 min; purity (UV) = 100%

Example 18:

2-(2,6-Dichloro-phenyl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4*H*)-thiopyran]-4-yl)ethanone (compound 119).

[0162] Prepared according to General Procedure A using 15mg of 2,6-dichlorotoluene (yield: 8%)

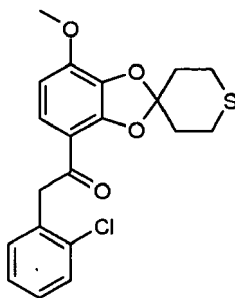


[0163] LC/MS (METHOD B): (m/z) 425.24 (MH⁺); RT = 5.28 min; purity (UV) = 100%

Example 19:

2-(2-chloro-phenyl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4*H*)-thiopyran]-4-yl)ethanone (compound 120)

[0164]



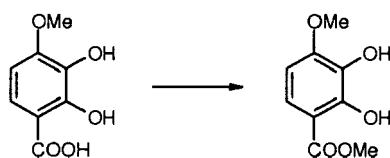
[0165] A solution of Methyl 7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4*H*)-thiopyran]-4-carboxylate (14 mg) in dry THF (500mL) was cooled to 0°C under Ar. 2-chlorobenzyl magnesiumchloride in diethyl ether (0.25M solution, 189μL) was added and the cooling was removed. After 2h at r.t. an additional portion of 2-chlorobenzyl magnesiumchloride in diethyl ether (0.25M solution, 189μL) was added. The mixture was stirred for 18h, added water and extracted with ethyl acetate. The combined organic phase was dried over MgSO₄ and evaporated to dryness under reduced pressure. Standard HPLC purification afforded compound **120** (9.2%).

[0166] LC/MS (METHOD B): (m/z) 391.22 (MH⁺); RT = 4.90 min; purity (UV) = 100%

Preparation 13:

Methyl 2,3-dihydroxy-4-methoxybenzoate (compound 513)

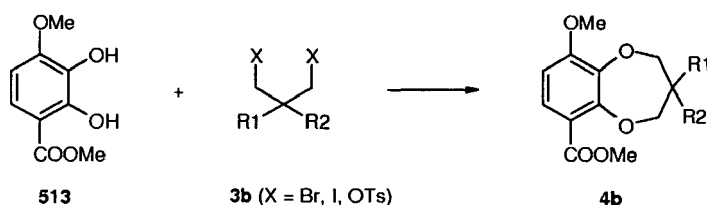
[0167]

**513**

[0168] A solution of commercially available 2,3-dihydroxy-4-methoxybenzoic acid (11.6 g, 63 mmol) in anhydrous MeOH (150 mL) was cooled in an ice-bath and conc. H_2SO_4 (8 mL) added dropwise. The reaction mixture was refluxed for 12 h, then cooled to rt and the solvent was removed under reduced pressure. H_2O (100 mL) and sat aq NaHCO_3 (50 mL) were added and extracted with EtOAc (3 x 100 mL). The combined organic phase was dried (Na_2SO_4), filtered and concentrated *in vacuo* to afford compound **513** as a pale yellow solid, which was used in the next step without further purification. LC-MS: $R_T = 2.31$ min.; m/z 197.3 (M-H)⁻. ^1H NMR (CDCl_3): δ 10.83 (1H, s), 7.41 (1H, d, J 9.0), 6.50 (1H, d, J 8.9), 5.45 (1H, s), 3.94 (3H, s), 3.93 (3H, s).

General procedure B:

[0169]



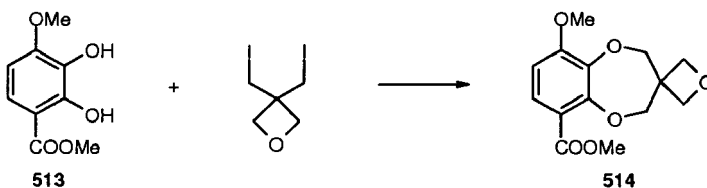
[0170] To a stirred solution of compound **513** and **3b** (1.1 eq) in anhydrous DMSO was added K_2CO_3 (2.5 eq) and the mixture stirred at 100 °C for 4-12 h, under inert atmosphere. After cooling to rt, ice-water mixture was added, stirred for 15 min. and then extracted with EtOAc (3 x 50 mL). The combined organic phases were dried (Na_2SO_4), filtered and concentrated under reduced pressure. The residue obtained was purified by flash column chromatography.

[0171] Using general procedure B the following compounds were obtained:

Preparation 14:

9-Methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-oxetane]-6-carboxylic acid methyl ester (compound **514**)

[0172]



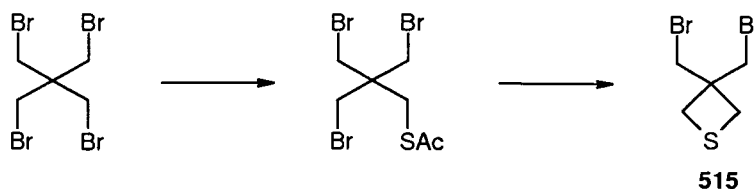
[0173] Following the general procedure, dialkylation of **513** (198 mg, 1 mmol) with commercial 3,3-bis(iodomethyl)oxetane (372 mg, 1.1 mmol) in DMSO (5 mL) in the presence of K_2CO_3 (345 mg, 2.5 mmol) afforded compound **514** as a white solid material after purification by column chromatography (50-65% EtOAc in light petroleum). LC-MS: $R_T = 2.40$ min.; m/z 281.26 (M+H)⁺. ^1H NMR (CDCl_3): δ 7.49 (1H, d, J 8.8), 6.62 (1H, d, J 8.8), 4.61 (2H, d, J 6.8), 4.58 (2H, d, J 6.8), 4.48 (4H, s), 3.90 (3H, s), 3.87 (3H, s).

Preparation 15:

9-Methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietane]-6-carboxylic acid methyl ester (compound 516)

Step A:

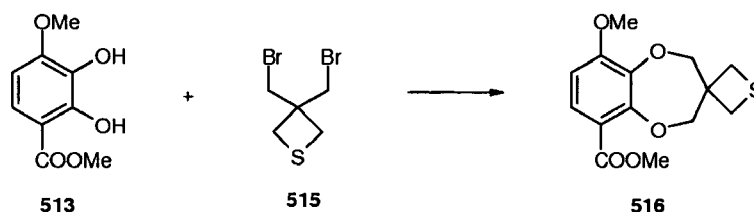
[0174]



[0175] 3,3-Bis(bromomethyl)thietane **515** was obtained in a two step process from 1,3-dibromo-2,2-bis(bromomethyl)propane following the literature procedure (Petrukina, M. A.; Henck, C.; Li, B.; Block, E.; Jin, J.; Zhang, S-Z.; Clerac, R. Inorg. Chem. 2005, 44, 77-84). Accordingly, a mixture of 1,3-dibromo-2,2-bis(bromomethyl)propane (7.76 g, 20 mmol) and KSAc (2.28 g, 20 mmol) in anhydrous THF (30 mL) was refluxed for 30 h. The ppt was filtered off and the filtrate concentrated and the residue obtained was purified by flash column chromatography (10-25% EtOAc in light petroleum) to afford thioacetic acid(2,2-bis(bromomethyl)-3-bromopropyl)ester as a pale yellow solid material. A mixture of thioacetic acid(2,2-bis(bromomethyl)-3-bromopropyl)ester (1.53 g, 4 mmol) and NaOMe (324 mg, 6 mmol) in anhydrous MeOH (10 mL) was stirred at 0 °C for 2 h. MeOH was removed *in vacuo*, coevaporated with toluene (2 x 2 mL) and the residue obtained was filtered through a short pad of silica gel to give 3,3-bis(bromomethyl)thietane **515** as a thick oil, which was used without further purification.

Step B:

[0176]



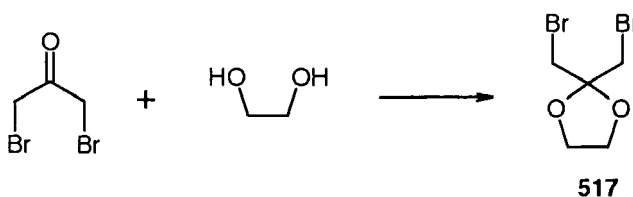
Following the general procedure, dialkylation of **513** (665 mg, 3.36 mmol) with **515** (962 mg, 3.7 mmol) in DMSO (15 mL) in the presence of K_2CO_3 (1.16 g, 8.4 mmol) afforded **516** as a white solid material after purification by column chromatography (40-60% EtOAc in light petroleum). LC-MS: $R_T = 3.17$ min.; m/z 297.19 ($M+H$)⁺. 1H NMR ($CDCl_3$): δ 7.49 (1H, d, J 8.8), 6.63 (1H, d, J 8.8), 4.30 (2H, s), 4.28 (2H, s), 3.90 (3H, s), 3.87 (3H, s), 3.11 (4H, s).

Preparation 16:

9-Methoxy-spiro[2H-1,5-benzodioxepin-3(4H),2'-(1,3-dioxolane)]-6-carboxylic acid methyl ester (compound 518)

Step A:

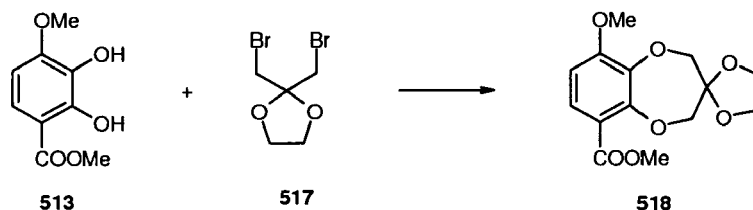
[0177]



[0178] 2,2-Bis(bromomethyl)-1,3-dioxolane **517** was obtained from dibromoacetone following the literature procedure (Valentin, M-L.; Bolte, J. Bull. Soc. Chim. Fr. 1995, 132, 1167-71). Accordingly, a solution of dibromoacetone (4.04 g, 18.7 mmol), ethylene glycol (2.32 g, 37.4 mmol) and *p*-TsOH (25 mg) in benzene (70 mL) was refluxed for 12 h, with azeotropic removal of water. The reaction mixture was concentrated under reduced pressure, Et₂O (50 mL) was added and the organic layer was washed with H₂O (2 x 50 mL), dried (Na₂SO₄), filtered and concentrated under reduced pressure. The residue obtained was purified by flash column chromatography (7-10% EtOAc in light petroleum) to afford 2,2-bis(bromomethyl)-1,3-dioxolane (**517**) as colourless liquid.

Step B:

[0179]



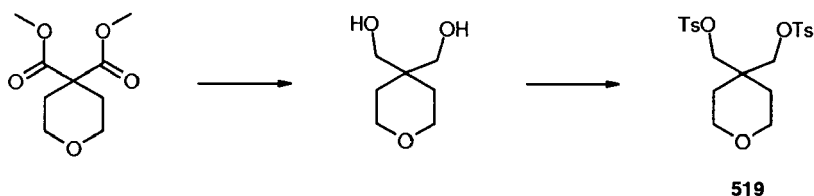
Following the general procedure, dialkylation of **513** (396 mg, 2 mmol) with **517** (572 mg, 2.2 mmol) in DMSO (10 mL) in the presence of K₂CO₃ (690 mg, 5 mmol) afforded **518** as a white solid material after purification by column chromatography (45-60% EtOAc in light petroleum). LC-MS: R_T = 2.70 min.; *m/z* 297.18 (M+H)⁺, 319.16 (M+Na)⁺. ¹H NMR (DMSO-*d*₆): δ 7.34 (1H, d, *J* 8.8), 6.76 (1H, d, *J* 8.8), 4.11 (2H, s), 4.09 (2H, s), 3.94 (4H, s), 3.80 (3H, s), 3.75 (3H, s).

Preparation 17:

9-Methoxy-spiro[2H-1,5-benzodioxepin-3(4H),4'-tetrahydropyran]-6-carboxylic acid methyl ester (compound **520**)

Step A:

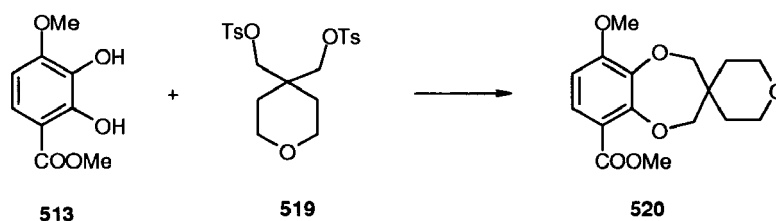
[0180]



[0181] 4,4-Bis(*p*-toluenesulphonyloxymethyl)tetrahydropyran (**519**) was obtained in a two step process from commercial tetrahydropyran-4,4-dicarboxylic acid dimethyl ester. Accordingly, to an ice-cooled solution of the diester (3.03 g, 15 mmol) in anhydrous toluene (45 mL) was added dropwise synhydride (70% in toluene, 19.5 mL, 66 mmol) and the mixture stirred at 120 °C for 3 h. The mixture was cooled to rt, H₂O (50 mL) was added slowly and concentrated over silica gel (35 g). Quick column chromatography with a linear gradient of MeOH in CH₂Cl₂ afforded 4,4-(bishydroxymethyl)tetrahydropyran as white solid material. A mixture of diol (1.46 g, 10 mmol) and TsCl (4.77 g, 25 mmol) in anhydrous pyridine (25 mL) was stirred at rt for 48 h. Solvent was removed in *vacuo* and co-evaporated with toluene (3 x 10 mL). CH₂Cl₂ (100 mL) and sat aq NaHCO₃ (100 mL) were added, the phases were separated and the organic phase was dried (Na₂SO₄), filtered and concentrated under reduced pressure. The residue obtained was flash chromatographed (35-45% EtOAc in light petroleum) to give 4,4-bis(*p*-toluenesulphonyloxymethyl)tetrahydropyran (**519**) as a white solid material.

Step B:

[0182]



[0183] Following the general procedure, dialkylation of compound **513** (95 mg, 0.48 mmol) with compound **519** (240 mg, 0.53 mmol) in DMSO (3 mL) in the presence of K_2CO_3 (166 mg, 1.2 mmol) afforded compound **520** as a white solid material after purification by column chromatography (45-65% EtOAc in light petroleum). LC-MS: $R_T = 2.40$ min.; 1H NMR (DMSO- d_6): δ 7.34 (1H, d, J 8.8), 6.78 (1H, d, J 8.8), 3.99 (4H, s), 3.80 (3H, s), 3.76 (3H, s), 3.61 (4H, t, J 5.5), 1.54 (4H, t, J 5.5).

Preparation 18:

9-Methoxy-2',2'-dimethyl-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxane]-6-carboxylic acid methyl ester (compound **522**)

Step A:

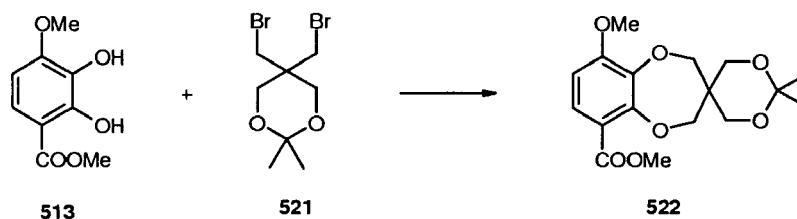
[0184]



[0185] A solution of 2,2-bis(bromomethyl)-1,3-propanediol (5.0 g, 19.1 mmol), anhydrous acetone (20 mL) and p -TsOH (100 mg) in benzene (75 mL) was refluxed for 12 h, with azeotropic removal of water. The reaction mixture was concentrated under reduced pressure, EtOAc (100 mL) was added and the organic layer was washed successively with H_2O (3 x 30 mL) and brine (30 mL), dried (Na_2SO_4), filtered and concentrated under reduced pressure. The white solid obtained was triturated with n -pentane (20 mL) to afford 5,5-bis(bromomethyl)-2,2-dimethyl-[1,3]dioxane (**521**) as colourless crystals.

Step B:

[0186]



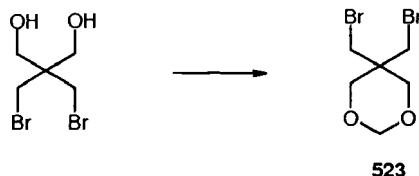
[0187] Following the general procedure, dialkylation of compound **513** (1.19 g, 6 mmol) with compound **521** (2.0 g, 6.62 mmol) in DMSO (30 mL) in the presence of K_2CO_3 (2.07 g, 15 mmol) afforded compound **522** as a white solid material after purification by column chromatography (30-40% EtOAc in light petroleum). LC-MS: $R_T = 2.99$ min.; m/z 339.31 ($M+H$) $^+$, 361.25 ($M+Na$) $^+$. 1H NMR ($CDCl_3$): δ 7.45 (1H, d, J 8.8), 6.59 (1H, d, J 8.8), 4.22 (2H, s), 4.18 (2H, s), 3.89 (3H, s), 3.86 (7H, s), 1.43 (6H, s).

Preparation 19:

9-Methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxane]-6-carboxylic acid methyl ester (compound 524)

Step A:

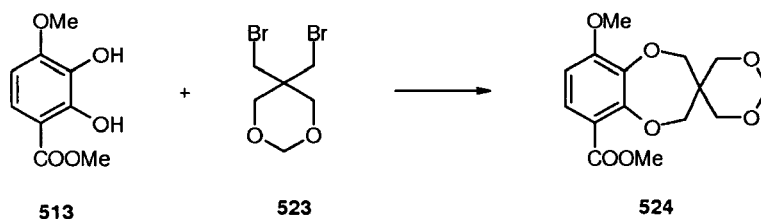
[0188]



[0189] 5,5-Bis(bromomethyl)-[1,3]dioxane (**523**) was obtained from 2,2-bis(bromomethyl)-1,3-propanediol following the literature procedure (Bitha, P.; Carvajal, S. G.; Citarella, R. V.; Delos Santos, E. F.; Durr, F. E.; Hlavka, J. J.; Lang, S. A., Jr.; Lindsay, H. L.; Thomas, J. P.; Wallace, R. E.; Yang-I, L. J. *Med. Chem.* 1989, 32(9), 2063-7 and Mitkin, O. D.; Wan, Y.; Kurchan, A. N.; Kutateladze, A. G. *Synthesis*, 2001, (8), 1133-42). Accordingly, a solution of 2,2-bis(bromomethyl)-1,3-propanediol (2.5 g, 9.55 mmol), formaldehyde (37% aqueous solution, 3.5 mL) and conc. HCl (2.0 mL) was refluxed for 12 h. After being cooled to rt, H₂O (25 mL) was added to the reaction mixture which was extracted with CH₂Cl₂ (2 x 25 mL). The combined organic layer successively washed with sat. aq. Na₂CO₃ (25 mL) and H₂O (25 mL), dried (Na₂SO₄), filtered and concentrated under reduced pressure. The colourless liquid obtained was found to be >95% pure by ¹H NMR, and it was used without further purification.

Step B:

[0190]



[0191] Following the general procedure, dialkylation of compound **513** (198 mg, 1 mmol) with compound **523** (301 mg, 1.1 mmol) in DMSO (5 mL) in the presence of K₂CO₃ (345 mg, 2.5 mmol) afforded compound **524** as a white solid material after purification by column chromatography (50-65% EtOAc in light petroleum). LC-MS: R_T = 2.65 min.; *m/z* 311.23 (M+H)⁺. ¹H NMR (DMSO-*d*₆): δ 7.37 (1H, d, *J* 8.8), 6.81 (1H, d, *J* 8.8), 4.80 (1H, d, *J* 6.4), 4.78 (1H, d, *J* 6.4), 4.07 (2H, s), 4.01 (2H, s), 3.81 (7H, s), 3.76 (3H, s).

Preparation 20:

9-Methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dithiane]-6-carboxylic acid methyl ester (compound 526)

Step A:

[0192]



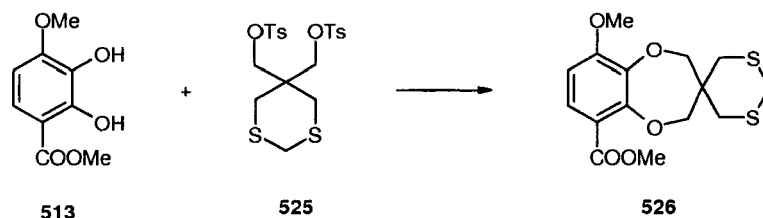
[0193] 4,4-Bis(hydroxymethyl)-[1,3]dithiane was obtained in a two step process from compound **523** following the literature procedure (Mitkin, O. D.; Wan, Y.; Kurchan, A. N.; Kutateladze, A. G. Synthesis, 2001, (8), 1133-42). Accordingly, a mixture of compound **523** (1.36 g, 5 mmol) and KSac (1.71 g, 15 mmol) in anhydrous DMF (10 mL) was stirred at rt for 30 h. The solvent was removed under reduced pressure and coevaporated with toluene (3 x 5 mL). Ice-water mixture was added and then extraction was performed with diisopropyl ether (2 x 25 mL). The combined organic phases were dried (Na₂SO₄), filtered and concentrated to afford 5,5-bis(acetylthiomethyl)-[1,3]dioxane (1.32 g, quantitative) as a pale yellow viscous oil. It was found to be >95% pure by ¹H NMR and it was used without further purification.

[0194] A solution of 5,5-bis(acetylthiomethyl)-[1,3]dioxane (1.32 g, 5 mmol) in aq HCl (2N, 25 mL) was refluxed for 16 h and then cooled to rt. The mixture was made alkaline by dropwise addition of aq Na₂CO₃ (2M) and then extracted with CH₂Cl₂ (3 x 40 mL). The combined organic layer dried (Na₂SO₄), filtered and concentrated under reduced pressure to a white solid material, which was triturated with hot *n*-hexane-diisopropyl ether (2:1, 15 mL). The white solid was filtered to provide 4,4-bis(hydroxymethyl)-[1,3]dithiane, which was used without further purification.

[0195] A mixture of 4,4-bis(hydroxymethyl)-[1,3]dithiane (405 mg, 2.25 mmol) and TsCl (1.29 g, 6.75 mmol) in anhydrous pyridine (4 mL) was stirred at rt for 48 h. Solvent was removed in *vacuo* and co-evaporated with toluene (3 x 3 mL). CH₂Cl₂ (40 mL) and sat aq NaHCO₃ (40 mL) were added, the phases separated and the organic phase dried (Na₂SO₄), filtered and concentrated under reduced pressure. The residue obtained was flash chromatographed (35-55% EtOAc in light petroleum) to provide 4,4-bis(*p*-toluenesulphonyloxymethyl)-[1,3]dithiane (**525**) as a white solid material.

Step B:

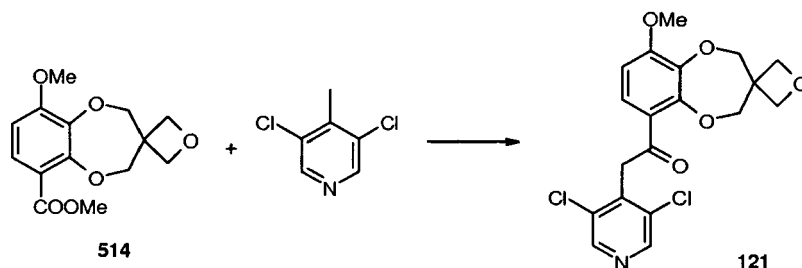
[0196]



Example 20

2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-oxetane]-6-yl}ethanone (compound 121)

[0200]

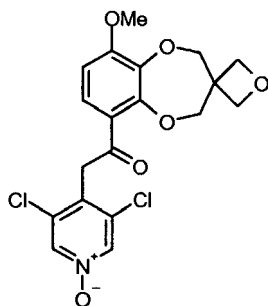


[0201] Following the general procedure, condensation of 3,5-dichloro-4-methylpyridine (169 mg, 1.04 mmol) with compound **514** (224 mg, 0.8 mmol) in THF (4 mL) in the presence of LiHMDS (2.4 mL, 2.4 mmol) afforded compound **121** as a white solid material after purification by column chromatography (65-80% EtOAc in light petroleum). LC-MS: $R_T = 3.24$ min.; m/z 410.07 ($M+H$)⁺, 408.19, 410.22 ($M-H$)⁻. ¹H NMR (DMSO-*d*₆): δ 8.66 (2H, s), 7.45 (1H, d, *J* 8.9), 6.91 (1H, d, *J* 9.0), 4.66 (2H, s), 4.57 (2H, s), 4.50 (2H, d, *J* 6.5), 4.46 (2H, d, *J* 6.5), 4.36 (2H, s), 3.85 (3H, s).

Example 21

2-(3,5-Dichloro-1-oxido-pyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-oxetane]-6-yl}ethanone (compound 122)

[0202]



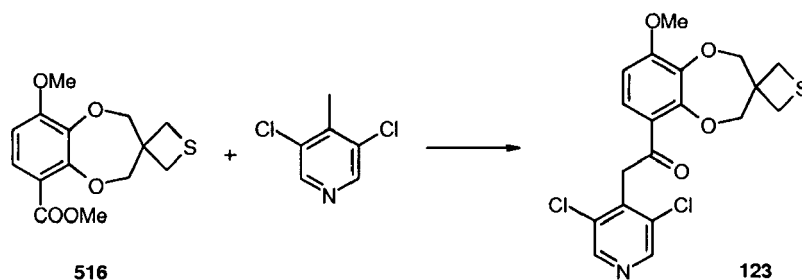
[0203] To a solution of 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-oxetane]-6-yl}ethanone [**121**] (20.5 mg, 50 μ mol) in CH₂Cl₂ (1 mL) was added 30% H₂O₂ (15 μ L) and methyltrioxorhenium(VII) (5 mg). The mixture was stirred for 18h, added MnO₂ (5mg) and was stirred for another hour. CH₂Cl₂ (10mL) was added and the organic phase was washed with water. The combined organic phase was dried over MgSO₄ and evaporated to dryness under reduced pressure. Standard HPLC purification afforded 14mg of the product.

[0204] LC/MS (METHOD B): (m/z) 426.18; 428.20 (MH^+); $R_T = 2.42$ min; purity (UV) = 100%

Example 22:

2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietane]-6-yl}ethanone (compound 123)

[0205]

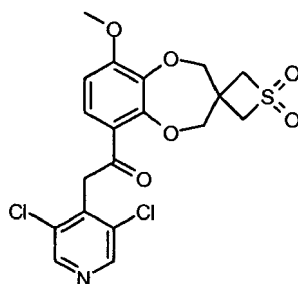


[0206] Following the general procedure, condensation of 3,5-dichloro-4-methylpyridine (366 mg, 2.26 mmol) with compound **516** (516 mg, 1.74 mmol) in THF (10 mL) in the presence of LiHMDS (5.2 mL, 5.2 mmol) afforded compound **123** as a white solid material after purification by column chromatography (55-65% EtOAc in light petroleum). LC-MS: $R_T = 4.23$ min.; m/z 426.24, 428.25 ($M+H$)⁺, 424.23 ($M-H$)⁻. ¹H NMR ($CDCl_3$): δ 8.51 (2H, s), 7.57 (1H, d, J 9.2), 6.71 (1H, d, J 9.2), 4.64 (2H, s), 4.46 (2H, s), 4.33 (2H, s), 3.93 (3H, s), 3.20 (2H, d, J 9.9), 3.12 (2H, d, J 9.9).

Example 23:

2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietane-1',1'-dioxide]-6-yl}ethanone (compound 124)

[0207]



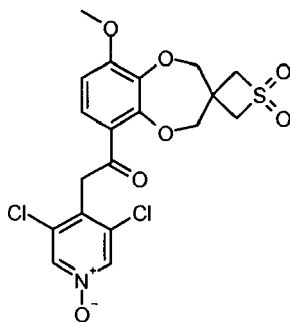
[0208] To a solution of 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietane]-6-yl}ethanone [**123**] (42.6 mg) in CH_2Cl_2 (1 mL) was added 30% H_2O_2 (38 μ L) and methyltrioxorhenium(VII) (5 mg). The mixture was stirred for 18h and subsequently washed with water. The organic phase was dried over $MgSO_4$ and evaporated to dryness under reduced pressure. Standard HPLC purification afforded 6mg of the product.

[0209] LC/MS (METHOD B): (m/z) 458.13, 460.10, 462.14 (MH)⁺; $RT = 3.30$ min; purity (UV) = 100%

Example 24:

2-(3,5-Dichloropyridin-1-oxido-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietane-1',1'-dioxide]-6-yl}ethanone (compound 125).

[0210]



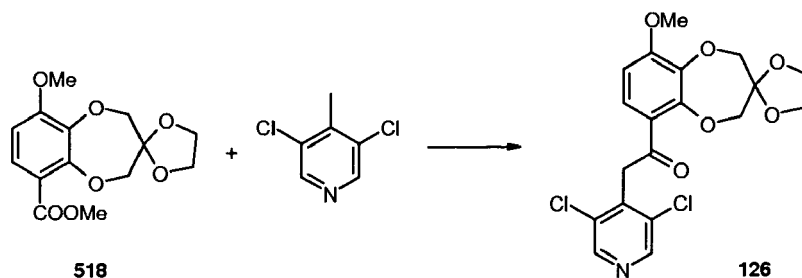
[0211] To a solution of 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietane]-6-yl} ethanone **[123]** (39.6 mg) in CH₂Cl₂ (1 mL) was added 30% H₂O₂ (76 μ L) and methyltrioxorhenium(VII) (5 mg). The mixture was stirred for 18h, and was subsequently added water. The organic phase was dried over MgSO₄ and evaporated to dryness under reduced pressure. Standard HPLC purification afforded 8.4mg of the product.

[0212] LC/MS (METHOD B): (*m/z*) 474.17, 476.16, 478.18 (MH⁺); RT = 2.39 min; purity (UV) = 100%

Example 25:

2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),2'-(1,3-dioxolane)]-6-yl}ethanone (compound **126**)

[0213]

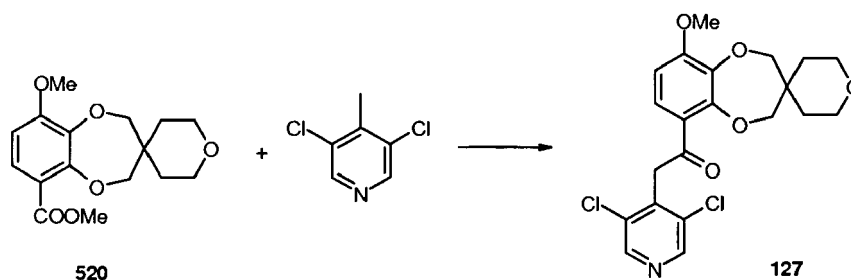


[0214] Following the general procedure, condensation of 3,5-dichloro-4-methylpyridine (53 mg, 0.33 mmol) with compound **518** (74 mg, 0.25 mmol) in THF (1.5 mL) in the presence of LiHMDS (0.75 mL, 0.75 mmol) afforded compound **126** as a white solid material after purification by column chromatography (60-80% EtOAc in light petroleum). LC-MS: *R_T* = 3.63 min.; *m/z* 426.18, 428.16 (M+H)⁺. ¹H NMR (DMSO-*d*₆): δ 8.64 (2H, s), 7.39 (1H, d, *J* 8.8), 6.86 (1H, d, *J* 8.9), 4.60 (2H, s), 4.34 (2H, s), 4.17 (2H, s), 3.97 (4H, s), 3.84 (3H, s).

Example 26:

2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),4'-tetrahydropyran]-6-yl}ethanone (compound **127**)

[0215]

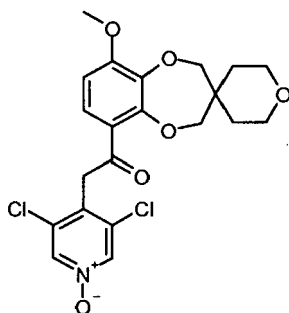


[0216] Following the general procedure, condensation of 3,5-dichloro-4-methylpyridine (59 mg, 0.36 mmol) with compound **520** (85 mg, 0.28 mmol) in THF (1.5 mL) in the presence of LiHMDS (0.85 mL, 0.85 mmol) afforded compound **127** as a white solid material after purification by column chromatography (60-70% EtOAc in light petroleum). LC-MS: *R_T* = 3.70 min.; *m/z* 438.21, 440.21 (M+H)⁺, 436.30, 438.27 (M-H)⁻. ¹H NMR (DMSO-*d*₆): δ 8.65 (2H, s), 7.42 (1H, d, *J* 8.9), 6.87 (1H, d, *J* 9.0), 4.63 (2H, s), 4.25 (2H, s), 4.07 (2H, s), 3.84 (3H, s), 3.64 (4H, t, *J* 5.3), 1.60 (4H, t, *J* 5.3).

Example 27:

2-(3,5-Dichloro-1-oxido-pyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),4'-tetrahydropyran]-6-yl}ethanone (compound **128**)

[0217]



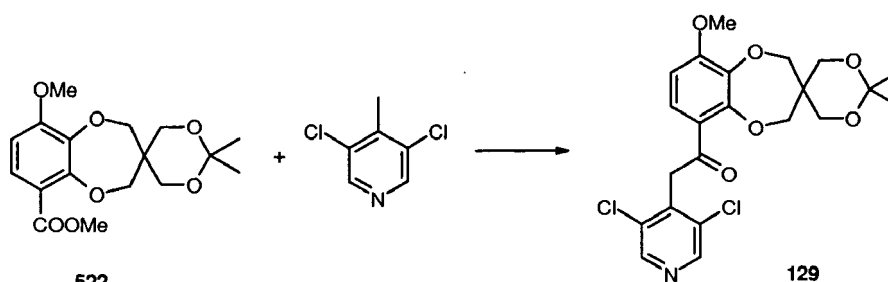
[0218] To a solution 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),4'-tetrahydropyran]-6-yl}ethanone **[127]** (66 mg, 150 μ mol) in CH_2Cl_2 (2 mL) was added 30% H_2O_2 (45 μ L) and methyltrioxorhenium(VII) (10 mg). The mixture was stirred for 18h, added MnO_2 (10mg) and was stirred for another hour. CH_2Cl_2 (10mL) was added and the organic phase was washed with water. The combined organic phase was dried over MgSO_4 and evaporated to dryness under reduced pressure. Standard HPLC purification afforded 5.6mg of the product.

[0219] LC/MS (METHOD B): (m/z) 454.32; 456.32 (MH^+); RT = 2.80 min; purity (UV) = 100%

[0220] Example 28:

2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-2',2'-dimethyl-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxane]-6-yl}ethanone (compound 129)

[0221]

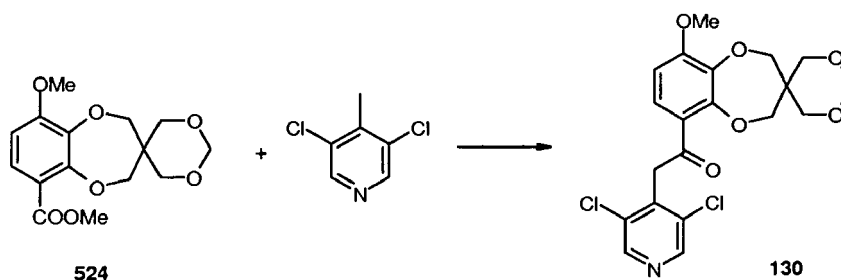


[0222] Following the general procedure, condensation of 3,5-dichloro-4-methylpyridine (93 mg, 0.58 mmol) with compound **522** (150 mg, 0.44 mmol) in THF (2 mL) in the presence of LiHMDS (1.3 mL, 1.3 mmol) afforded compound **129** as a white solid material after purification by column chromatography (55-60% EtOAc in light petroleum). LC-MS: R_T = 3.95 min.; m/z 468.19, 470.23 ($\text{M}+\text{H}^+$), 466.36, 468.33 ($\text{M}-\text{H}^-$). ^1H NMR ($\text{DMSO}-d_6$): δ 8.65 (2H, s), 7.42 (1H, d, J 8.8), 6.89 (1H, d, J 9.1), 4.63 (2H, s), 4.33 (2H, s), 4.08 (2H, s), 3.84 (3H, s), 3.82 (4H, s), 1.39 (3H, s), 1.37 (3H, s).

Example 29:

2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxane]-6-yl}ethanone (compound 130)

[0223]

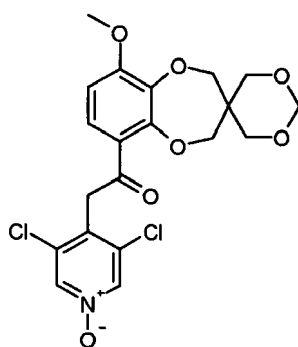


[0224] Following the general procedure, condensation of 3,5-dichloro-4-methylpyridine (211 mg, 1.3 mmol) with compound **524** (310 mg, 1.0 mmol) in THF (6 mL) in the presence of LiHMDS (3 mL, 3.0 mmol) afforded compound **130** as a white solid material after purification by column chromatography (60-80% EtOAc in light petroleum). LC-MS: R_T = 3.64 min.; m/z 440.16, 442.17 ($M+H$)⁺, 438.27, 440.29 ($M-H$)⁻. ¹H NMR (DMSO-*d*₆): δ 8.65 (2H, s), 7.43 (1H, d, *J* 8.9), 6.89 (1H, d, *J* 8.9), 4.84 (1H, d, *J* 6.1), 4.79 (1H, d, *J* 6.1), 4.64 (2H, s), 4.34 (2H, s), 4.08 (2H, s), 3.90 (2H, d, *J* 11.4), 3.85 (3H, s), 3.82 (2H, d, *J* 11.2).

Example 30:

2-(3,5-Dichloro-1-oxido-pyridine-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxane]-6-yl}ethanone (compound 131)

[0225]



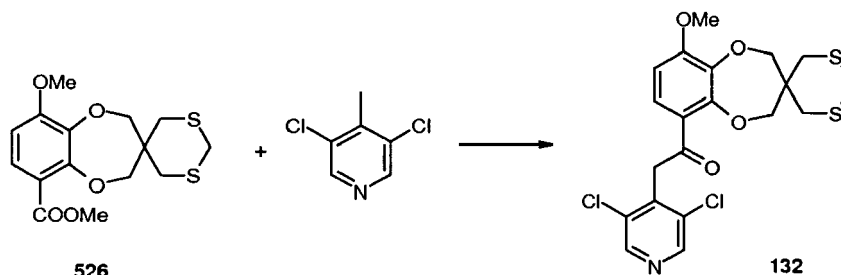
[0226] To a solution of 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxane]-6-yl}ethanone [130] (100 mg) in CH₂Cl₂ (2 mL) was added 30% H₂O₂ (120 μ L) and methyltrioxorhenium(VII) (5 mg). The mixture was stirred for 18h and subsequently washed with water. The organic phase was dried over MgSO₄ and evaporated to dryness under reduced pressure. Standard HPLC purification afforded 85mg of the product.

[0227] LC/MS (METHOD B): (m/z) 456.23 (MH^+); R_T = 2.55 min; purity (UV) = 95%

Example 31:

2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dithiane]-6-yl}ethanone (compound 132)

[0228]



[0229] Following the general procedure, condensation of 3,5-dichloro-4-methylpyridine (154 mg, 0.95 mmol) with compound **526** (250 mg, 0.73 mmol) in THF (4 mL) in the presence of LiHMDS (2.2 mL, 2.2 mmol) afforded compound **132** as a white solid material after purification by column chromatography (45-55% EtOAc in light petroleum). LC-MS: R_T = 4.39 min.; m/z 472.15, 474.14 ($M+H$)⁺. ¹H NMR (DMSO-*d*₆): δ 8.65 (2H, s), 7.42 (1H, d, *J* 8.9), 6.88 (1H, d, *J* 8.9), 4.64 (2H, s), 4.43 (2H, s), 4.24 (2H, s), 3.85 (3H, s), 3.84 (2H, s), 2.91 (4H, s).

Example 32:

PDE4 assay

[0230] Human recombinant PDE4 (Genbank accession no NM_006203) was incubated for 1 hour, with the test compound at concentrations up to 10 μ M, with cAMP (1x10⁻⁵M), and with a low amount (0.021 MBq) of radioactively labelled cAMP. At the end of the incubation, the cleavage of the substrate was evaluated by the binding of the AMP product to SPA beads, which generate chemoluminescence when bound to the radioactive tracer. The AMP product inhibited the binding of the radioactive tracer to the beads, and the luminescent signal was competed. The results were calculated as the molar concentrations resulting in 50% inhibition of the substrate cleavage compared to controls samples, and are expressed as IC₅₀ (M).

[0231] The results are shown in Table 1 below.

Table 1

Compound	IC ₅₀ (PDE4)
101	26nM
102	41nM
103	26nM
104	10nM
105	310nM
106	127nM
107	590nM
108	220nM
110	11nM
111	18nM
112	29nM
113	53nM
114	45nM
115	1410nM
116	1290nM
117	207nM
118	1800nM
119	71nM
120	310nM
121	52nM
122	50nM
123	17nM
124	104nM
125	200nM
126	67nM
127	22nM
128	11nM
129	
130	40nM

EP 2 114 951 B9

(continued)

Compound	IC ₅₀ (PDE4)
131	43nM
132	26nM

Example 33:

[0232] TNF- α release

[0233] Human peripheral blood mononuclear cells (PBMC) were isolated from buffy coats. The blood is mixed with saline at a ratio of 1:1, and the PBMC were isolated using Lymphoprep tubesTM (Nycomed, Norway). The PBMC were suspended in RPMI1640 with 2 % foetal calf serum (FCS), pen/strep and 2 mM L-glutamine at a concentration of 5 x 10⁵ c/ml. The cells were pre-incubated for 30 minutes with the test compounds in 96 well tissue culture plates and stimulated for 18 hours with lipopolysaccharide 1 mg/ml (Sigma). The level of TNF- α was measured in the culture supernatant by enzyme immunoassays using primary and secondary biotinylated antibodies from R&D systems. Results are expressed as IC₅₀ values calculated from inhibition curves using as positive controls the secretion in LPS stimulated wells and as negative controls the secretion in unstimulated cells.

[0234] The results are shown in Table 2 below.

Table 2

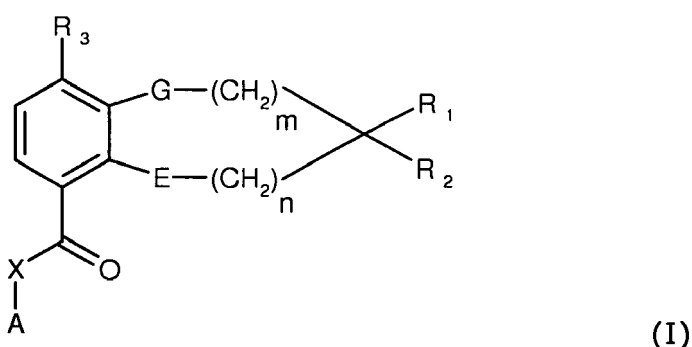
Compound	IC ₅₀ (TNF- α)
101	28nM
102	88nM
103	23nM
104	44nM
105	800nM
106	158nM
107	910nM
108	167nM
110	17nM
111	19nM
112	34nM
113	82nM
114	95nM
115	471nM
116	2940nM
117	286nM
118	5520nM
119	508nM
120	1250nM
121	64nM
122	20nM
123	35nM
124	135nM
125	135nM

(continued)

Compound	IC ₅₀ (TNF- α)
126	111nM
127	27nM
128	9nM
130	50nM
131	16nM
132	29nM

Claims

1. A compound of general formula I



wherein m and n independently represent 0, 1, 2, 3, 4, 5, 6, or 7;

and wherein G and E independently represent sulphur, oxygen, -N(R₅)-, or -N(R₅)C(O)-,

and R₁ and R₂, together with the carbon atom to which they are attached, form a heterocyclic ring comprising one or two heteroatoms selected from oxygen, sulphur, -S(O)-, -S(O)₂-, -N=, -N(R₅)-, one or more carbon atoms in said heterocyclic ring being optionally substituted with one or more, same or different substituents selected from R₄;

R₃ is halogen, hydroxy, C₁₋₆-alkyl, C₂₋₆-alkenyl, C₂₋₆-alkynyl, halo-C₁₋₆-alkyl, C₁₋₆-alkoxy, halo-C₁₋₆-alkoxy, C₁₋₆-alkylthio, formyl, C₁₋₆-alkoxycarbonyl, C₁₋₆-alkylcarbonyl, or aminocarbonyl;

R₄ is hydrogen, amino, thioxo, C₁₋₆-alkyl, halo-C₁₋₆-alkyl, hydroxy-C₁₋₆-alkyl, C₁₋₆-alkoxy, halogen, oxo or hydroxy;

R₅ is hydrogen, C₁₋₆-alkyl, halo-C₁₋₆-alkyl, C₁₋₆-alkylcarbonyl, hydroxy-C₁₋₆-alkyl, C₁₋₆-alkoxycarbonyl, C₁₋₆-alkylsulfonyl, C₁₋₆-alkylaminosulfonyl or aminosulfonyl;

X is a bond, -CH₂-, or -NH-;

A is aryl comprising 6-10 carbon atoms, C₃₋₁₀-cycloalkyl, C₃₋₁₀-cycloalkenyl, aryl-C₁₋₆-alkyl, heteroaryl comprising 1-10 carbon atoms and 1-5 hetero atoms, heteroaryl-C₁₋₆-alkyl, the heteroaryl comprising 1-10 carbon atoms and 1-5 hetero atoms, hetero-C₃₋₁₀-cycloalkyl comprising 1-3 hetero atoms or hetero-C₃₋₁₀-cycloalkenyl comprising 1-3 hetero atoms, optionally substituted with one or more, same or different substituents selected from R₄;

and pharmaceutically acceptable salts, hydrates, N-oxides, or solvates thereof.

2. A compound according to claim 1, wherein E and G are both oxygen.

3. A compound according to claim 1 or 2, wherein m and n are both one.

4. A compound according to claim 1 or 2, wherein m and n are both zero.

5. A compound according to any one of the preceding claims, wherein R₁ and R₂, together with the carbon atom to which they are attached, form a 4-, 5- or 6-membered heterocyclic ring.

6. A compound according to claim 5, wherein the heterocyclic ring is tetrahydropyran, oxetane, [1,3]dioxolane, [1,3]di-

oxane, tetrahydrothiopyran, tetrahydrothiopyran-1,1-dioxide, tetrahydrothiopyran-1-oxide, piperidine, tetrahydrothiophene, [1,3]-dithiane, thietane, [1,3]-dithiane-1,3-dioxide, thietane-1-oxide, or thiethane-1,1-dioxide.

7. A compound according to claim 5, wherein the heterocyclic ring comprises one heteroatom.
8. A compound according to claim 5, wherein the heterocyclic ring comprises two heteroatoms.
9. A compound according to claim 7 or 8, wherein the heteroatom(s) is/are oxygen.
10. A compound according to claim 7 or 8, wherein the heteroatom(s) is/are sulphur, -S(O)-, or -S(O)₂-.
11. A compound according to any one of the preceding claims, wherein A is heteroaryl comprising 1-10 carbon atoms and 1-5 hetero atoms or heteroaryl-C₁₋₆-alkyl, the heteroaryl comprising 1-10 carbon atoms and 1-5 hetero atoms.
12. A compound according to claim 11, wherein A is pyridyl, pyrazinyl or quinolyl.
13. A compound according to any one of claims 1-10, wherein A is phenyl.
14. A compound according to any one of the preceding claims, wherein A is substituted with halogen.
15. A compound according to any one of the preceding claims, wherein R₃ is C₁₋₆alkoxy, C₁₋₆halogenalkyl, or halogen.
16. A compound according to claim 15, wherein R₃ is methoxy or ethoxy.
17. A compound according to any one of the preceding claims, wherein X is -CH₂-.
18. A compound according to any one of claims 1-16, wherein X is -NH-.
19. A compound according to claim 12, wherein A is 4-(3,5-dichloropyridyl).
20. A compound according to any one of the preceding claims, wherein R₄ is hydrogen.
21. A compound according to any one of the preceding claims, selected from the group consisting of 2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-yl)ethanone (compound 101),
N-(3,5-Dichloropyridine-4-yl)-7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-carboxamide (compound 102),
2-(3,5-Dichloro-1-oxido-pyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-yl)ethanone (compound 103),
2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-4',5'-dihydro-spiro[1,3-benzodioxole-2,3'-(2H)-thiophen]-4-yl)ethanone (compound 104),
2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 105),
2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-1'-[methoxycarbonyl]-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 106),
2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-1'-[methylsulfonyl]-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 107),
2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-1'-acetyl-spiro[1,3-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 108),
2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-1'-methyl-spiro[1,5-benzodioxole-2,4'-piperidine]-4-yl)ethanone (compound 109),
2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran]-4-yl)ethanone (compound 110),
2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran 1'-oxide]-4-yl)ethanone (compound 111),
2-(3,5-Dichloropyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4H)-thiopyran-1',1'-dioxide]-4-yl)ethanone (compound 112),
2-(3,5-Dichloro-1-oxido-pyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran-1',1'-dioxide]-4-yl)ethanone (compound 113),

2-(3-bromopyridine-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4*H*)-thiopyran]-4-yl)ethanone (compound 114),
 2-(3-Bromo-pyrazin-2-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4*H*)-thiopyran]-4-yl)ethanone (compound 115),
 2-(pyrazin-2-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4*H*)-thiopyran]-4-yl)ethanone (compound 116),
 2-(pyridin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4*H*)-thiopyran]-4-yl)ethanone (compound 117), 2-(quinolin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4*H*)-thiopyran]-4-yl)ethanone (compound 118),
 2-(2,6-Dichloro-phenyl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4*H*)-thiopyran]-4-yl)ethanone (compound 119),
 2-(2-Chloro-phenyl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxole-2, 4'-(4*H*)-thiopyran]-4-yl)ethanone (compound 120),
 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4*H*),3'-oxetane]-6-yl}ethanone (compound 121),
 2-(3,5-Dichloro-1-oxido-pyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4*H*),3'-oxetane]-6-yl}ethanone (compound 122),
 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4*H*),3'-thietane]-6-yl}ethanone (compound 123),
 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4*H*),3'-thietane-1',1'-dioxide]-6-yl}ethanone (compound 124),
 2-(3,5-Dichloropyridin-1-oxido-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4*H*),3'-thietane-1',1'-dioxide]-6-yl}ethanone (compound 125),
 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4*H*),2'-(1,3-dioxolane)]-6-yl}ethanone (compound 126),
 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4*H*),4'-tetrahydropyran]-6-yl}ethanone (compound 127),
 2-(3,5-Dichloro-1-oxido-pyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4*H*),4'-tetrahydropyran]-6-yl}ethanone (compound 128),
 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-2',2'-dimethyl-spiro[2H-1,5-benzodioxepin-3(4*H*),5'-[1,3]dioxane]-6-yl}ethanone (compound 129),
 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4*H*),5'-[1,3]dioxane]-6-yl}ethanone (compound 130), 2-(3,5-Dichloro-1-oxido-pyridine-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4*H*),5'-[1,3]dioxane]-6-yl}ethanone (compound 131), and
 2-(3,5-Dichloropyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4*H*),5'-[1,3]dithiane]-6-yl}ethanone (compound 132),
 and pharmaceutically acceptable salts, hydrates, N-oxides, or solvates thereof.

- 22.** A compound according to any one of claims 1-21 for use in therapy.
- 23.** A pharmaceutical composition comprising a compound according to any one of claims 1-21 together with a pharmaceutically acceptable excipient or vehicle or pharmaceutically acceptable carrier(s).
- 24.** A pharmaceutical composition according to claim 23 together with one or more other therapeutically active compound(s).
- 25.** A compound according to any one of claims 1-21 for use in the prevention, treatment or amelioration of dermal diseases or conditions, or acute or chronic cutaneous wound disorders.
- 26.** The compound according to claim 25, for use in the prevention, treatment or amelioration of a dermal disease or condition selected from the group consisting of proliferative and inflammatory skin disorders, psoriasis, cancer, epidermal inflammation, alopecia, skin atrophy, steroid induced skin atrophy, skin ageing, photo skin ageing, acne, dermatitis, atopic dermatitis, seborrheic dermatitis, contact dermatitis, urticaria, pruritis and eczema.

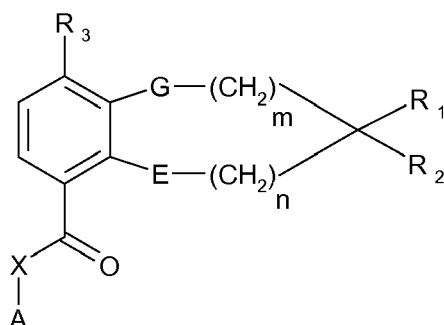
Patentansprüche

1. Verbindung der allgemeinen Formel I

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(I)

worin m und n unabhängig für 0, 1, 2, 3, 4, 5, 6 oder 7 stehen;

und worin G und E unabhängig für Schwefel, Sauerstoff, -N(R₅)- oder -N(R₅)C(O)-stehen,

und R₁ und R₂ gemeinsam mit dem Kohlenstoffatom, an welches sie gebunden sind, einen heterocyclischen Ring bilden, welcher ein oder zwei Heteroatome umfasst, die unter Sauerstoff, Schwefel, -S(O)-, -S(O)₂-, -N=, -N(R₅)-ausgewählt sind, wobei ein oder mehrere Kohlenstoffatome im heterocyclischen Ring gegebenenfalls mit einem oder mehreren identischen oder verschiedenen unter R₄ ausgewählten Substituenten substituiert sind;

R₃ ist Halogen, Hydroxy, C₁₋₆-Alkyl, C₂₋₆-Alkenyl, C₂₋₆-Alkinyl, Halogen-C₁₋₆-alkyl, C₁₋₆-Alkoxy, Halogen-C₁₋₆-alkoxy, C₁₋₆-Alkylthio, Formyl, C₁₋₆-Alkoxycarbonyl, C₁₋₆-Alkylcarbonyl oder Aminocarbonyl;

R₄ ist Wasserstoff, Amino, Thioxo, C₁₋₆-Alkyl, Halogen-C₁₋₆-alkyl, Hydroxy-C₁₋₆-alkyl, C₁₋₆-Alkoxy, Halogen, Oxo oder Hydroxy;

R₅ ist Wasserstoff, C₁₋₆-Alkyl, Halogen-C₁₋₆-alkyl, C₁₋₆-Alkylcarbonyl, Hydroxy-C₁₋₆-alkyl, C₁₋₆-Alkoxycarbonyl, C₁₋₆-Alkylsulfonyl, C₁₋₆-Alkylaminosulfonyl oder Aminosulfonyl;

X ist eine Bindung, -CH₂- oder -NH-;

A ist 6-10 Kohlenstoffatome umfassendes Aryl, C₃₋₁₀-Cycloalkyl, C₃₋₁₀-Cycloalkenyl, Aryl-C₁₋₆-alkyl, 1-10 Kohlenstoffatome und 1-5 Heteroatome umfassendes Heteroaryl, Heteroaryl-C₁₋₆-alkyl, wobei das Heteroaryl 1-10 Kohlenstoffatome und 1-5 Heteroatome umfasst, 1-3 Heteroatome umfassendes Hetero-C₃₋₁₀-cycloalkyl, oder 1-3 Heteroatome umfassendes Hetero-C₃₋₁₀-cycloalkenyl, die gegebenenfalls mit einem oder mehreren identischen oder verschiedenen unter R₄ ausgewählten Substituenten substituiert sind;

und pharmazeutisch akzeptable Salze, Hydrate, N-Oxide oder Solvate davon.

2. Verbindung gemäß Anspruch 1, worin E und G beide Sauerstoff sind.

3. Verbindung gemäß Anspruch 1 oder 2, worin m und n beide Eins sind.

4. Verbindung gemäß Anspruch 1 oder 2, worin m und n beide Null sind.

5. Verbindung gemäß einem der vorhergehenden Ansprüche, worin R₁ und R₂ gemeinsam mit dem Kohlenstoffatom, an welches sie gebunden sind, einen 4-, 5- oder 6-gliedrigen heterocyclischen Ring bilden.

6. Verbindung gemäß Anspruch 5, worin der heterocyclische Ring Tetrahydropyran, Oxetan, [1,3]-Dioxolan, [1,3]-Dioxan, Tetrahydrothiopyran, Tetrahydrothiopyran-1,1-dioxid, Tetrahydrothiopyran-1-oxid, Piperidin, Tetrahydrothiophen, [1,3]-Dithian, Thietan, [1,3]-Dithian-1,3-dioxid, Thietan-1-oxid oder Thiethan-1,1-dioxid ist.

7. Verbindung gemäß Anspruch 5, worin der heterocyclische Ring ein Heteroatom umfasst.

8. Verbindung gemäß Anspruch 5, worin der heterocyclische Ring zwei Heteroatome umfasst.

9. Verbindung gemäß Anspruch 7 oder 8, worin das/die Heteroatom(e) Sauerstoff ist/sind.

10. Verbindung gemäß Anspruch 7 oder 8, worin das/die Heteroatom(e) Schwefel, -S(O)- oder -S(O)₂- ist/sind.

11. Verbindung gemäß einem der vorhergehenden Ansprüche, worin A 1-10 Kohlenstoffatome und 1-5 Heteroatome umfassendes Heteroaryl, oder Heteroaryl-C₁₋₆-alkyl, wobei das Heteroaryl 1-10 Kohlenstoffatome und 1-5 Heteroatome umfasst, ist.
- 5 12. Verbindung gemäß Anspruch 11, worin A Pyridyl, Pyrazinyl oder Chinolyl ist.
13. Verbindung gemäß einem der Ansprüche 1-10, worin A Phenyl ist.
14. Verbindung gemäß einem der vorhergehenden Ansprüche, worin A mit Halogen substituiert ist.
- 10 15. Verbindung gemäß einem der vorhergehenden Ansprüche, worin R₃ C₁₋₆-Alkoxy, C₁₋₆-Halogenalkyl oder Halogen ist.
16. Verbindung gemäß Anspruch 15, worin R₃ Methoxy oder Ethoxy ist.
- 15 17. Verbindung gemäß einem der vorhergehenden Ansprüche, worin X -CH₂- ist.
18. Verbindung gemäß einem der Ansprüche 1-16, worin X -NH- ist.
- 20 19. Verbindung gemäß Anspruch 12, worin A 4-(3,5-Dichlorpyridyl) ist.
20. Verbindung gemäß einem der vorhergehenden Ansprüche, worin R₄ Wasserstoff ist.
21. Verbindung gemäß einem der vorhergehenden Ansprüche, ausgewählt unter
25 2-(3,5-Dichlorpyridin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2,4'-(4H)-pyran]-4-yl)ethanon (Verbindung 101),
N-(3,5-Dichlorpyridin-4-yl)-7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2,4'-(4H)-pyran]-4-carboxamid (Verbindung 102),
2-(3,5-Dichlor-1-oxido-pyridin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2,4'-(4H)-pyran]-4-yl)ethanon (Verbindung 103),
30 2-(3,5-Dichlorpyridin-4-yl)-1-(7-methoxy-4',5'-dihydro-spiro[1,3-benzodioxol-2,3'-(2H)-thiophen]-4-yl)ethanon (Verbindung 104),
2-(3,5-Dichlorpyridin-4-yl)-1-(7-methoxy-spiro[1,3-benzodioxol-2,4'-piperidin]-4-yl)ethanon (Verbindung 105),
2-(3,5-Dichlorpyridin-4-yl)-1-(7-methoxy-1'-[methoxycarbonyl]-spiro[1,3-benzodioxol-2,4'-piperidin]-4-yl)ethanon (Verbindung 106),
35 2-(3,5-Dichlorpyridin-4-yl)-1-(7-methoxy-1'-[methylsulfonyl]-spiro[1,3-benzodioxol-2,4'-piperidin]-4-yl)ethanon (Verbindung 107),
2-(3,5-Dichlorpyridin-4-yl)-1-(7-methoxy-1'-acetyl-spiro[1,3-benzodioxol-2,4'-piperidin]-4-yl)ethanon (Verbindung 108),
2-(3,5-Dichlorpyridin-4-yl)-1-(7-methoxy-1'-methyl-spiro[1,5-benzodioxol-2,4'-piperidin]-4-yl)ethanon (Verbindung 109),
40 2-(3,5-Dichlorpyridin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2, 4'-(4H)-thiopyran]-4-yl)ethanon (Verbindung 110),
2-(3,5-Dichlorpyridin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2, 4'-(4H)-thiopyran 1'-oxide]-4-yl)ethanon. (Verbindung 111),
45 2-(3,5-Dichlorpyridin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2, 4'-(4H)-thiopyran-1',1'-dioxide]-4-yl)ethanon (Verbindung 112),
2-(3,5-Dichlor-1-oxido-pyridin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2,4'-(4H)-thiopyran-1',1'-dioxide]-4-yl)ethanon (Verbindung 113),
2-(3-Brompyridin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2, 4'-(4H)-thiopyran]-4-yl)ethanon (Verbindung 114),
50 2-(3-Brom-pyrazin-2-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2, 4'-(4H)-thiopyran]-4-yl)ethanon (Verbindung 115),
2-(Pyrazin-2-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2, 4'-(4H)-thiopyran]-4-yl)ethanon (Verbindung 116),
55 2-(Pyridin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2, 4'-(4H)-thiopyran]-4-yl)ethanon (Verbindung 117),
2-(Chinolin-4-yl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2, 4'-(4H)-thiopyran]-4-yl)ethanon (Verbindung 118),

2-(2,6-Dichlor-phenyl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzo-dioxol-2, 4'-(4H)-thiopyran]-4-yl)ethanon (Verbindung 119),
 2-(2-Chlor-phenyl)-1-(7-methoxy-2',3',5',6'-tetrahydro-spiro[1,3-benzodioxol-2, 4'-(4H)-thiopyran]-4-yl)ethanon (Verbindung 120),
 2-(3,5-Dichlorpyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-oxetane]-6-yl}ethanon (Verbindung 121),
 2-(3,5-Dichlor-1-oxido-pyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-oxetan]-6-yl}ethanon (Verbindung 122),
 2-(3,5-Dichlorpyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietan]-6-yl}ethanon (Verbindung 123),
 2-(3,5-Dichlorpyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietan-1', 1'-dioxid]-6-yl}ethanon (Verbindung 124),
 2-(3,5-Dichlorpyridin-1-oxido-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),3'-thietan-1',1'-dioxid]-6-yl}ethanon (Verbindung 125),
 2-(3,5-Dichlorpyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),2'-(1,3-dioxolan)]-6-yl}ethanon (Verbindung 126),
 2-(3,5-Dichlorpyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),4'-tetrahydropyran]-6-yl}ethanon (Verbindung 127),
 2-(3,5-Dichlor-1-oxido-pyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),4'-tetrahydropyran]-6-yl}ethanon (Verbindung 128),
 2-(3,5-Dichlorpyridin-4-yl)-1-{9-methoxy-2',2'-dimethyl-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxane]-6-yl}ethanon (Verbindung 129),
 2-(3,5-Dichlorpyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxan]-6-yl}ethanon (Verbindung 130),
 2-(3,5-Dichlor-1-oxido-pyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dioxan]-6-yl}ethanon (Verbindung 131), und
 2-(3,5-Dichlorpyridin-4-yl)-1-{9-methoxy-spiro[2H-1,5-benzodioxepin-3(4H),5'-[1,3]dithian]-6-yl}ethanon (Verbindung 132),
 und pharmazeutisch akzeptable Salze, Hydrate, N-Oxide oder Solvate davon.

22. Verbindung gemäß einem der Ansprüche 1-21 zur Verwendung in der Therapie.

23. Pharmazeutische Zusammensetzung, umfassend eine Verbindung gemäß einem der Ansprüche 1-21 zusammen mit einem pharmazeutisch akzeptablen Exzipienten oder Vehikel oder mit einem oder mehreren pharmazeutisch akzeptablen Trägern.

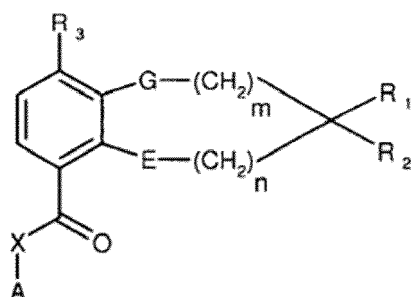
24. Pharmazeutische Zusammensetzung gemäß Anspruch 23 gemeinsam mit einer oder mehreren therapeutisch wirksamen Verbindungen.

25. Verbindung gemäß einem der Ansprüche 1-21 zur Verwendung in der Prävention, Behandlung oder Besserung von dermalen Erkrankungen oder Zuständen oder akuten oder chronischen kutanen Wund-Erkrankungen.

26. Verbindung nach Anspruch 25 zur Verwendung bei der Prävention, Behandlung oder Besserung einer dermalen Erkrankung oder eines dermalen Zustands, ausgewählt unter proliferativen und entzündlichen Hauterkrankungen, Psoriasis, Krebs, epidermaler Entzündung, Alopezie, Hautatrophie, steroidinduzierter Hautatrophie, Hautalterung, lichtinduzierter Hautalterung, Akne, Dermatitis, atopischer Dermatitis, seborrhöischer Dermatitis, Kontaktdermatitis, Urtikaria, Pruritis und Ekzemen.

Revendications

1. Composé de formule générale I



(I)

où m et n représentent indépendamment 0, 1, 2, 3, 4, 5, 6 ou 7 ;

et où G et E représentent indépendamment le soufre, l'oxygène, -N(R₅)- ou -N(R₅)C(O)-,

et R₁ et R₂, avec l'atome de carbone auquel ils sont liés, forment un cycle hétérocyclique comprenant un ou deux hétéroatomes choisis parmi l'oxygène, le soufre, -S(O)-, -S(O)₂-, -N=, -N(R₅)-, un ou plusieurs atomes de carbone dans ledit cycle hétérocyclique étant éventuellement substitués avec un ou plusieurs substituants identiques ou différents choisis parmi R₄ ;

R₃ est halogène, hydroxy, C₁₋₆-alkyle, C₂₋₆-alcényle, C₂₋₆-alcynyle, halo-C₁₋₆-alkyle, C₁₋₆-alcoxy, halo-C₁₋₆-alcoxy, C₁₋₆-alkylthio, formyle, C₁₋₆-alcoxy-carbonyl, C₁₋₆-alkylcarbonyl, ou aminocarbonyl ;

R₄ est l'hydrogène, amino, thioxo, C₁₋₆-alkyle, halo-C₁₋₆-alkyle, hydroxy-C₁₋₆-alkyle, C₁₋₆-alcoxy, halogène, oxo ou hydroxy ;

R₅ est l'hydrogène, C₁₋₆-alkyle, halo-C₁₋₆-alkyle, C₁₋₆-alkylcarbonyl, hydroxy-C₁₋₆-alkyle, C₁₋₆-alcoxycarbonyl, C₁₋₆-alkylsulfonyl, C₁₋₆-alkylaminosulfonyl ou aminosulfonyl ;

X est une liaison, -CH₂- ou -NH- ;

A est aryle comprenant 6-10 atomes de carbone, C₃₋₁₀-cycloalkyle, C₃₋₁₀-cycloalcényle, aryl-C₁₋₆-alkyle, hétéroaryle comprenant 1-10 atomes de carbone et 1-5 hétéroatomes, hétéroaryl-C₁₋₆-alkyle, l'hétéroaryle comprenant 1-10 atomes de carbone et 1-5 hétéroatomes, hétéro-C₃₋₁₀-cycloalkyle comprenant 1-3 hétéroatomes ou hétéro-C₃₋₁₀-cycloalcényle comprenant 1-3 hétéroatomes, éventuellement substitué avec un ou plusieurs substituants identiques ou différents choisis parmi R₄ ;

et ses sels, hydrates, N-oxydes ou solvates pharmaceutiquement acceptables.

2. Composé selon la revendication 1, où E et G sont l'un et l'autre l'oxygène.

3. Composé selon la revendication 1 ou 2, où m et n sont l'un et l'autre un.

4. Composé selon la revendication 1 ou 2, où m et n sont l'un et l'autre zéro.

5. Composé selon l'une quelconque des revendications précédentes, où R₁ et R₂, avec l'atome de carbone auquel ils sont liés, forment un cycle hétérocyclique à 4, 5 ou 6 chaînons.

6. Composé selon la revendication 5, où le cycle hétérocyclique est tétrahydropyrane, oxétane, [1,3]dioxolane, [1,3]dioxane, tétrahydrothiopyrane, tétrahydrothiopyrane-1,1-dioxyde, tétrahydrothiopyrane-1-oxyde, pipéridine, tétrahydrothiophène, [1,3]-dithiane, thiétane, [1,3]-dithiane-1,3-dioxyde, thiétane-1-oxyde, ou thiéthane-1,1-dioxyde.

7. Composé selon la revendication 5, où le cycle hétérocyclique comprend un hétéroatome.

8. Composé selon la revendication 5, où le cycle hétérocyclique comprend deux hétéroatomes.

9. Composé selon la revendication 7 ou 8, où le ou les hétéroatomes est/sont l'oxygène.

10. Composé selon la revendication 7 ou 8, où le ou les hétéroatomes est/sont le soufre, -S(O)- ou -S(O)₂-.

11. Composé selon l'une quelconque des revendications précédentes, où A est hétéroaryle comprenant 1-10 atomes de carbone et 1-5 hétéroatomes ou hétéroaryl-C₁₋₆-alkyle, l'hétéroaryle comprenant 1-10 atomes de carbone et 1-5 hétéroatomes.

12. Composé selon la revendication 11, où A est pyridyle, pyrazinyle ou quinolyle.

13. Composé selon l'une quelconque des revendications 1-10, où A est phényle.

14. Composé selon l'une quelconque des revendications précédentes, où A est substitué avec halogène.

5 15. Composé selon l'une quelconque des revendications précédentes, où R₃ est C₁₋₆alcoxy, C₁₋₆halogénoalkyle ou halogène.

16. Composé selon la revendication 15, où R₃ est méthoxy ou éthoxy.

10 17. Composé selon l'une quelconque des revendications précédentes, où X est -CH₂-.

18. Composé selon l'une quelconque des revendications 1-16, où X est -NH-.

19. Composé selon la revendication 12, où A est 4-(3,5-dichloropyridyle).

15 20. Composé selon l'une quelconque des revendications précédentes, où R₄ est l'hydrogène.

21. Composé selon l'une quelconque des revendications précédentes, choisi dans le groupe consistant en la 2-(3,5-dichloropyridin-4-yl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyran]-4-yl)éthanone (composé 101),

le N-(3,5-dichloropyridin-4-yl)-7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyrane]-4-carboxamide (composé 102),

la 2-(3,5-dichloro-1-oxydo-pyridin-4-yl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-pyrane]-4-yl)éthanone (composé 103),

25 la 2-(3,5-dichloropyridin-4-yl)-1-(7-méthoxy-4',5'-dihydro-spiro[1,3-benzodioxole-2,3'-(2H)-thiophène]-4-yl)éthanone (composé 104),

la 2-(3,5-dichloropyridin-4-yl)-1-(7-méthoxy-spiro[1,3-benzodioxole-2,4'-pipéridine]-4-yl)éthanone (composé 105),

la 2-(3,5-dichloropyridin-4-yl)-1-(7-méthoxy-1'-[méthoxycarbonyl]-spiro[1,3-benzodioxole-2,4'-pipéridine]-4-yl)éthanone (composé 106),

30 la 2-(3,5-dichloropyridin-4-yl)-1-(7-méthoxy-1'-[méthylsulfonyl]-spiro[1,3-benzodioxole-2,4'-pipéridine]-4-yl)éthanone (composé 107),

la 2-(3,5-dichloropyridin-4-yl)-1-(7-méthoxy-1'-acétyl-spiro[1,3-benzodioxole-2,4'-pipéridine]-4-yl)éthanone (composé 108),

35 la 2-(3,5-dichloropyridin-4-yl)-1-(7-méthoxy-1'-méthyl-spiro[1,5-benzodioxole-2,4'-pipéridine]-4-yl)éthanone (composé 109),

la 2-(3,5-dichloropyridin-4-yl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyran]-4-yl)éthanone (composé 110),

la 2-(3,5-dichloropyridin-4-yl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyrane-1'-oxyde]-4-yl)éthanone (composé 111),

40 la 2-(3,5-dichloropyridin-4-yl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyrane-1',1'-dioxide]-4-yl)éthanone (composé 112),

la 2-(3,5-dichloro-1-oxydo-pyridin-4-yl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyrane-1',1'-dioxide]-4-yl)éthanone (composé 113),

la 2-(3-bromopyridin-4-yl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyrane]-4-yl)éthanone (composé 114),

45 la 2-(3-bromo-pyrazin-2-yl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyrane]-4-yl)éthanone (composé 115),

la 2-(pyrazin-2-yl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyrane]-4-yl)éthanone (composé 116),

50 la 2-(pyridin-4-yl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyrane]-4-yl)éthanone (composé 117),

la 2-(quinolin-4-yl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyrane]-4-yl)éthanone (composé 118),

la 2-(2,6-dichloro-phényl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyrane]-4-yl)éthanone (composé 119),

55 la 2-(2-chloro-phényl)-1-(7-méthoxy-2',3',5',6'-tétrahydro-spiro[1,3-benzodioxole-2,4'-(4H)-thiopyrane]-4-yl)éthanone (composé 120),

la 2-(3,5-dichloropyridin-4-yl)-1-{9-méthoxy-spiro[2H-1,5-benzodioxépin-3(4H),3'-oxétane]-6-yl}éthanone (compo-

sé 121),

la 2-(3,5-dichloro-1-oxydo-pyridin-4-yl)-1-{9-méthoxy-spiro[2H-1,5-benzodioxépin-3(4H),3'-oxétane]-6-yl}éthanone (composé 122),

la 2-(3,5-dichloropyridin-4-yl)-1-{9-méthoxy-spiro[2H-1,5-benzodioxépin-3(4H),3'-thiétane]-6-yl}éthanone (composé 123),

la 2-(3,5-dichloropyridin-4-yl)-1-{9-méthoxy-spiro[2H-1,5-benzodioxépin-3(4H),3'-thiétane-1',1'-dioxyde]-6-yl}éthanone (composé 124),

la 2-(3,5-dichloropyridin-4-yl)-1-{9-méthoxy-spiro[2H-1,5-benzodioxépin-3(4H),3'-thiétane-1',1'-dioxyde]-6-yl}éthanone (composé 125),

la 2-(3,5-dichloropyridin-4-yl)-1-{9-méthoxy-spiro[2H-1,5-benzodioxépin-3(4H),2'-(1,3-dioxolane)]-6-yl}éthanone (composé 126),

la 2-(3,5-dichloropyridin-4-yl)-1-{9-méthoxy-spiro[2H-1,5-benzodioxépin-3(4H),4'-tétrahydropyran]-6-yl}éthanone (composé 127),

la 2-(3,5-dichloro-1-oxydo-pyridin-4-yl)-1-{9-méthoxy-spiro[2H-1,5-benzodioxépin-3(4H),4'-tétrahydropyran]-6-yl}éthanone (composé 128),

la 2-(3,5-dichloropyridin-4-yl)-1-{9-méthoxy-2',2'-diméthyl-spiro[2H-1,5-benzodioxépin-3(4H),5'-[1,3]dioxane]-6-yl}éthanone (composé 129),

la 2-(3,5-dichloropyridin-4-yl)-1-{9-méthoxy-spiro[2H-1,5-benzodioxépin-3(4H),5'-[1,3]dioxane]-6-yl}éthanone (composé 130),

la 2-(3,5-dichloro-1-oxydo-pyridin-4-yl)-1-{9-méthoxy-spiro[2H-1,5-benzodioxépin-3(4H),5'-[1,3]dioxane]-6-yl}éthanone (composé 131), et

la 2-(3,5-dichloropyridin-4-yl)-1-{9-méthoxy-spiro[2H-1,5-benzodioxépin-3(4H),5'-[1,3]dithiane]-6-yl}éthanone (composé 132),

et ses sels, hydrates, N-oxydes ou solvates pharmaceutiquement acceptables.

22. Composé selon l'une quelconque des revendications 1-21 destiné à être utilisé en thérapie.

23. Composition pharmaceutique comprenant un composé selon l'une quelconque des revendications 1-21 avec un excipient ou véhicule pharmaceutiquement acceptable ou un ou des vecteurs pharmaceutiquement acceptables.

24. Composition pharmaceutique selon la revendication 23 avec un ou plusieurs autres composés thérapeutiquement actifs.

25. Composé selon l'une quelconque des revendications 1-21 destiné à être utilisé dans la prévention, le traitement ou l'amélioration des maladies ou affections dermatiques ou des troubles de type plaie cutanée aigus ou chroniques.

26. Composé selon la revendication 25, destiné à être utilisé dans la prévention, le traitement ou l'amélioration d'une maladie ou affection dermatique choisie dans le groupe consistant en les troubles cutanées prolifératifs et inflammatoires, le psoriasis, le cancer, l'inflammation épidermique, l'alopécie, l'atrophie cutanée, l'atrophie cutanée induite par des stéroïdes, le vieillissement cutané, le photovieillissement cutané, l'acné, la dermatite, la dermatite atopique, la dermatite séborrhéique, la dermatite de contact, l'urticaire, le prurit et l'eczéma.

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