

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property
Organization
International Bureau



(43) International Publication Date
20 October 2005 (20.10.2005)

PCT

(10) International Publication Number
WO 2005/097763 A3

(51) International Patent Classification⁷: C07D 417/12,
413/12, A61K 31/4245, A61P 3/10, 3/06

(21) International Application Number:
PCT/US2005/010855

(22) International Filing Date: 30 March 2005 (30.03.2005)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:
60/558,420 1 April 2004 (01.04.2004) US

(71) Applicant (for all designated States except US): AVEN-
TIS PHARMACEUTICALS INC. [US/US]; 300 Somer-
set Corporate Boulevard, Bridgewater, New Jersey 08807-
2854 (US).

(72) Inventors; and

(75) Inventors/Applicants (for US only): MCGARRY,
Daniel, G. [GB/US]; 205 Carol Jean Way, Branch-
burg, New Jersey 08876 (US). GOERLITZER, Jochen
[DE/DE]; Stegstrasse 60, D-60594 Frankfurt am Main
(DE). KEIL, Stefanie [DE/DE]; Am Kreishaus 12, 65719
Hofheim (DE). CHANDROSS, Karen [US/US]; 13
Staudt Court, Somerset, NJ 08873 (US). MERRILL, Jean
[US/US]; 101 Alpine Lane, Whippany, New Jersey 07981
(US). WENDLER, Wolfgang [DE/DE]; Haintchener Str.
12a, 65618 Selters (DE).

(74) Agents: STRUPCZEWSKI, Joseph et al.; sanofi-aventis,
Route 202-206, P. O. Box 6800, Bridgewater, NJ 08807-
0800 (US).

(81) Designated States (unless otherwise indicated, for every
kind of national protection available): AE, AG, AL, AM,
AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CII, CN,
CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI,
GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE,
KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD,
MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG,
PH, PI, PT, RO, RU, SC, SD, SE, SG, SK, SI, SM, SY, TJ,
TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA,
ZM, ZW.

(84) Designated States (unless otherwise indicated, for every
kind of regional protection available): ARIPO (BW, GH,
GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM,
ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM),
European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI,
FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO,
SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN,
GQ, GW, ML, MR, NE, SN, TD, TG).

Declaration under Rule 4.17:

— of inventorship (Rule 4.17(iv)) for US only

Published:

— with international search report

(88) Date of publication of the international search report:
15 December 2005

For two-letter codes and other abbreviations, refer to the "Guid-
ance Notes on Codes and Abbreviations" appearing at the begin-
ning of each regular issue of the PCT Gazette.

(54) Title: 1,3,4-OXADIAZOL-2-ONES AS PPAR DELTA

(57) Abstract: The present invention is directed to 1, 3, 4-oxadiazolones, and their pharmaceutically acceptable salts stereoisomers, tautomers, or solvates thereof. The compounds of this invention are modulators of PPARdelta and therefore useful as pharmaceutical agents, especially for the treatment of demyelinating diseases and disorders of fatty acid metabolism and glucose utilization.

WO 2005/097763 A3

5

10 **1, 3, 4-OXADIAZOL-2-ONES AS PPAR DELTA MODULATORS AND THEIR USE
THEREOF**

FIELD OF THE INVENTION

This invention relates to novel compounds and pharmaceutical formulations that act as
15 selective PPARdelta ligand receptor binders, which are useful in modulating PPARdelta
receptors for the treatment of diseases mediated by nuclear hormone receptors. The
PPARdelta ligand receptor binders of this invention are useful as agonists or antagonists of the
PPARdelta receptor.

20 **BACKGROUND OF THE INVENTION**

The peroxisome proliferator-activated receptors (PPARs) comprise a subfamily of the nuclear receptor superfamily. Four closely related isoforms have been identified and cloned and are commonly known as PPARalpha, PPARgamma-1, PPARgamma-2 and PPARdelta. Each receptor subtype has a signature DNA binding domain (DBD) and a ligand-binding domain (LBD), both being necessary for ligand activated gene expression. PPARs bind as heterodimers with a retinoid X receptor. See J. Berger and D. E. Miller, *Annu. Rev. Med.*, 2002, 53, 409-435.

PPARdelta (also known as PPARbeta) is expressed in a broad range of mammalian tissue, but little information regarding its biological functions or the full array of genes regulated by the receptor have been elucidated. However, it has recently been found that agonists may be useful to treat conditions such as dyslipedemia and certain dermatological conditions, while antagonists may be useful to treat osteoporosis or colorectal cancer (D.

Sternbach, in *Annual Reports in Medicinal Chemistry, Volume 38*, A. M. Doherty, ed., Elsevier Academic Press, 2003 pp. 71-80).

PPARdelta appears to be significantly expressed in the CNS; however much of its function there still remains undiscovered. Of singular interest however, is the discovery that 5 PPARdelta was expressed in rodent oligodendrocytes, the major lipid producing cells of the CNS (J. Granneman, et al., *J. Neurosci. Res.*, 1998, 51, 563-573). Moreover, it was also found that a PPARdelta selective agonist was found to significantly increase oligodendroglial myelin gene expression and myelin sheath diameter in mouse cultures (I. Saluja et al., *Glia*, 2001, 33, 194-204). Thus, PPARdelta activators may be of use for the treatment of 10 demyelinating and dysmyelinating diseases.

Demyelinating conditions are manifested in loss of myelin- the multiple dense layers of lipids and protein which cover many nerve fibers. These layers are provided by oligodendroglia in the central nervous system (CNS), and Schwann cells in the peripheral nervous system (PNS). In patients with demyelinating conditions, demyelination may be 15 irreversible; it is usually accompanied or followed by axonal degeneration, and often by cellular degeneration. Demyelination can occur as a result of neuronal damage or damage to the myelin itself--whether due to aberrant immune responses, local injury, ischemia, metabolic disorders, toxic agents, or viral infections (Prineas and McDonald, *Demyelinating Diseases. In Greenfield's Neuropathology*, 6.sup.th ed. (Edward Arnold: New York, 1997) 813-811, Beers 20 and Berkow, eds., *The Merck Manual of Diagnosis and Therapy*, 17.sup.th ed. (Whitehouse Station, N.J.: Merck Research Laboratories, 1999) 1299, 1437, 1473-76, 1483).

Central demyelination (demyelination of the CNS) occurs in several conditions, often of uncertain etiology, that have come to be known as the primary demyelinating diseases. Of these, multiple sclerosis (MS) is the most prevalent. Other primary demyelinating diseases 25 include adrenoleukodystrophy (ALD), adrenomyeloneuropathy, AIDS-vacuolar myopathy, HTLV-associated myopathy, Leber's hereditary optic atrophy, progressive multifocal leukoencephalopathy (PML), subacute sclerosing panencephalitis, Guillain-Barre syndrome and tropical spastic paraparesis. In addition, there are acute conditions in which demyelination can occur in the CNS, e.g., acute disseminated encephalomyelitis (ADEM) and acute viral 30 encephalitis. Furthermore, acute transverse myelitis, a syndrome in which an acute spinal cord transection of unknown cause affects both gray and white matter in one or more adjacent thoracic segments, can also result in demyelination. Also, disorders in which myelin forming glial cells are damaged including spinal cord injuries, neuropathies and nerve injury.

Selective PPARdelta modulators may also be useful for treating or preventing other disease states see, for example, Joel Berger et al., Annu. Rev. Med. 2002, 53, 409 – 435; Timothy Wilson et al. J. Med. Chem., 2000, Vol. 43, No. 4, 527-550; Steven Kliewer et al., Recent Prog Horm Res. 2001; 56: 239-63; Jean-Charles Fruchart, Bart Staels and Patrick 5 Duriez: PPARS, Metabolic Disease and Arteriosclerosis, Pharmacological Research, Vol. 44, No. 5, 345-52; 2001; Sander Kersten, Beatrice Desvergne & Walter Wahli: Roles of PPARs in health and disease, Nature, vol 405, 25 may 2000; 421-4; Ines Pineda Torra, Giulia Chinetti, Caroline Duval, Jean-Charles Fruchart and Bart Staels: Peroxisome proliferator-activated receptors: from transcriptional control to clinical practice, Curr Opin Lipidol 12: 2001, 245-10 254).

Compounds acting as PPARdelta modulators may be particularly suitable for the treatment and/or prevention of disorders of fatty acid metabolism and glucose utilization disorders in which insulin resistance is involved.

15 Diabetes mellitus, especially type 2 diabetes, including the prevention of the sequelae associated therewith. Particular aspects in this connection are hyperglycemia, improvement in insulin resistance, improvement in glucose tolerance, protection of the pancreatic β cells, prevention of macro- and microvascular disorders.

Dyslipidemias and their sequelae such as, for example, atherosclerosis, coronary heart disease, cerebrovascular disorders etc, especially those (but not 20 restricted thereto) which are characterized by one or more of the following factors: high plasma triglyceride concentrations, high postprandial plasma triglyceride concentrations, low HDL cholesterol concentrations, low ApoA lipoprotein concentrations, high LDL cholesterol concentrations, small dense LDL cholesterol particles, high ApoB lipoprotein concentrations.

25 Various other conditions which may be associated with the metabolic syndrome, such as: obesity (excess weight), including central obesity, thromboses, hypercoagulable and prothrombotic states (arterial and venous), high blood pressure, heart failure such as, for example (but not restricted thereto), following myocardial infarction, hypertensive heart disease or cardiomyopathy.

30 Other disorders or conditions in which inflammatory reactions or cell differentiation, may for example be involved are: atherosclerosis such as, for example (but not restricted thereto), coronary sclerosis including angina pectoris or myocardial infarction, stroke, vascular restenosis or reocclusion, chronic inflammatory bowel diseases, such as, for example,

Crohn's disease and ulcerative colitis, pancreatitis, other inflammatory states, retinopathy, adipose cell tumors, lipomatous carcinomas such as, for example, liposarcomas, solid tumors and neoplasms such as, for example (but not restricted thereto), carcinomas of the gastrointestinal tract, of the liver, of the biliary tract and of the pancreas, endocrine tumors, 5 carcinomas of the lungs, of the kidneys and the urinary tract, of the genital tract, prostate carcinomas etc., acute and chronic myeloproliferative disorders and lymphomas angiogenesis, neurodegenerative disorders, Alzheimer's disease, Parkinson's disease, erythematous squamous dermatoses such as, for example, psoriasis, acne vulgaris.

Other skin disorders and dermatological conditions modulated by PPARdelta:

10 eczemas and neurodermitis, dermatitis such as, for example, seborrheic dermatitis or photodermatitis, keratitis and keratoses such as, for example, seborrheic keratoses, senile keratoses, actinic keratoses, photo-induced keratoses or keratosis follicularis keloids and keloid prophylaxis, warts, including condylomata or condylomata acuminata, human papilloma viral (HPV) infections such as, for example, venereal papillomata, viral warts such 15 as, for example, molluscum contagiosum, leukoplakia papular dermatoses such as, for example, Lichen planus, skin cancer such as, for example, basal-cell carcinomas, melanomas or cutaneous T-cell lymphomas, localized benign epidermal tumors such as, for example, keratoderma, epidermal naevi and chilblains.

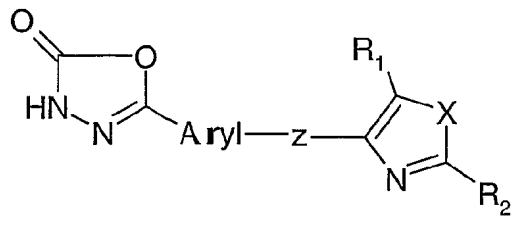
20 Various other conditions potentially modulated by PPARdelta including syndrome X, polycystic ovary syndrome (PCOS), asthma osteoarthritis, lupus erythematosus (LE) or inflammatory rheumatic disorders such as, for example, rheumatoid arthritis, vasculitis, wasting (cachexia), gout ischemia/reperfusion syndrome and acute respiratory distress syndrome (ARDS).

25

SUMMARY OF THE INVENTION

The present invention is directed to compound of formula I.

30



wherein

ARYL is phenyl or pyridinyl, wherein said phenyl or pyridinyl is optionally substituted with one or more substituents selected from the group consisting of halogen, C₁₋₆alkyl, C₂₋₆alkenyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl; C₁₋₆alkylthio, hydroxy, hydroxyC₁₋₆alkyl, C₁₋₄acyloxy, nitro, cyano, C₁₋₆alkylsulfonyl, amino, C₁₋₆alkylamino and C₁₋₆alkoxycarbonyl;

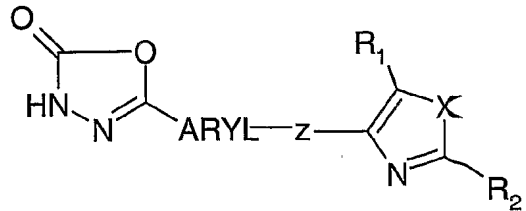
5 Z is -O(CH₂)_n-, -SO₂(CH₂)_n-, -(CH₂)_n-Y-(CH₂)_n-, -(CH₂)_n-CO-, -O(CH₂)_n-CO- or -(CH₂)_n-Y-(CH₂)_n-CO- wherein Y is NR₃, O or S and R₃ is selected from the group consisting of H, C₁₋₆alkyl C₃₋₈cycloalkyl, C₁₋₆alkylC₃₋₈cycloalkyl and benzyl and n is independently an integer from 1 to 5;

10 X is NR₃, O or S wherein R₃ is as defined above;
R₁ is H, halogen, C₁₋₆alkyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl; hydroxyC₁₋₆alkyl, nitro, cyano, and C₁₋₆alkylamino; and
R₂ is substituted or unsubstituted phenyl, pyridinyl or thiienyl wherein the substituents are selected from the group consisting of halogen, C₁₋₆alkyl, C₂₋₆alkenyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl, C₁₋₆alkylthio, hydroxy, hydroxyC₁₋₆alkyl, C₁₋₄acyloxy, nitro, cyano, C₁₋₆alkylsulfonyl, amino, C₁₋₆alkylamino and C₁₋₆alkoxycarbonyl;

15 with the proviso that when Z is -O(CH₂)_n or -SO₂(CH₂)_n and ARYL is phenyl then R₂ is other than phenyl;
20 or a stereoisomer, a tautomer or a solvate thereof or a pharmaceutically acceptable salt thereof.

The present invention is also directed to pharmaceutical compositions of formula I, and methods of using said compounds and compositions for modulating PPARdelta in a subject in need of such modulation by administering a compound which preferentially modulates the activity of PPARdelta.

25 Another aspect of this invention is disclosed a method of treating a disease in a mammal wherein the disease is capable of being, modulated by PPARdelta ligand binding activity, which comprises administering to said mammal having said disease a therapeutically effective amount of a compound of formula I.



I

wherein

ARYL is phenyl or pyridinyl, wherein said phenyl or pyridinyl is optionally substituted with one or more substituents selected from the group consisting of halogen, C₁₋₆alkyl, C₂₋₆alkenyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl; C₁₋₆alkylthio, hydroxy, hydroxyC₁₋₆alkyl, C₁₋₄acyloxy, nitro, cyano, C₁₋₆alkylsulfonyl, amino, C₁₋₆alkylamino and C₁₋₆alkoxycarbonyl;

10 Z is -O(CH₂)_n-, -SO₂(CH₂)_n-, -(CH₂)_n-Y-(CH₂)_n-, -(CH₂)_n-CO-, -O(CH₂)_n-CO-
or -(CH₂)_n-Y-(CH₂)_n-CO- wherein Y is NR₃, O or S and R₃ is selected from the group
consisting of H, C₁₋₆alkyl C₃₋₈cycloalkyl, C₁₋₆alkylC₃₋₈cycloalkyl and benzyl and n is
independently an integer from 1 to 5;

X is NR₃, O or S wherein R₃ is as defined above;

15 R₁ is H, halogen, C₁₋₆alkyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl; hydroxyC₁₋₆alkyl, nitro, cyano,
and C₁₋₆alkylamino; and

R₂ is substituted or unsubstituted phenyl, pyridinyl or thiophenyl wherein the
substituents are selected from the group consisting of halogen, C₁₋₆alkyl, C₂₋₆alkenyl,
C₁₋₆alkoxy, C₁₋₆perfluoroalkyl, C₁₋₆alkylthio, hydroxy, hydroxyC₁₋₆alkyl, C₁₋₄acyloxy,
nitro, cyano, C₁₋₆alkylsulfonyl, amino, C₁₋₆alkylamino and C₁₋₆alkoxycarbonyl; or a
stereoisomer, a tautomer or a solvate thereof or a pharmaceutically acceptable salt
thereof.

DETAILED DESCRIPTION OF THE INVENTION

The terms as used herein have the following meanings:

25 As used herein, the expression "C₁₋₆ alkyl" includes methyl and ethyl groups, and straight-chained or branched propyl, butyl, pentyl and hexyl groups. Particular alkyl groups are methyl, ethyl, n-propyl, isopropyl and tert-butyl. Derived expressions such as "C₁₋₆alkoxy", "C₁₋₆alkoxyC₁₋₆alkyl", "hydroxyC₁₋₆alkyl", "C₁₋₆alkylcarbonyl", "C₁₋₆alkoxycarbonylC₁₋₆alkyl", "C₁₋₆alkoxycarbonyl", "aminoC₁₋₆alkyl", "C₁₋₆alkylcarbamoylC₁₋₆alkyl", "C₁₋₆dialkylcarbamoylC₁₋₆alkyl" "mono- or di-C₁₋₆alkylaminoC₁₋₆alkyl",
30

aminoC₁₋₆alkylcarbonyl", "diphenylC₁₋₆alkyl", "arylC₁₋₆alkyl", "arylcarbonylC₁₋₆alkyl" and "aryloxyC₁₋₆alkyl" are to be construed accordingly.

As used herein, the expression "C₂₋₆alkenyl" includes ethenyl and straight-chained or branched propenyl, butenyl, pentenyl and hexenyl groups. Similarly, the expression "C₂₋₆alkynyl" includes ethynyl and propynyl, and straight-chained or branched butynyl, pentynyl and hexynyl groups.

As used herein, the term "C₁₋₄acyloxy" denotes an acyl radical attached to an oxygen atom, some examples include but not limited to acyloxy, propionyloxy, butanoyloxy, iso-butanoyloxy, sec-butanoyloxy, t-butanoyloxy and the like.

10 As used herein "aryl" represents a carbocyclic aromatic ring system such as phenyl, biphenyl, naphthyl, anthracenyl, phenanthrenyl, fluorenyl, indenyl, pentalenyl, azulenyl, biphenylenyl and the like. Aryl is also intended to include the partially hydrogenated derivatives of the carbocyclic aromatic systems enumerated above. Non-limiting examples of such partially hydrogenated derivatives are 1,2,3,4-tetrahydronaphthyl, 1,4-dihydronaphthyl
15 and the like.

As used herein "aryloxy" represents a group --O-aryl wherein aryl is as defined above.

As used herein "heteroaryl" (on its own or in any combination, such as "heteroaryloxy", or "heteroaryl alkyl")--a 5-10 membered aromatic ring system in which one or more rings contain one or more heteroatoms selected from the group consisting of N, O or 20 S, such as, but not limited, to pyrrole, pyrazole, furan, thiophene, quinoline, isoquinoline, quinazolinyl, pyridine, pyrimidine, oxazole, thiazole, thiadiazole, tetrazole, triazole, imidazole, or benzimidazole.

As used herein "heterocyclic or heterocyclyl" (on its own or in any combination, such as "heterocyclalkyl")--a saturated or partially unsaturated 4-10 membered ring system in which one or more rings contain one or more heteroatoms selected from the group consisting of N, O, or S; such as, but not limited to, pyrrolidine, piperidine, piperazine, morpholine, tetrahydro pyran, or imidazolidine.

As used herein, the expression "C₁₋₆ perfluoroalkyl" means that all of the hydrogen atoms in said alkyl group are replaced with fluorine atoms. Illustrative examples include 30 trifluoromethyl and pentafluoroethyl, and straight-chained or branched heptafluoropropyl, nonafluorobutyl, undecafluoropentyl and tridecafluorohexyl groups. Derived expression, "C₁₋₆ perfluoroalkoxy", is to be construed accordingly.

As used herein, the expression "C₃₋₈cycloalkyl" means cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cycloheptyl and cyclooctyl.

As used herein, the expression "C₃₋₈cycloalkylC₁₋₆alkyl" means that the C₃₋₈cycloalkyl as defined herein is further attached to C₁₋₆alkyl as defined herein. Representative examples 5 include cyclopropylmethyl, 1-cyclobutylethyl, 2-cyclopentylpropyl, cyclohexylmethyl, 2-cycloheptylethyl and 2-cyclooctylbutyl and the like.

As used herein "halogen" or "halo" means chloro, fluoro, bromo, and iodo.

As used herein "C₁₋₆alkylsulfonyl" in the present context designates a group --S(=O)₂C₁₋₆alkyl wherein C₁₋₆alkyl is as defined above. Representative examples include, but 10 are not limited to, methylsulfonyl, ethylsulfonyl, n-propylsulfonyl, isopropylsulfonyl, butylsulfonyl, iso-butylsulfonyl, sec-butylsulfonyl, tert-butylsulfonyl, n-pentylsulfonyl, isopentylsulfonyl, neopentylsulfonyl, tert-pentylsulfonyl, n-hexylsulfonyl, isoheptylsulfonyl and the like.

As used herein "arylsulfonyl" represents a group --S(=O)₂aryl wherein aryl is as 15 defined above.

As used herein "heteroarylsulfonyl" represents a group --S(=O)₂heteroaryl wherein heteroaryl is as defined above.

The expression "stereoisomers" is a general term used for all isomers of the individual molecules that differ only in the orientation of their atoms in space. Typically it includes 20 mirror image isomers that are usually formed due to at least one asymmetric center, (enantiomers). Where the compounds according to the invention possess two or more asymmetric centers, they may additionally exist as diastereoisomers, also certain individual molecules may exist as geometric isomers (cis/trans). It is to be understood that all such isomers and mixtures thereof in any proportion are encompassed within the scope of the 25 present invention.

"Substituted" means substituted by 1 to 2 substituents independently selected from the group consisting of C₁₋₆ alkyl, C₁₋₆ perfluoroalkyl, hydroxy, -CO₂H, an ester, an amide, C₁ -C₆ alkoxy, C₁ -C₆ perfluoroalkoxy, -NH₂, Cl, Br, I, F, -NH-lower alkyl, and -N(lower alkyl)₂.

The compounds and salts of the present invention may exist in several tautomeric 30 forms, including the enol and imine form, and the keto and enamine form and geometric isomers and mixtures thereof. All such tautomeric forms are included within the scope of the present invention. Tautomers exist as mixtures of a tautomeric set in solution. In solid form,

usually one tautomer predominates. Even though one tautomer may be described, the present invention includes all tautomers of the present compounds.

As used herein the term "modulator" refers to a chemical compound with capacity to either enhance (e.g., "agonist" activity) or inhibit (e.g., "antagonist" activity) a functional property of biological activity or process (e.g., enzyme activity or receptor binding); such enhancement or inhibition may be contingent on the occurrence of a specific event, such as activation or repression of a signal transduction pathway and/or may be manifest only in particular cell types and may result in a measurable biological change.

As used herein, "patient" means a warm blooded animal, such as for example rat, mice, dogs, cats, guinea pigs, and primates such as humans.

As used herein, the expression "pharmaceutically acceptable carrier" means a non-toxic solvent, dispersant, excipient, adjuvant, or other material which is mixed with the compound of the present invention in order to permit the formation of a pharmaceutical composition, i.e., a dosage form capable of administration to the patient. One example of such a carrier is a pharmaceutically acceptable oil typically used for parenteral administration.

The term "pharmaceutically acceptable salts" as used herein means that the salts of the compounds of the present invention can be used in medicinal preparations. Other salts may, however, be useful in the preparation of the compounds according to the invention or of their pharmaceutically acceptable salts. Suitable pharmaceutically acceptable salts of the compounds of this invention include acid addition salts which may, for example, be formed by mixing a solution of the compound according to the invention with a solution of a pharmaceutically acceptable acid such as hydrochloric acid, hydrobromic acid, sulfuric acid, methanesulfonic acid, 2-hydroxyethanesulfonic acid, p-toluenesulfonic acid, fumaric acid, maleic acid, hydroxymaleic acid, malic acid, ascorbic acid, succinic acid, glutaric acid, acetic acid, salicylic acid, cinnamic acid, 2-phenoxybenzoic acid, hydroxybenzoic acid, phenylacetic acid, benzoic acid, oxalic acid, citric acid, tartaric acid, glycolic acid, lactic acid, pyruvic acid, malonic acid, carbonic acid or phosphoric acid. The acid metal salts such as sodium monohydrogen orthophosphate and potassium hydrogen sulfate can also be formed. Also, the salts so formed may present either as mono- or di- acid salts and can exist either as hydrated or can be substantially anhydrous. Furthermore, where the compounds of the invention carry an acidic moiety, suitable pharmaceutically acceptable salts thereof may include alkali metal salts, e.g. sodium or potassium salts; alkaline earth metal salts, e.g. calcium or magnesium salts; and salts formed with suitable organic ligands, e.g. quaternary ammonium salts.

The term "therapeutically effective amount" as used herein means an amount of the compound which is effective in treating the named disorder or condition.

The invention also provides pharmaceutical compositions comprising one or more of the compounds according to this invention in association with a pharmaceutically acceptable carrier. Preferably these compositions are in unit dosage forms such as tablets, pills, capsules, powders, granules, sterile parenteral solutions or suspensions, metered aerosol or liquid sprays, drops, ampoules, auto-injector devices or suppositories; for oral, parenteral, intranasal, sublingual or rectal administration, or for administration by inhalation or insufflation.

Alternatively, the compositions may be presented in a form suitable for once-weekly or once-monthly administration; for example, an insoluble salt of the active compound, such as the

decanoate salt, may be adapted to provide a depot preparation for intramuscular injection. An erodible polymer containing the active ingredient may be envisaged. For preparing solid compositions such as tablets, the principal active ingredient is mixed with a pharmaceutical carrier, e.g. conventional tableting ingredients such as corn starch, lactose, sucrose, sorbitol, talc, stearic acid, magnesium stearate, dicalcium phosphate or gums, and other pharmaceutical diluents, e.g. water, to form a solid preformulation composition containing a homogeneous mixture of a compound of the present invention, or a pharmaceutically acceptable salt thereof.

When referring to these preformulation compositions as homogeneous, it is meant that the active ingredient is dispersed evenly throughout the composition so that the composition may be readily subdivided into equally effective unit dosage forms such as tablets, pills and capsules. This solid preformulation composition is then subdivided into unit dosage forms of the type described above containing from 0.1 to about 500 mg of the active ingredient of the present invention. Flavored unit dosage forms contain from 1 to 100 mg, for example 1, 2, 5,

10, 25, 50 or 100 mg, of the active ingredient. The tablets or pills of the novel composition can be coated or otherwise compounded to provide a dosage form affording the advantage of

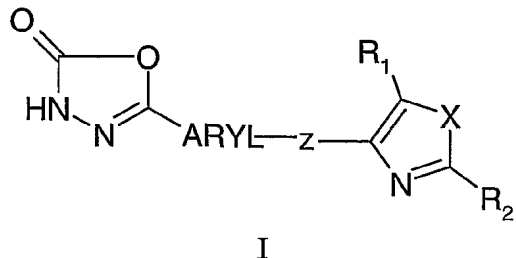
prolonged action. For example, the tablet or pill can comprise an inner dosage and an outer dosage component, the latter being in the form of an envelope over the former. The two components can be separated by an enteric layer which serves to resist disintegration in the stomach and permits the inner component to pass intact into the duodenum or to be delayed in release. A variety of materials can be used for such enteric layers or coatings, such materials including a number of polymeric acids and mixtures of polymeric acids with such materials as shellac, cetyl alcohol and cellulose acetate.

The liquid forms in which the novel compositions of the present invention may be incorporated for administration orally or by injection include aqueous solutions, suitably flavored syrups, aqueous or oil suspensions, and flavored emulsions with edible oils such as cottonseed oil, sesame oil, coconut oil or peanut oil, as well as elixirs and similar pharmaceutical vehicles. Suitable dispersing or suspending agents for aqueous suspensions include synthetic and natural gums such as tragacanth, acacia, alginate, dextran, sodium carboxymethylcellulose, methylcellulose, polyvinyl-pyrrolidone or gelatin.

In the treatment of various disease states as described herein, a suitable dosage level is about 0.01 to 250 mg/kg per day, preferably about 0.05 to 100 mg/kg per day, and especially about 0.05 to 20 mg/kg per day. The compounds may be administered on a regimen of 1 to 4 times per day.

As used in the examples and preparations that follow, the terms used therein shall have the meanings indicated: "kg" refers to kilograms, "g" refers to grams, "mg" refers to milligrams, "g" refers to micrograms, "pg" refers to picograms, "mol" refers to moles, "mmol" refers to millimoles, "nmole" refers to nanomoles, "L" refers to liters, "mL" or "ml" refers to milliliters, "μL" refers to microliters, "°C" refers to degrees Celsius, "R_f" refers to retention factor, "mp" or "m.p." refers to melting point, "dec" refers to decomposition, "bp" or "b.p." refers to boiling point, "mm of Hg" refers to pressure in millimeters of mercury, "cm" refers to centimeters, "nm" refers to nanometers, "[α]²⁰_D" refers to specific rotation of the D line of sodium at 20°C obtained in a 1 decimeter cell, "c" refers to concentration in g/mL, "THF" refers to tetrahydrofuran, "DMF" refers to dimethylformamide, "NMP" refers to 1-methyl-2-pyrrolidinone, "brine" refers to a saturated aqueous sodium chloride solution, "M" refers to molar, "mM" refers to millimolar, "M" refers to micromolar, "nM" refers to nanomolar, "TLC" refers to thin layer chromatography, "HPLC" refers to high performance liquid chromatography, "HRMS" refers to high resolution mass spectrum, "CIMS" refers to chemical ionization mass spectrometry, "ESI" refers to electrospray ionization mass spectrometry, "t_R" refers to retention time, "lb" refers to pounds, "gal" refers to gallons, "L.O.D." refers to loss on drying, "μCi" refers to microcuries, "i.p." refers to intraperitoneally, "i.v." refers to intravenously.

In one aspect of this invention there is disclosed novel compounds having the general structure shown in formula I:



wherein

ARYL is phenyl or pyridinyl, wherein said phenyl or pyridinyl is optionally substituted with

5 one or more substituents selected from the group consisting of halogen, C₁₋₆alkyl, C₂₋₆alkenyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl; C₁₋₆alkylthio, hydroxy, hydroxyC₁₋₆alkyl, C₁₋₄acyloxy, nitro, cyano, C₁₋₆alkylsulfonyl, amino, C₁₋₆alkylamino and C₁₋₆alkoxycarbonyl;

Z is -O(CH₂)_n-, -SO₂(CH₂)_n-, -(CH₂)_n-Y-(CH₂)_n-, -(CH₂)_n-CO-, -O(CH₂)_n-CO-,

10 or -(CH₂)_n-Y-(CH₂)_n-CO- wherein Y is NR₃, O or S and R₃ is selected from the group consisting of H, C₁₋₆alkyl C₃₋₈cycloalkyl, C₁₋₆alkylC₃₋₈cycloalkyl and benzyl and n is independently an integer from 1 to 5;

X is NR₃, O or S wherein R₃ is as defined above;

R₁ is H, halogen, C₁₋₆alkyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl; hydroxyC₁₋₆alkyl, nitro, cyano, and C₁₋₆alkylamino; and

R₂ is substituted or unsubstituted phenyl, pyridinyl or thienyl wherein the substituents are selected from the group consisting of halogen, C₁₋₆alkyl, C₂₋₆alkenyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl, C₁₋₆alkylthio, hydroxy, hydroxyC₁₋₆alkyl, C₁₋₄acyloxy, nitro, cyano, C₁₋₆alkylsulfonyl, amino, C₁₋₆alkylamino and C₁₋₆alkoxycarbonyl;

20 with the proviso that when Z is -O(CH₂)_n- or -SO₂(CH₂)_n-, and ARYL is phenyl then R₂ is other than phenyl;

or a stereoisomer, a tautomer or a solvate thereof or a pharmaceutically acceptable salt thereof.

In a further aspect of this embodiment, is disclosed a compound wherein ARYL is

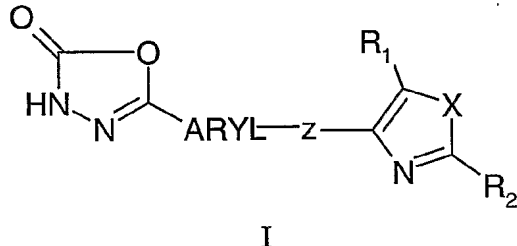
25 phenyl and X is O or S.

In another aspect of this embodiment, is disclosed a compound wherein X is O.

A compound exemplary of this embodiment is 5-(4-{2-[5-methyl-2-(4-trifluoromethyl-phenyl)-thiazol-4-yl]-ethoxy}-phenyl)-3H-[1,3,4]oxadiazol-2-one.

In another embodiment of the present invention, is disclosed a pharmaceutical composition comprising an effective amount of a compound of formula I and a pharmaceutical acceptable carrier.

In another embodiment of the present invention, is disclosed a method of treating a disease in a mammal wherein the disease is capable of being, modulated by PPARdelta ligand binding activity, which comprises administering to said mammal having said disease a therapeutically effective amount of a compound of formula I.



10 wherein

ARYL is phenyl or pyridinyl, wherein said phenyl or pyridinyl is optionally substituted with one or more substituents selected from the group consisting of halogen, C₁₋₆alkyl, C₂₋₆alkenyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl; C₁₋₆alkylthio, hydroxy, hydroxyC₁₋₆alkyl, C₁₋₄acyloxy, nitro, cyano, C₁₋₆alkylsulfonyl, amino, C₁₋₆alkylamino and C₁₋₆alkoxycarbonyl;

15 Z is -O(CH₂)_n-, -SO₂(CH₂)_n-, -(CH₂)_n-Y-(CH₂)_n-, -(CH₂)_n-CO-, -O(CH₂)_n-CO- or -(CH₂)_n-Y-(CH₂)_n-CO- wherein Y is NR₃, O or S and R₃ is selected from the group consisting of H, C₁₋₆alkyl C₃₋₈cycloalkyl, C₁₋₆alkylC₃₋₈cycloalkyl and benzyl and n is independently an integer from 1 to 5;

20 X is NR₃, O or S wherein R₃ is as defined above;

R₁ is H, halogen, C₁₋₆alkyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl; hydroxyC₁₋₆alkyl, nitro, cyano, and C₁₋₆alkylamino; and

25 R₂ is substituted or unsubstituted phenyl, pyridinyl or thienyl wherein the substituents are selected from the group consisting of halogen, C₁₋₆alkyl, C₂₋₆alkenyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl, C₁₋₆alkylthio, hydroxy, hydroxyC₁₋₆alkyl, C₁₋₄acyloxy, nitro, cyano, C₁₋₆alkylsulfonyl, amino, C₁₋₆alkylamino and C₁₋₆alkoxycarbonyl; or a stereoisomer, a tautomer or a solvate thereof or a pharmaceutically acceptable salt thereof.

In a further aspect of this embodiment, of the method of the invention is disclosed a compound wherein ARYL is phenyl.

In another aspect of this embodiment, of this method of the invention is disclosed a compound wherein ARYL is phenyl and R₂ is phenyl.

In a further aspect of this embodiment, of this method of the invention is disclosed a compound wherein ARYL is phenyl, Z is -O(CH₂)_n- and R₂ is phenyl.

5 In yet another aspect of this embodiment, of this method of the invention is disclosed a compound wherein ARYL is phenyl, Z is -O(CH₂)_n-, X is O or S and R₂ is phenyl.

In another aspect of this embodiment, of this method of the invention a compound wherein ARYL is phenyl, Z is -O(CH₂)_n-, X is O or S, R₁ is C₁-alkyl and R₂ is phenyl.

10 In a further aspect of this embodiment, of this method of the invention is disclosed a compound wherein X is O.

In yet another aspect of this invention, of this method of the invention is disclosed a compound wherein X is S.

15 In a further aspect of this embodiment, is disclosed a method wherein said disease is a demyelinating disease selected from the group consisting of multiple sclerosis, Charcot-Marie-Tooth disease, Pelizaeus-Merzbacher disease, encephalomyelitis, neuromyelitis optica, adrenoleukodystrophy, Guillain-Barre syndrome and disorders in which myelin forming glial cells are damaged including spinal cord injuries, neuropathies and nerve injury.

In another aspect of this embodiment, is disclosed a method wherein the demyelinating disease is multiple sclerosis.

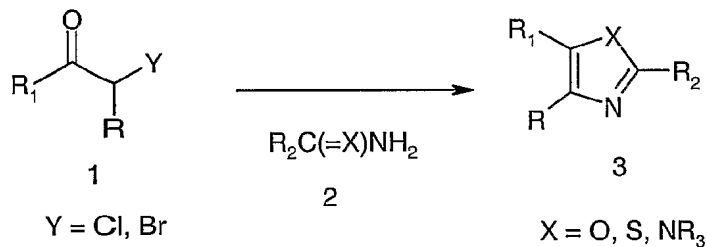
20 In still another aspect of this invention is disclosed a method wherein said disease is selected from the group consisting of obesity, hypertriglyceridemia, hyperlipidemia, hypoalphalipoproteinemia, hypercholesterolemia, dyslipidemia, Syndrome X, Type II diabetes mellitus and complications thereof selected from the group consisting of neuropathy, nephropathy, retinopathy and cataracts, hyperinsulinemia, impaired glucose tolerance, insulin 25 resistance, atherosclerosis, hypertension, coronary heart disease, peripheral vascular disease or congestive heart failure.

30 The compounds disclosed herein can be synthesized according to the following procedures of Schemes, wherein the Aryl, X, Z and R substituents are as identified for formula (I), above unless otherwise noted. If necessary, in the following synthetic schemes, reactive functional groups present in the compounds described in this invention may be protected by suitable protecting groups. The protecting group may be removed at a later stage of the synthesis. Procedures for protecting reactive functional groups and their subsequent

removal may be found in T. W. Greene and P. G. M. Wuts, *Protective Groups in Organic Synthesis*, Wiley and Sons, 1991.

Scheme A shows the synthesis of the appropriate imidazole, oxazole or thiazole, intermediates for compounds of formula I wherein X is O, S or NR₃. The heterocycles can be 5 prepared using methods known in the chemical literature (for reviews see Katritzky, A.R.; Rees, C.W. Eds. *Comprehensive Heterocyclic Chemistry*, Vol. 5; Pergamon Press (1984); Katritzky, A.R.; Rees, C.W.; Scriven, E.F.V. Eds. *Comprehensive Heterocyclic Chemistry II*; Vols 3 & 4, Pergamon Press (1996)). Specifically, said oxazoles, imidazoles and thiazoles 10 can be prepared by fusion of an appropriate α halo-ketone 1, respectively, with an amide, amidine or a thioamide (general formula 2), at temperatures ranging from about 40 °C to 150 °C to give the intermediate heterocycles 3.

Scheme A



15

In Scheme B the general synthesis of compounds of formula I wherein Z is $-\text{O}(\text{CH}_2)_n-$ is shown. Accordingly, in Step B1 the appropriately substituted carboxylic acid ester 4, which can be synthesized as illustrated in Scheme A is reduced to the alcohol 5 by methods that are 20 well known in the art. For example, the reduction may be effected by aluminum hydrides such as lithium aluminum hydrides or diisobutylaluminum hydride in an inert solvent. In Step B2, the alcohol functional group in compound 5, is converted to a leaving group to give compound 6, wherein Lg is a leaving group such as halogen, or sulfonate esters, for example mesylates or tosylates. Conversion to the leaving group can be accomplished by reaction of 25 the alcohol with reagents such as N-bromosuccinimide in the presence of triphenylphosphine to produce a compound wherein the leaving group is bromide, or reaction with thionyl chloride to give a compound wherein the leaving group is chloride. If a sulfonate ester is desired, reaction of compound 5 with an appropriate sulfonyl chloride in the presence of a suitable

base would produce the desired sulfonate ester. For example, reaction of compound 5 with methanesulfonyl chloride in the presence of an organic base such as triethylamine or pyridine in an inert solvent would give compound 6 wherein the leaving group is OSO_2CH_3 .

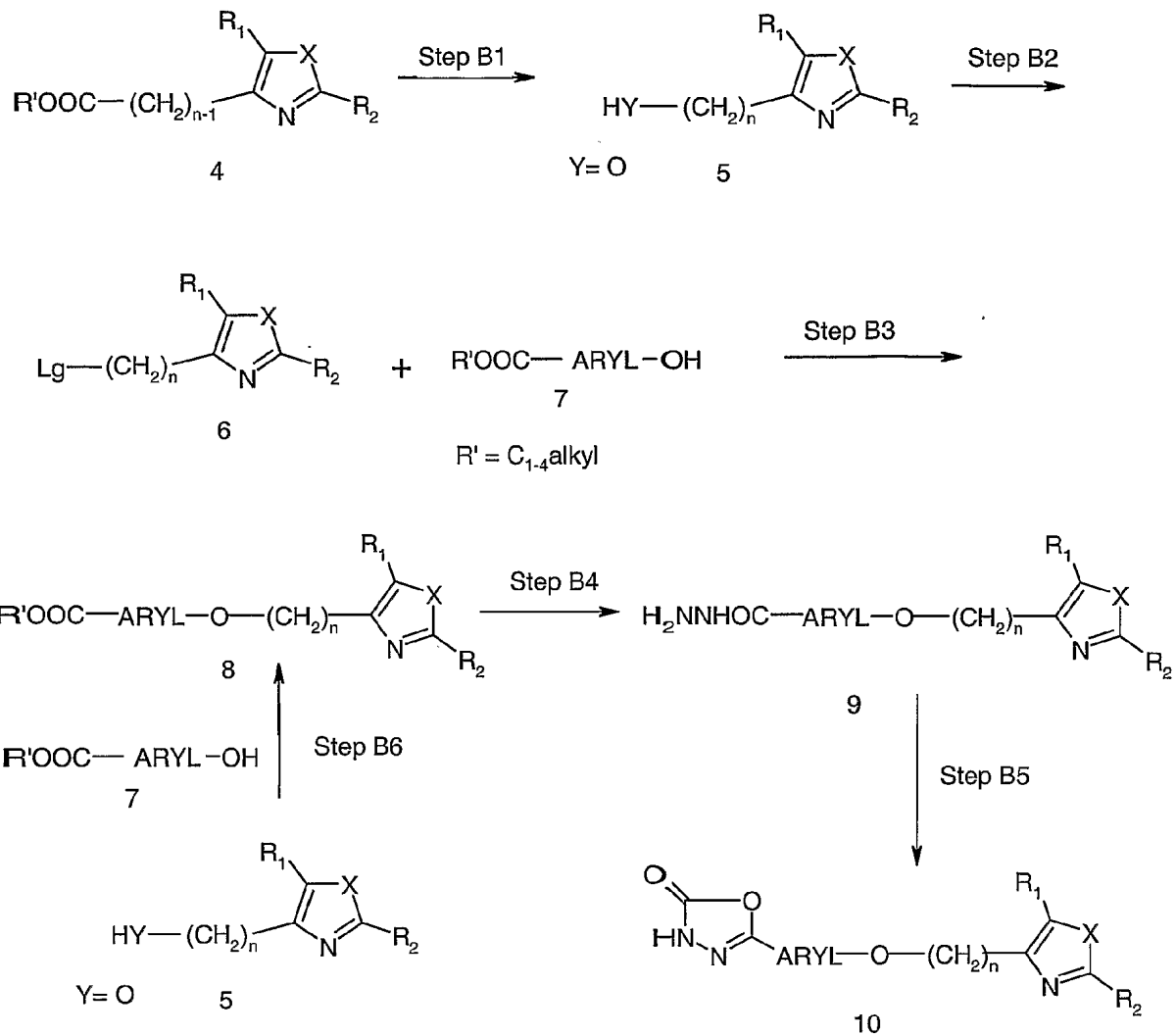
5 In Step B3 an appropriately substituted hydroxy aryl ester, 7 is reacted with the heterocycle, 6 to displace the leaving group to afford coupled ester, 8. The displacement reaction is run under conditions well known in the art. Typically the reaction is run in the presence of a base such as sodium hydride or other inorganic bases such as alkali carbonates or alkali hydroxides in an inert solvent. The temperature of the reaction, although not critical, is from 0°C to the reflux temperature of the inert solvent.

10 Compound 8, in Step B4 is then treated with hydrazine either neat or in a suitable organic solvent at elevated temperatures to give the acid hydrazide, 9. Typically the reaction is run at a temperature of between 50°C and the reflux temperature of the organic solvent.

15 Cyclization of the acid hydrazide 9, in Step B5, to the target 1,3,4-oxadiazol-2-ones, 10 is accomplished by treatment of compound 9 with a chloroformate in the presence of an organic base such as pyridine followed by treatment with a strong, hindered amine base such as 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU) in a suitable organic solvent such as acetonitrile in a sealed tube at elevated temperature. Typically, the reaction can be run from 100°C to 200°C. The 1,3,4-oxadiazol-2-ones may also be synthesized by reacting compound 9 with phosgene. See Stempel, A., et al., *J. Org. Chem.* 1955, 20, 412.

20 In Step B6, an alternative synthesis of the coupled ester, 8 is illustrated. Accordingly, the alcohol, 5 can be reacted with the hydroxy aryl ester, 8 in the presence of a triaryl or trialkylphosphine, such as triphenylphosphine or tri- n-butylphosphine and diethylazodicarboxylate in an inert solvent, for example THF or dichloromethane to afford the coupled ester 8. Typically the reaction is run at a temperature between room temperature and 25 the reflux temperature of the inert solvent.

Scheme B



Scheme C illustrates the synthesis of the compound of formula I wherein Z is $-(CH_2)_n-Y-$.

5 $(CH_2)_n$. The scheme is most useful to synthesize compounds wherein n represents 1 or 2 in the alkylene chain attached to ARYL. In Step C1 compound 5 (Y= O) is converted to compound 6 (wherein Lg is chloro or bromo) as described in Scheme B, Step B2. Compound 6 is then reacted with thiourea, compound 11, under conditions similar to those found in Treau, M. et al. *Heterocycles*, 2001, 55 (9), 1727-1735, to produce the thiol, 5a.

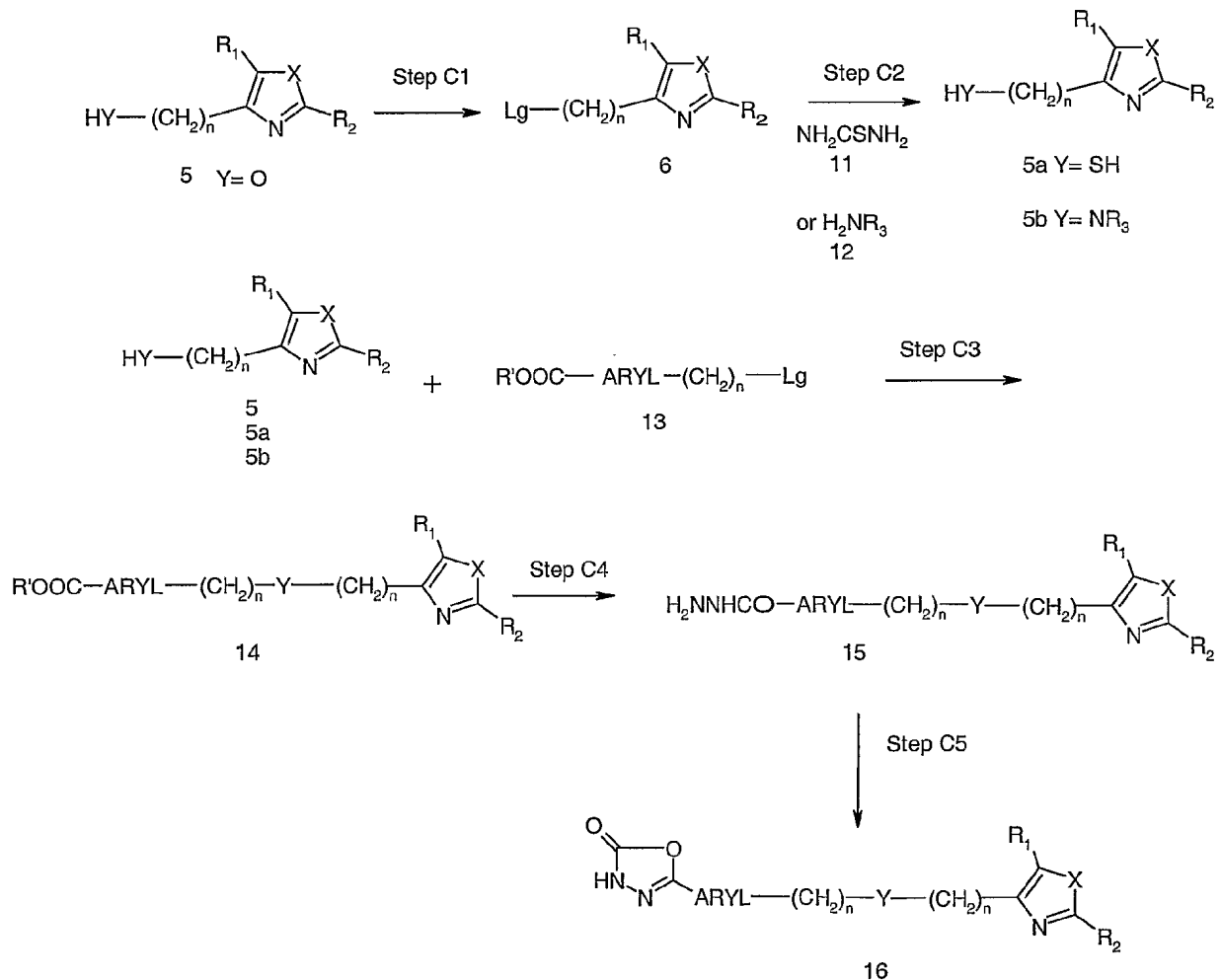
When compound 6 is reacted with a primary amine 12, the aminoalkyl heterocycle 5b is produced. This displacement of a leaving group by an amine is well known to those skilled in the art. Typically, the displacement reaction is run in a polar organic solvent in the presence of an organic base that acts as an acid scavenger. Although not critical the

displacement reaction is run at a temperature of between ambient to reflux temperature of the solvent.

In Step C3 compounds 5, 5a and 5b can be reacted with compound 13 to afford the coupled arylester, 14, wherein Y is O, S or NR₃. Thus, when compounds 5 (Y= O) and 5a (Y= S) are reacted with 13 to displace the leaving group, the reaction will typically be run in the presence of a strong base, for example sodium hydride, in a polar aprotic solvent, such as DMF or DMSO at temperatures of about between 0°C to 150°C. When compound 5b (Y= NR₃) is reacted with 14, conditions identical to those described above in Step C2 for the primary amine are used.

Synthesis of the desired 1,3,4-oxadiazol-2-ones 16, from compound 14 is accomplished in the two steps (C4 and C5) exactly as described in Scheme B, Steps B4 and B5.

Scheme C



In Scheme D an alternative approach to compounds of formula I wherein Z is $-(CH_2)_n-$ $Y-(CH_2)_n-$ is shown. The scheme is most useful to synthesize compounds wherein n represents 3 to 5 in the alkylene chain attached to ARYL.

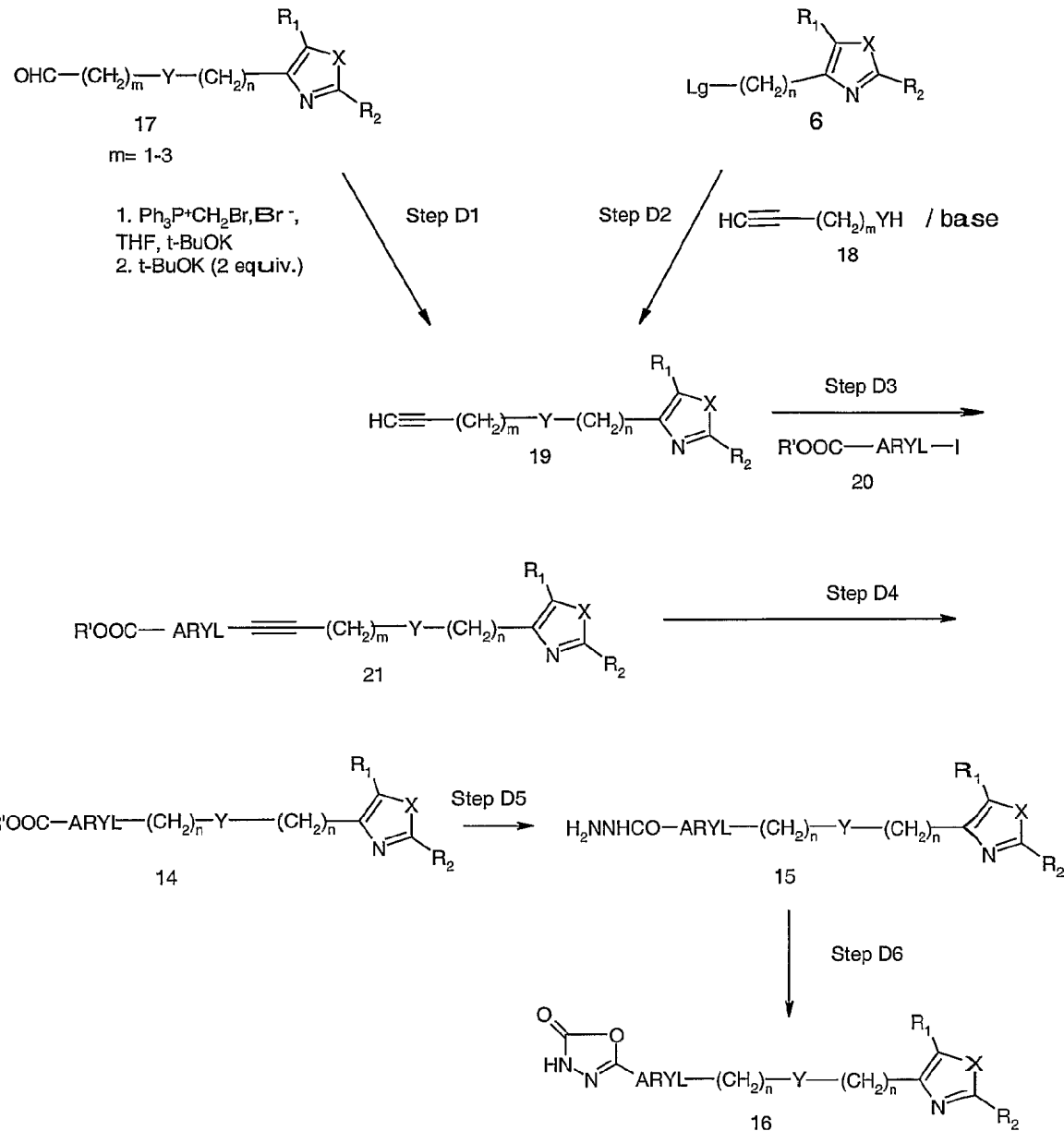
In Step D1 the terminal aldehyde compound 17, which can be synthesized by the 5 method described in Scheme A, is converted in a two-step reaction sequence to the terminal acetylene, 19. Thus, reaction of 17 with bromomethylenetriphenylphosphorane (first step) with potassium t-BuOK produces an intermediate bromoolefin (not shown), which is subsequently treated with 2 equivalents of t-BuOK (second step) to the form the acetylene, 19. The reaction sequence for the conversion is described in Pianetti, P., *Tet. Letters*, 1986, 48, 10 5853-5856. Also, see Corey, E. J., et al. *J. Am. Chem. Soc.*, 1969, 91, 4318-4320.

Alternatively, as shown in Step D2 intermediates of the type 19 can be prepared by displacement of a leaving group from an intermediate such as 6 (see Scheme C) using a nucleophile, such as 18, wherein a terminal acetylene is incorporated.

In Step D3, Sonogashira coupling of acetylenic intermediate, 19 with the aryl iodide, 15 20 is effected in the presence of tetrakistriphenylphosphinepalladium (0), cuprous iodide and a suitable organic base in an inert solvent to yield the coupled terminal acetylene 21. The reduction of the acetylene, 21 can then be accomplished in Step D4 by catalytic hydrogenation of compound 21 to give the saturated ester 14. Typically, the reduction can be accomplished by use of catalysts such as palladium on carbon or chlorotris(triphenylphosphine)rhodium(I) 20 in an inert organic solvent with hydrogen at pressures between 30 to 300 p.s.i.. The reduction can be run at a temperature between room temperature and 175°C.

Synthesis of the desired 1,3,4-oxadiazol-2-ones 16, from compound 14 is accomplished in two steps (D5 and D6) exactly as described in Scheme B, Steps B4 and B5.

Scheme D



Scheme E illustrates a particular synthesis of compounds of formula I wherein Z is —

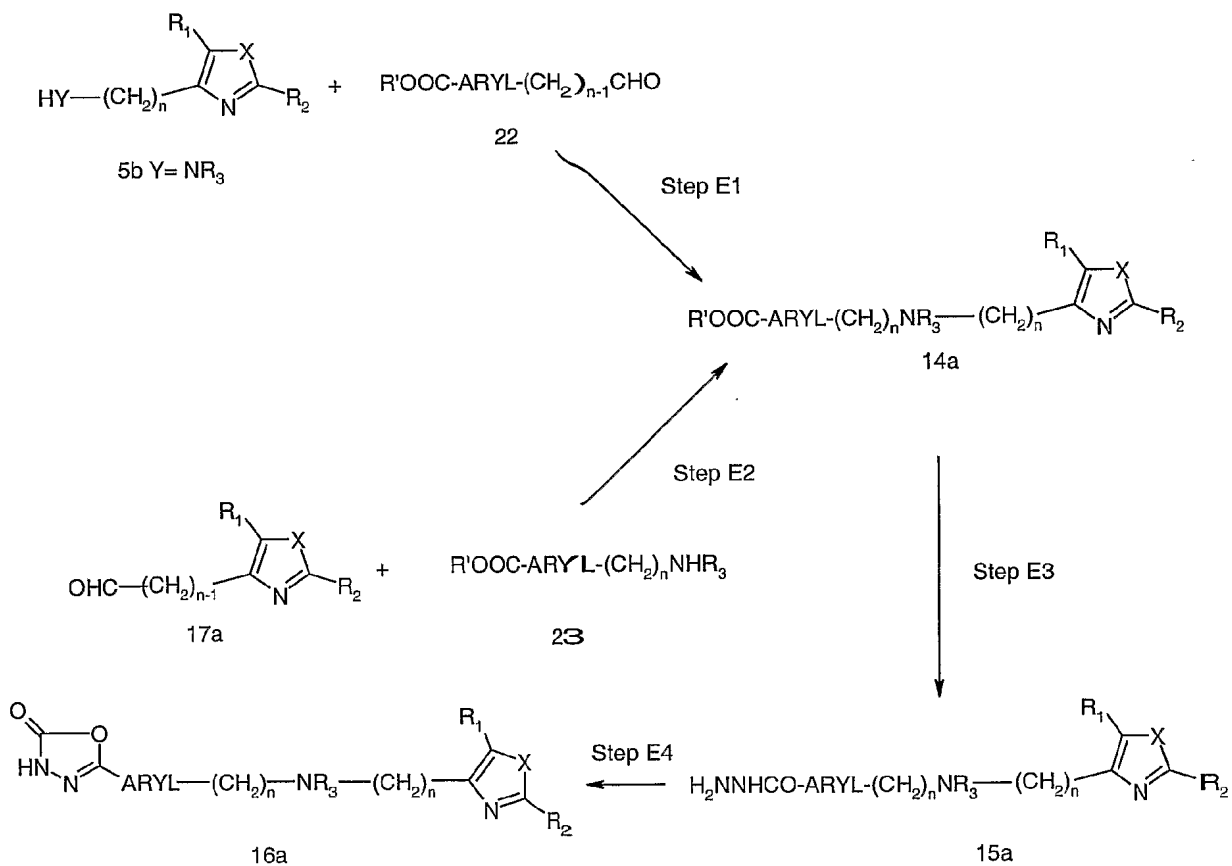
5 (CH₂)_nNR₃(CH₂)_n. In this approach, the linker Z is constructed by a reductive amination of an aldehyde with an amine. For example, in Step E1 treatment of 5b (wherein Y = NR₃) with an aldehyde, such as 4-formyl-benzoic acid methyl ester (n = 1) compound 22, in a polar solvent, usually an alcohol or an alcohol/THF mixture, followed by treatment with a reducing agent such as sodium triacetoxyborohydride provides the required intermediate 14a (n = 1).

10 Similarly, in Step E2 treatment of an aldehyde such as 17a with an amine, such as 4-aminoalkyl benzoic acid methyl ester (n = 1), compound 23, provides 14a, wherein n is 1 and

R_3 is H for $-(CH_2)_nNR_3$. Compound 14a in steps E3 and E4 is converted to 1,3,4,-oxadiazol-2-ones 16a as described in Scheme B, Steps B4 and B5.

More generally, appropriate amines ($R'COOC-ARYL-(CH_2)_nNHR_3$) are prepared from the corresponding nitriles or nitro compounds by catalytic hydrogenation or from acetylenic amines and an aryl iodide or bromide by Sonogashira coupling followed by catalytic hydrogenation as described in Scheme D.

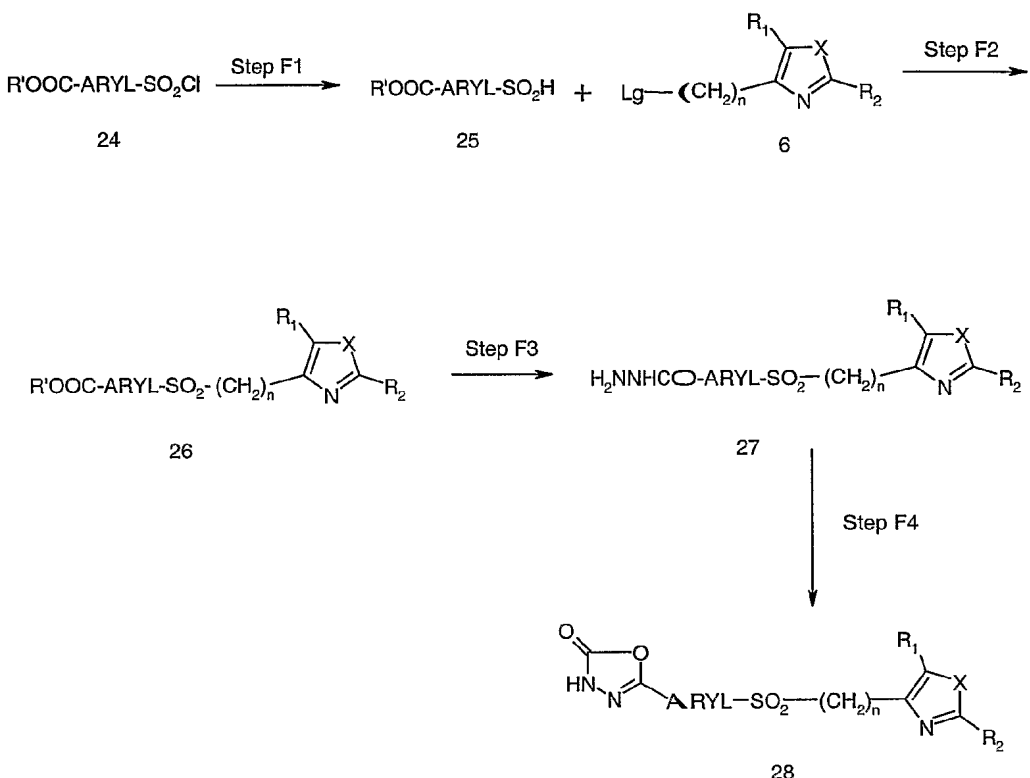
Scheme E



10

Scheme F illustrates the synthesis of compounds of formula I wherein Z is $-SO_2$ $(CH_2)_n-$. In Step F1 treatment of an aryl sulfonyl chloride, 24 with aqueous sodium sulfite provides the sulfinic acid, 25. Reaction of 25, as in Step F2, with an intermediate such as 6 in a polar solvent such as DMF, acetonitrile or ethanol in the presence of a base such as DBU, pyridine, sodium methoxide or sodium hydroxide provides intermediate 26. Intermediate 26 is converted to the corresponding 1,3,4-oxadiazol-2-one, 28 in Steps F3 and F4 as illustrated in Scheme B, Steps B4 and B5.

Scheme F



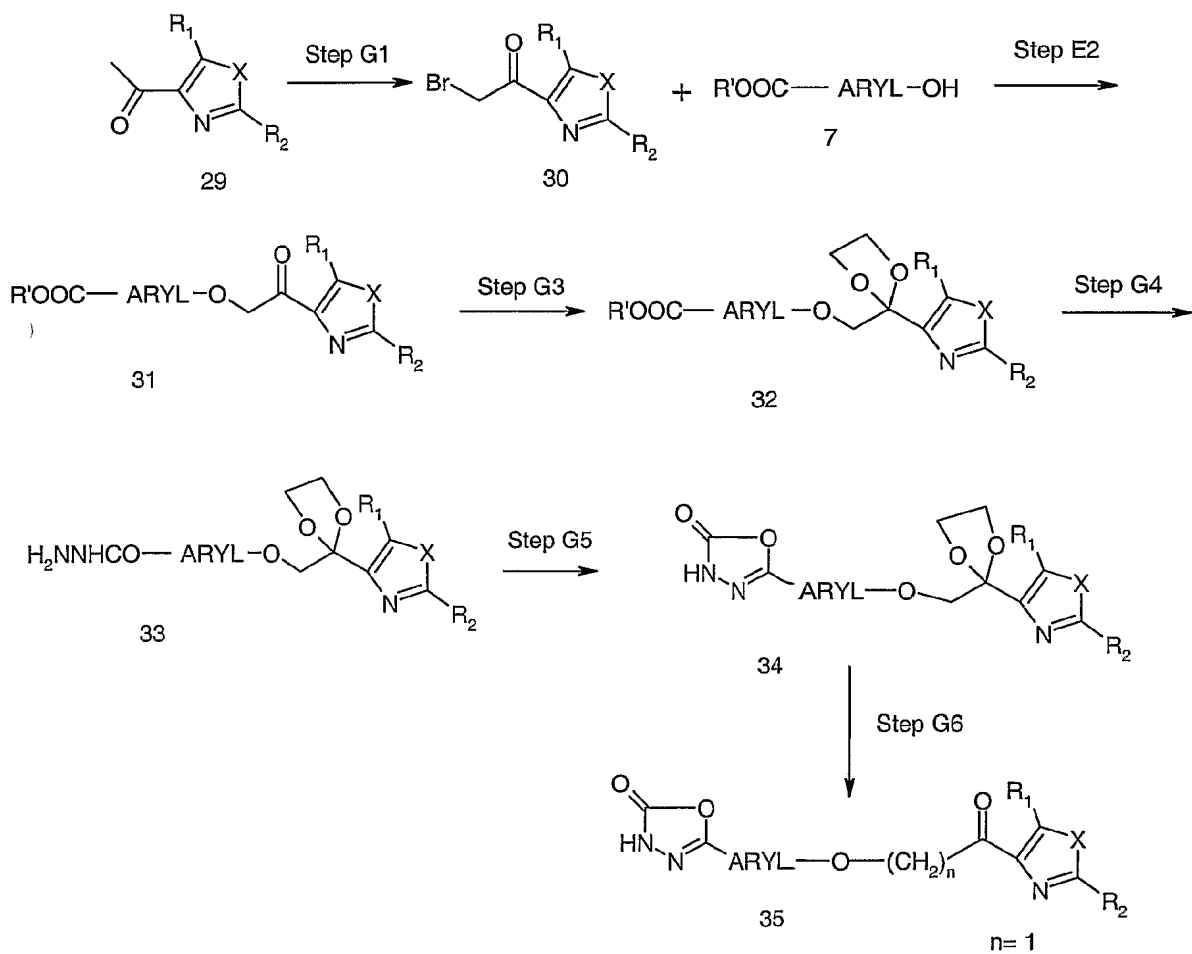
5 Scheme G illustrates the synthesis of compound of formula I wherein Z is—
O(CH₂)_nCO-. The scheme illustrates the case wherein n is 1. The starting 2-acyl heterocycle,
29 can be synthesized from the corresponding carboxylic acid (prepared by the method
illustrated in Scheme A) by addition of an appropriate Grignard reagent to an intermediate N-
methoxy N-methyl carboxamide (Khlestkin, V.K. et al.; Current Organic Chemistry, 2003,
10 7(10), 967-993. and Singh, J. et al., Journal für Praktische Chemie, 2000, 342, 340-347).
Preparation of the intermediate N-methoxy-N-methyl carboxamide is most conveniently
carried out by reaction of the acid with N-methoxy-N-methyl hydroxylamine hydrochloride in
the presence of a peptide coupling reagent such as EDC, DCC, DMPU and a tertiary amine
base such as diisopropylethylamine or triethylamine.

15 Thus in hand, 29 is brominated to produce the bromoketone 30, as shown in Step G1. The bromination can be accomplished by well-known methods, for example reaction of 29 with pyridinium bromide perbromide in acetic acid or reaction of 29 with Br_2 in an inert organic solvent such as dichloromethane. The resulting bromoketone 30, in Step G2, is reacted with the arylhydroxy ester 7 under conditions described in Scheme B (Step B3) to

afford the coupled ester 31. The ketone functionality in 31 is protected as a ketal 32, as shown in Step G3 by methods well known in the art. Compound 32 is then converted to the 1,3,4-oxadizol-2-one ketal 34 in Steps G4 to G5 by the standard sequence as described in Scheme B (B4 and B5). Finally, in Step G6, the ketal functionality in 34 is cleaved, for example, with mineral acid in THF-methanol-water or other methods known in the art to afford the target structure 35.

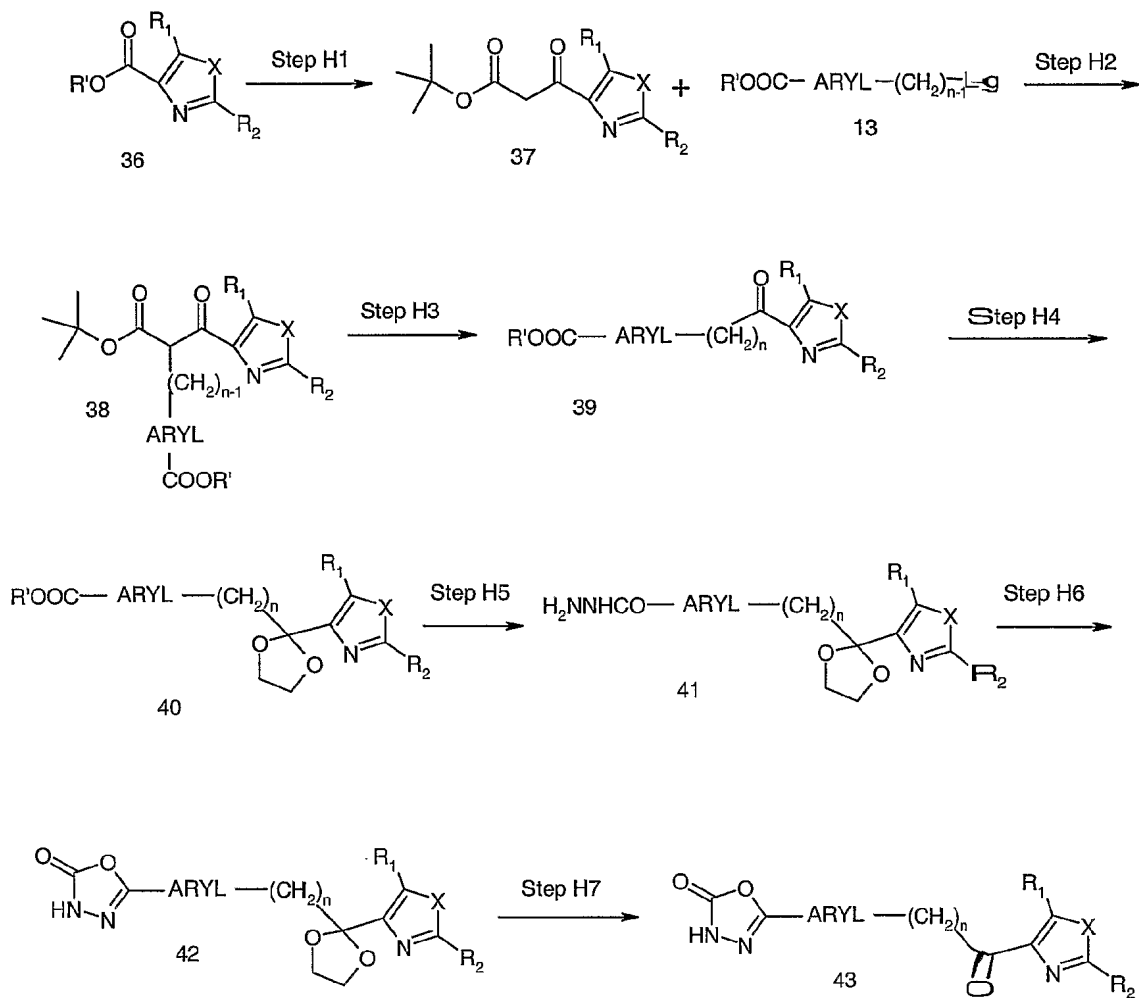
It would be evident to one skilled in the art that the above procedure of Scheme G could be used to synthesize analogs where n is 2-5 for compound 35 by starting with a bromoketone, compound 30, with a larger bromoalkanoyl substituent ($\text{Br}(\text{CH}_2)_n\text{CO}-$, wherein 10 n is 2 to 5).

Scheme G



Scheme H illustrates the procedure for the preparation of compounds of formula I wherein Z is $-(\text{CH}_2)_n\text{CO}-$. In Step H1, the appropriate methoxycarbonyl-substituted heterocycle, 36 is treated with 2 equivalents of the lithium enolate of t-butylacetate in a solvent such as THF or DME at a temperature ranging from -78°C to room temperature to provide the ketoacetate intermediate 37. In Step H2, treatment of 37 with a base such as sodium hydride in an inert solvent at a temperature between -10°C and room temperature followed by alkylation of the resulting anion with an electrophile such as 13 yields the advanced intermediate ketodiester 38. The decarboxylation shown in Step H3 and can be accomplished by first treatment of 38 with TFA in an inert solvent such as dichloromethane followed by thermolysis at a temperature between 70°C and 150°C to provide intermediate ketoester 39. The ketone functionality in 39 is protected as a ketal 40, as shown in Step H4 by methods well known in the art. Compound 40 is then converted to the 1,3,4-oxadiazol-2-one ketal 42 in Steps H5 to H6 by the standard sequence as described in Scheme B (B4 and B5). Finally, in Step H7, the ketal functionality in 42 is cleaved, as described above in Scheme G, Step G6 to afford the desired 1,3,4-oxadiazol-2-one, compound 43.

Scheme H



5

Biological Examples:

The following test protocols are used to ascertain the biological properties of the compounds of this invention. The following examples are being presented to further illustrate the invention. However, they should not be construed as limiting the invention in any manner.

10

Determination of EC₅₀ values in the cell based PPARdelta-GAL4 assay:Principle

The potency of substances, which bind to human PPAR delta and activate it in an agonistic manner, is analyzed using a stably transfected HEK cell line (HEK= human embryo kidney) which is referred to here as PPAR delta reporter cell line. The PPAR delta reporter

cell line contains two genetic elements, a luciferase reporter element (pdeltaM-GAL4-Luc-Zeo) and a PPAR delta fusion protein (GR-GAL4-humanPPAR delta-LBD), which mediates expression of the luciferase reporter element depending on a PPAR delta ligand. The stably and constitutively expressed fusion protein GR-GAL4-humanPPAR delta-LBD binds in the 5 cell nucleus of the PPAR delta reporter cell line via the GAL4 protein portion to the GAL4 DNA binding motifs 5'-upstream of the luciferase reporter element which is stably integrated in the genome of the cell line. There is only little expression of the luciferase reporter gene in the absence of a PPAR delta ligand if fatty acid-depleted fetal calf serum (cs-FCS) is used in the assay. PPAR delta ligands bind and activate the PPAR delta fusion protein and thereby 10 stimulate expression of the luciferase reporter gene. The luciferase, which is formed can be detected by means of chemiluminescence via an appropriate substrate.

Construction of the PPAR delta reporter cell line:

15 The production of the stable PPAR delta reporter cell line is based on a stable HEK-cell clone which was stably transfected with a luciferase reporter element. This step was already described above in the section "construction of the PPAR alpha reporter cell line". In a second step, the PPAR delta fusion protein (GR-GAL4-humanPPAR delta-LBD) was stably introduced into this cell clone. For this purpose, the cDNA coding for the N-terminal 20 76 amino acids of the glucocorticoid receptor (Accession # P04150) was linked to the cDNA section coding for amino acids 1-147 of the yeast transcription factor GAL4 (Accession # P04386). The cDNA of the ligand-binding domain of the human PPAR delta receptor (amino acids S139-Y441; Accession # L07592) was cloned in at the 3'-end of this GR-GAL4 construct. The fusion construct prepared in this way (GR-GAL4-humanPPAR delta-LBD) was recloned into the plasmid pcDNA3 (Invitrogen) in order to enable constitutive expression by 25 the cytomegalovirus promoter. This plasmid was linearized with a restriction endonuclease and stably transfected into the previously described cell clone containing the luciferase reporter element. The resulting PPAR delta reporter cell line which contains a luciferase reporter element and constitutively expresses the PPAR delta fusion protein (GR-GAL4-human PPAR delta-LBD) was isolated by selection with zeocin (0.5 mg/ml) and G418 30 (0.5 mg/ml).

Assay procedure and evaluation:

The activity of PPAR delta agonists is determined in a 3-day assay, which is described below:

5

Day 1

The PPAR delta reporter cell line is cultivated to 80% confluence in DMEM (# 41965-039, Invitrogen) which is mixed with the following additions: 10% cs-FCS (fetal calf serum; #SH-30068.03, Hyclone), 0.5 mg/ml zeocin (#R250-01, Invitrogen), 0.5 mg/ml G418 (#10131-027, Invitrogen), 1% penicillin-streptomycin solution (#15140-122, Invitrogen) and 2 mM L-glutamine (#25030-024, Invitrogen). The cultivation takes place in standard cell culture bottles (# 353112, Becton Dickinson) in a cell culture incubator at 37°C in the presence of 5% CO₂. The 80%-confluent cells are washed once with 15 ml of PBS (#14190-094, Invitrogen), treated with 3 ml of trypsin solution (#25300-054, Invitrogen) at 37°C for 2 min, taken up in 5 ml of the DMEM described and counted in a cell counter. After dilution to 500.000 cells/ml, 35,000 cells are seeded in each well in a volume of 180 µL of a 96 well microtiter plate with a clear plastic base (#3610, Corning Costar). The plates are incubated in the cell culture incubator at 37°C and 5% CO₂ for 24 h.

15

Day 2

PPAR delta agonists to be tested are dissolved in DMSO in a concentration of 10 mM. This stock solution is diluted in DMEM (#41965-039, Invitrogen) which is mixed with 5% cs-FCS (#SH-30068.03, Hyclone), 2 mM L-glutamine (#25030-024, Invitrogen) and the previously described antibiotics (zeocin, G418, penicillin and streptomycin).

20

Test substances are tested in 11 different concentrations in the range from 10 µM to 100 pM. More potent compounds are tested in concentration ranges from 1 µM to 10 pM or between 100 nM and 1 pM.

25

The medium of the PPAR delta reporter cell line seeded on day 1 is completely removed by aspiration or not, and the test substances diluted in medium are immediately added to the cells. The dilution and addition of the substances is carried out by a robot (Beckman FX). The final volume of the test substances diluted in medium is 100 µl per well of a 96 well microtiter plate. The DMSO concentration in the assay is less than 0.1 % v/v

in order to avoid cytotoxic effects of the solvent.

Each plate was charged with a standard PPAR delta agonist, which was likewise diluted in 11 different concentrations, in order to demonstrate the functioning of the assay in each individual plate. The assay plates are incubated in an incubator at 37°C and 5% CO₂ for 24 h.

5 Alternatively, 20µL of a 10x final concentration of the test substance is added directly to the 180 µL containing the plated cells. The test substances are tested in 8 different concentrations, in triplicate, in this assay plate set-up.

Day 3

10 The PPAR delta reporter cells treated with the test substances are removed from the incubator, and the medium is aspirated off. The cells are lysed by pipetting 50 µl of Bright Glo reagent (from Promega) into each well of a 96 well microtiter plate. After incubation at room temperature in the dark for 10 minutes, the microtiter plates are measured in the luminometer (Trilux from Wallac). The measuring time for each well of a microtiter plate is 1
15 sec.

Evaluation:

20 The raw data from the luminometer are transferred into a Microsoft Excel file. Dose-effect plots and EC₅₀ values of PPAR agonists are calculated using the XL.Fit program as specified by the manufacturer (IDBS).

PPARdelta EC₅₀ values in the range of 1nM to >10 µM were measured for the PPAR modulators of the examples in this application. Compounds of the invention of formula I can act as agonists or antagonists. The assay to determine partial agonist or antagonist activity is described below.

25 Determination of Effectiveness of Partial Agonists or Antagonists At the PPARdelta Receptor

This assay determines if compounds act as partial agonists or antagonists at the PPARdelta receptor.

30 The plating and harvesting of the assay plates is as described in Day 1 and 3 above.

Day 2

The partial agonist or antagonist and a known selective agonist are diluted in DMEM (#41965-039, Invitrogen), which is mixed with 10% cs-FCS (#SH-30068.03, Hyclone), 2 mM

L-glutamine (#25030-024, Invitrogen) and the previously described antibiotics (zeocin, G418, penicillin and streptomycin) to 20X desired final concentrations. Ten microliters of the partial agonist or antagonist is added to the cell-containing assay plate. The assay plates are incubated in an incubator at 37°C and 5% CO₂ for 30 minutes. Ten microliters of the 20X

5 known selective agonists are then added, after the partial agonist or antagonist pre-incubation. The assay plates are incubated in an incubator at 37°C and 5% CO₂ for 24 h. The effect on the known selective agonists EC₅₀'s is determined for each partial agonist or antagonist concentration.

10 **SPA PPARdelta-LBD Binding Assay**

Stock Solutions:

15 1 M Tris (pH=8.0 or pH=7.6)(Gene Medicine Stock Room)

2 M KCl (Powder in N2140)

Tween 20

100 mM DTT

20 13.9 uM GW2331 in EtOH HOT

10 mM GW 2331 in DMSO COLD

PPAR-alpha (Conc. varies)

Ex: .884 µg/µl

25 **Wash Buffer:(Store at 4°C. Buffer is good for one week)**

30 10 mM Tris(pH=7.6 or 8) 10 ml

50 mM Kcl 25 ml

0.05 % Tween 20 0.5 ml

Millipore Water 964.5

Check PH=7.6

Binding Buffer:(Prepare fresh binding buffer every time)

35 Wash Buffer 50 ml

10 mM DTT 5.5ml

Preparation of Reaction Reagents for 1 Plate:

40 **Glutathione Coated SPA beads**

Each SPA bead bottle contains 500 mg beads

Reconstitute 500mg of SPA beads in 5 ml of wash buffer, and will be good for few weeks)
Store at 4°C

Prepare diluted SPA beads in the binding buffer.

5

Adding 1 ml of above reconstituted SPA beads to 60 ml of Binding buffer
adding 20 µl of above diluted beads to each well of a 96-well plate.

Use 2ml of above diluted beads for each plate (no dead volume included).

10

3H GW-2331 plus GST-PPAR delta-LBD (for one 96-well plate no dead volume) 13.9 µM
40nM / well

3.0 ml/plate (Including dead volume)

15 If 3H-GW2331 specific activity is 1mci/ml(From Amersham), dilute 17 µl of 3H GW-2331
into 3.0 ml of Binding Buffer = .08 µM

If protein concentration is 1mg/mL, add 21 µl of proteins into 3.0 ml of binding buffer.

20 In Summary: ONE 96-well plate: 3000 µL Binding Buffer + 17 µL of 3H-GW2331+21 µL of
GST-PPAR-delta(1mg/ml)

Control Plates

25 A 96-well Mother Plate(For 2 control Plates)

In column #1:

Add 5 µl of cold GW2331(10 mM) to the wells E-H.

Add 45 µL of DMSO to the wells A-H.

30

In column #12 (3-fold dilution):

Add 10 µL of cold GW2331(10 mM) to the well A.

then add 90 µl of DMSO to the well A, mix well the solution.

add 20 μ l of DMSO to the wells B-H.
take 10 μ l solution from the well A to B, mix well,
then take 10 μ l solution from B to C, mix well,
then take 10 μ l solution from C to D, mix well.

5

Finally, take 10 μ l from F to H.

A control Plate (for 8 Reaction Plates)

10 A control plate is 1:10 dilution of the mother plate. The dilution buffer is the wash buffer.

Sample plates

To fresh CPC library plate, add 90 μ l of DMSO

15

Take 10 μ l of DMSO dilution and add it to 90 μ l of wash buffer in a sample plate

Reaction Plates:

20 Add 20 μ l of SPA beads and 30 μ l of 3H-GW2331 with GST-PPAR-delta to each well of a Reaction Plate.

Add 5 μ l compounds from each well of the sample plate into columns 2 to 11 of a reaction plate.

25 Add 5 μ l compounds from column 1 and column 12 of the control plate to the column 1 and column 12 of the reaction plate.

96-well SPA Protocol:

Let reaction plates equilibrate for 20 minutes to 2 hours.

30

Seal the plates before counting in a Microbeta counter (Wallac).

Calculate IC₅₀.

In the SPA PPAR delta-LBD Binding Assay IC₅₀ values in the range of 1 nM to >10 μ M were measured for the PPAR modulators of the examples in this application. Compounds of the invention of formula I can act as agonists or antagonists.

5

RAT/MICE Oligodendrocyte culturesPreparation of cells:

1. Primary rat oligodendrocyte progenitor cells are obtained from the neocortex of newborn (postnatal days 2-3) rats or mice and are enriched, after removal of microglia, by mechanical separation from the astrocytic monolayer using a modification of the technique originally described by McCarthy and de Vellis (1980).
- 10 2. Remove the meninges from neonatal rat brain and mechanically dissociate tissue. Plate cells on T75 flasks and feed cells with DMEM/F12 + 10% FBS.
3. Collect oligodendrocytes growing on the astrocyte bed layer by shaking-off method
15 fourteen days after the original prep date. Centrifuge the suspension and resuspend the cell pellet in serum free media (SFM; DMEM combined with 25 μ g/ml transferring, 30 nM triiodothyronine, 20 nM hydrocortisone, 20 nM progesterone, 10 nM biotin, 1x trace elements, 30 nM selenium, 1 μ g/ml putrescine, 0.1% BSA, 5 U/ml PenStrep, 10 μ g/ml insulin) supplemented with the following growth factors: Platelet derived
20 growth factor- α (PDGF) and fibroblast growth factor-2 (FGF).
4. Plate the cells on PDL-coated dishes and incubate at 37°C with 6-7% CO₂.
5. Media components are replaced every 48 hr to keep the cells in a progenitor state.

Progenitor cell passaging to increase cell numbers for screening assays:

- 25 1. When the culture are confluent, rinse the culture with PBS, add trypsin and incubate for ~2-3 min at 37°C.
2. Neutralize and centrifuge the cell suspension at 900g for 5 min.
3. Resuspend the cell pellet in SFM + PDGF/FGF.
4. Feed the cells with fresh growth factors every 48 hrs to keep enrich for rapidly
30 dividing progenitor cells.
5. Cells are passaged no more than 4-5 times prior to experimental assays.
6. All experiments involving oligodendrocyte progenitor cells were done using cells that were continuously maintained under these conditions. Greater than 95% of all cells

were A2B5 immunopositive and expressed 2' 3' -cyclic nucleotide 3' – phosphodiesterase II mRNA.

7. To generate mature oligodendrocytes, 24 h after plating progenitor cells were switched to SFM supplemented with or without IGF-I and grown under these conditions for 7 d prior to experimental assays.
8. Alternatively, the enriched rat Central Glia-4 (CG4) progenitor cell line may be used, which is maintained in base media (DMEM, with 2 mM glutamine, 1mM sodium pyruvate, biotin (40 nM), insulin (1 μ M) and N1) supplemented with 30% conditioned media from the B-104 neuroblastoma cell line. To induce differentiation, CG4 cells are switched to base media with 1% fetal calf serum (removed after 2 days) and insulin (500 nM). A2B5 and MBP immunoreactivity is used to confirm >95% enrichment in immature and mature cultures, respectively.

Rat/Mouse Culture Compound Treatment:

1. Put 10,000 – 15,000 cells /well in 24-well PDL coated plates and culture the cells in presence of mitogen (10 ng/ml) overnight.
2. In the presence of mitogen:
 - a. Next day, remove the old medium and add compounds in fresh medium (with mitogen)
 - b. Compound dose response evaluations are performed at 6 different concentrations (10 μ M, 1 μ M, 100 nM, 10 nM, 1 nM, and 0.1 nM);
 - c. Triplicates wells are run for each compound concentration.
3. In the absence of mitogen:
 - a. Next day, remove the old medium and add compounds in fresh medium (without mitogen)
 - b. Compound dose response evaluations are performed at 6 concentrations (10 μ M, 1 μ M, 100 nM, 10 nM, 1 nM, and 0.1 nM);
 - c. Triplicates wells are run for each compound concentration.
4. Culture the treated cells for 7 d prior to using in experimental assays.

HUMAN Oligodendrocyte cultures

Preparation of cells:

1. Human neurospheres collected from E19.5 – E22 human embryo cortex) are cultured for 2 weeks in progenitor media: DMEM/F12 containing 100 μ g/ml transferring, 30

nM triiodothyronine, 20 nM hydrocortisone, 20 nM progesterone, 10 nM biotin, 1x trace elements, 30 nM selenium, 60 uM putrescine, 0.1% BSA, 5 U/ml PenStrep, 25 μ g/ml insulin) supplemented with PDGF and FGF.

2. Neurospheres are dissociated with 20 U/ml papain at 37⁰C for 30-50 min.
- 5 3. Cells are plated onto PDL coated dishes at density of 50,000-100,000 cell/well in progenitor media containing PDGF/FGF and incubated at 37⁰C with 5-6% CO₂.
4. Media and growth factors are replenished every 48 hr.

Human Culture Compound Treatment:

- 10 1. 24 to 48 hr after plating remove the old medium and add compounds in fresh medium (with mitogen)
2. Compound dose response evaluations are performed at 3-6 different concentrations (10 μ M, 1 μ M, 100 nM, 10 nM, 1 nM, and 0.1 nM)
3. Triplicates wells are run for each compound concentration.
- 15 5. Culture the treated cells for 7 d prior to using in experimental assays.

RAT/MOUSE/HUMAN Oligodendrocyte Specific Immunostaining:

Following compound exposure, oligodendrocyte-specific antibodies are used to assess ability of compound to accelerate/promote oligodendrocyte differentiation (for example, O4, O1, or 20 myelin basic protein immunoreactivity is over time between compound treated and untreated cultures).

- 25 1. Cells are plated onto poly-D-lysine treated 4-well chamber slides at 5x10³ to 20x10³ cells/well and grown as described above. Sequential staining is performed on oligodendrocyte populations with increasing degrees of cellular differentiation, as determined by days *in vitro* without PDGF and FGF.
2. Live staining for 30 min at 37⁰C is used to detect oligodendrocyte stage specific cell surface marker expression (including A2B5, O4, and O1).
3. Subsequently, cells are fixed with 4% paraformaldehyde, 10 min, room temperature.
4. Fixed staining procedures are used to detect oligodendrocyte stage specific marker expression (including myelin basic protein, MBP).
- 30 5. Rinse with PBS.
6. Permeabilize with 0.1% Triton/0.01% NaAz diluted in 1X PBS for 10 min, room temperature.

7. Block with 5-10% goat serum in antibody dilution buffer (0.1% Triton-X 100 and 1% IgG-free bovine serum albumin; also used to dilute antibodies), 15 min, room temperature.
8. Add primary antibody diluted in antibody dilution buffer.
9. Incubate overnight, gently rocking, 4° C.
- 5 10. Next day, rinse with PBS 1X 5 min, followed by 3X 15 min each, room temperature.
11. Incubate with appropriate secondary antibodies, 45 min, room temperature.
12. Cell nuclei are stained with 4,6-diamidino-2-phenylindole (DAPI), 15 min, room temperature.
- 10 13. Rinse several times with PBS and evaluate using fluorescent microscopy.
14. The following conditions are compared over time and at different compound doses: PDGF/FGF alone, SFM alone, SFM-IGF1 alone, PDGF/FGF and compound, SFM and compound.

RAT/MOUSE/HUMAN Bromodeoxyuridine (BrdU) immunostaining:

15 To confirm that compounds do not promote cell proliferation.

1. Oligodendrocyte progenitor cells are labeled with 10 µM BrdU for 20 hr and then fixed with either 70% ethanol or 4% paraformaldehyde.
2. The cells are incubated successively with biotinylated mouse anti-BrdU and Streptavidin-Peroxidase, with three intervening washes with PBS.
- 20 3. Colormetric visualization of the BrdU immunoreactivity is developed with DAB and total cell numbers are assessed using the counter-stain hematoxylin.
4. BrdU immunopositive cells are counted by two independent observers.

RAT/MOUSE/HUMAN Culture Image analysis: Fluorescent microscopy is used to quantitate the extent of oligodendrocyte differentiation after compound exposure. This assay demonstrates that selective agonists accelerate/promote oligodendrocytes differentiation.

- 25 1. Manual Cell Counting: Four fields are randomly selected for each experimental condition and 500-600 cells are counted in each field. The percentage of MBP (or O4) immunopositive cells (mature process bearing cells with or without myelin sheets) versus DAPI positive cells (total cell number) cells are compared in the control and drug-treated groups.
- 30 2. Automated Cell Counting: Fluorescent microscopy was used to quantitate the extent of oligodendrocyte differentiation after compound exposure. Six fields/well were

randomly selected to assess the number of differentiating oligodendrocytes among the total population (~8 to 15x10³ cells are counted/well). Immunofluorescence images were obtained using a Zeiss AxioVision digital imaging system, with a Zeiss AxioCam HRc cooled CCD camera connected to the same microscope. All microscopic imaging parameters were set for acquiring images for the analysis of cellular immunofluorescence intensity. The percentage of MBP positive (differentiated) cells versus total cells (DAPI nuclear stained) was compared in the control versus drug-treated groups. Cellular autofluorescence was undetectable under the imaging conditions.

5 10 3. Human oligodendrocyte differentiation assay: manually count total number of O4 immunopositive cells/well (bipolar and multipolar).

15 RAT/MOUSE/HUMAN Quantitative Polymerase Chain Reaction (PCR): To evaluate compound induced PPAR delta pathway activation and the extent of oligodendrocyte maturation (changes in mRNA levels).

1 2 3 4 5 20 25 30 35 40 45 50 55 60 65 70 75 80 85 90 95 100 105 110 115 120 125 130 135 140 145 150 155 160 165 170 175 180 185 190 195 200 205 210 215 220 225 230 235 240 245 250 255 260 265 270 275 280 285 290 295 300 305 310 315 320 325 330 335 340 345 350 355 360 365 370 375 380 385 390 395 400 405 410 415 420 425 430 435 440 445 450 455 460 465 470 475 480 485 490 495 500 505 510 515 520 525 530 535 540 545 550 555 560 565 570 575 580 585 590 595 600 605 610 615 620 625 630 635 640 645 650 655 660 665 670 675 680 685 690 695 700 705 710 715 720 725 730 735 740 745 750 755 760 765 770 775 780 785 790 795 800 805 810 815 820 825 830 835 840 845 850 855 860 865 870 875 880 885 890 895 900 905 910 915 920 925 930 935 940 945 950 955 960 965 970 975 980 985 990 995 1000 1005 1010 1015 1020 1025 1030 1035 1040 1045 1050 1055 1060 1065 1070 1075 1080 1085 1090 1095 1100 1105 1110 1115 1120 1125 1130 1135 1140 1145 1150 1155 1160 1165 1170 1175 1180 1185 1190 1195 1200 1205 1210 1215 1220 1225 1230 1235 1240 1245 1250 1255 1260 1265 1270 1275 1280 1285 1290 1295 1300 1305 1310 1315 1320 1325 1330 1335 1340 1345 1350 1355 1360 1365 1370 1375 1380 1385 1390 1395 1400 1405 1410 1415 1420 1425 1430 1435 1440 1445 1450 1455 1460 1465 1470 1475 1480 1485 1490 1495 1500 1505 1510 1515 1520 1525 1530 1535 1540 1545 1550 1555 1560 1565 1570 1575 1580 1585 1590 1595 1600 1605 1610 1615 1620 1625 1630 1635 1640 1645 1650 1655 1660 1665 1670 1675 1680 1685 1690 1695 1700 1705 1710 1715 1720 1725 1730 1735 1740 1745 1750 1755 1760 1765 1770 1775 1780 1785 1790 1795 1800 1805 1810 1815 1820 1825 1830 1835 1840 1845 1850 1855 1860 1865 1870 1875 1880 1885 1890 1895 1900 1905 1910 1915 1920 1925 1930 1935 1940 1945 1950 1955 1960 1965 1970 1975 1980 1985 1990 1995 2000 2005 2010 2015 2020 2025 2030 2035 2040 2045 2050 2055 2060 2065 2070 2075 2080 2085 2090 2095 2100 2105 2110 2115 2120 2125 2130 2135 2140 2145 2150 2155 2160 2165 2170 2175 2180 2185 2190 2195 2200 2205 2210 2215 2220 2225 2230 2235 2240 2245 2250 2255 2260 2265 2270 2275 2280 2285 2290 2295 2300 2305 2310 2315 2320 2325 2330 2335 2340 2345 2350 2355 2360 2365 2370 2375 2380 2385 2390 2395 2400 2405 2410 2415 2420 2425 2430 2435 2440 2445 2450 2455 2460 2465 2470 2475 2480 2485 2490 2495 2500 2505 2510 2515 2520 2525 2530 2535 2540 2545 2550 2555 2560 2565 2570 2575 2580 2585 2590 2595 2600 2605 2610 2615 2620 2625 2630 2635 2640 2645 2650 2655 2660 2665 2670 2675 2680 2685 2690 2695 2700 2705 2710 2715 2720 2725 2730 2735 2740 2745 2750 2755 2760 2765 2770 2775 2780 2785 2790 2795 2800 2805 2810 2815 2820 2825 2830 2835 2840 2845 2850 2855 2860 2865 2870 2875 2880 2885 2890 2895 2900 2905 2910 2915 2920 2925 2930 2935 2940 2945 2950 2955 2960 2965 2970 2975 2980 2985 2990 2995 3000 3005 3010 3015 3020 3025 3030 3035 3040 3045 3050 3055 3060 3065 3070 3075 3080 3085 3090 3095 3100 3105 3110 3115 3120 3125 3130 3135 3140 3145 3150 3155 3160 3165 3170 3175 3180 3185 3190 3195 3200 3205 3210 3215 3220 3225 3230 3235 3240 3245 3250 3255 3260 3265 3270 3275 3280 3285 3290 3295 3300 3305 3310 3315 3320 3325 3330 3335 3340 3345 3350 3355 3360 3365 3370 3375 3380 3385 3390 3395 3400 3405 3410 3415 3420 3425 3430 3435 3440 3445 3450 3455 3460 3465 3470 3475 3480 3485 3490 3495 3500 3505 3510 3515 3520 3525 3530 3535 3540 3545 3550 3555 3560 3565 3570 3575 3580 3585 3590 3595 3600 3605 3610 3615 3620 3625 3630 3635 3640 3645 3650 3655 3660 3665 3670 3675 3680 3685 3690 3695 3700 3705 3710 3715 3720 3725 3730 3735 3740 3745 3750 3755 3760 3765 3770 3775 3780 3785 3790 3795 3800 3805 3810 3815 3820 3825 3830 3835 3840 3845 3850 3855 3860 3865 3870 3875 3880 3885 3890 3895 3900 3905 3910 3915 3920 3925 3930 3935 3940 3945 3950 3955 3960 3965 3970 3975 3980 3985 3990 3995 4000 4005 4010 4015 4020 4025 4030 4035 4040 4045 4050 4055 4060 4065 4070 4075 4080 4085 4090 4095 4100 4105 4110 4115 4120 4125 4130 4135 4140 4145 4150 4155 4160 4165 4170 4175 4180 4185 4190 4195 4200 4205 4210 4215 4220 4225 4230 4235 4240 4245 4250 4255 4260 4265 4270 4275 4280 4285 4290 4295 4300 4305 4310 4315 4320 4325 4330 4335 4340 4345 4350 4355 4360 4365 4370 4375 4380 4385 4390 4395 4400 4405 4410 4415 4420 4425 4430 4435 4440 4445 4450 4455 4460 4465 4470 4475 4480 4485 4490 4495 4500 4505 4510 4515 4520 4525 4530 4535 4540 4545 4550 4555 4560 4565 4570 4575 4580 4585 4590 4595 4600 4605 4610 4615 4620 4625 4630 4635 4640 4645 4650 4655 4660 4665 4670 4675 4680 4685 4690 4695 4700 4705 4710 4715 4720 4725 4730 4735 4740 4745 4750 4755 4760 4765 4770 4775 4780 4785 4790 4795 4800 4805 4810 4815 4820 4825 4830 4835 4840 4845 4850 4855 4860 4865 4870 4875 4880 4885 4890 4895 4900 4905 4910 4915 4920 4925 4930 4935 4940 4945 4950 4955 4960 4965 4970 4975 4980 4985 4990 4995 5000 5005 5010 5015 5020 5025 5030 5035 5040 5045 5050 5055 5060 5065 5070 5075 5080 5085 5090 5095 5100 5105 5110 5115 5120 5125 5130 5135 5140 5145 5150 5155 5160 5165 5170 5175 5180 5185 5190 5195 5200 5205 5210 5215 5220 5225 5230 5235 5240 5245 5250 5255 5260 5265 5270 5275 5280 5285 5290 5295 5300 5305 5310 5315 5320 5325 5330 5335 5340 5345 5350 5355 5360 5365 5370 5375 5380 5385 5390 5395 5400 5405 5410 5415 5420 5425 5430 5435 5440 5445 5450 5455 5460 5465 5470 5475 5480 5485 5490 5495 5500 5505 5510 5515 5520 5525 5530 5535 5540 5545 5550 5555 5560 5565 5570 5575 5580 5585 5590 5595 5600 5605 5610 5615 5620 5625 5630 5635 5640 5645 5650 5655 5660 5665 5670 5675 5680 5685 5690 5695 5700 5705 5710 5715 5720 5725 5730 5735 5740 5745 5750 5755 5760 5765 5770 5775 5780 5785 5790 5795 5800 5805 5810 5815 5820 5825 5830 5835 5840 5845 5850 5855 5860 5865 5870 5875 5880 5885 5890 5895 5900 5905 5910 5915 5920 5925 5930 5935 5940 5945 5950 5955 5960 5965 5970 5975 5980 5985 5990 5995 6000 6005 6010 6015 6020 6025 6030 6035 6040 6045 6050 6055 6060 6065 6070 6075 6080 6085 6090 6095 6100 6105 6110 6115 6120 6125 6130 6135 6140 6145 6150 6155 6160 6165 6170 6175 6180 6185 6190 6195 6200 6205 6210 6215 6220 6225 6230 6235 6240 6245 6250 6255 6260 6265 6270 6275 6280 6285 6290 6295 6300 6305 6310 6315 6320 6325 6330 6335 6340 6345 6350 6355 6360 6365 6370 6375 6380 6385 6390 6395 6400 6405 6410 6415 6420 6425 6430 6435 6440 6445 6450 6455 6460 6465 6470 6475 6480 6485 6490 6495 6500 6505 6510 6515 6520 6525 6530 6535 6540 6545 6550 6555 6560 6565 6570 6575 6580 6585 6590 6595 6600 6605 6610 6615 6620 6625 6630 6635 6640 6645 6650 6655 6660 6665 6670 6675 6680 6685 6690 6695 6700 6705 6710 6715 6720 6725 6730 6735 6740 6745 6750 6755 6760 6765 6770 6775 6780 6785 6790 6795 6800 6805 6810 6815 6820 6825 6830 6835 6840 6845 6850 6855 6860 6865 6870 6875 6880 6885 6890 6895 6900 6905 6910 6915 6920 6925 6930 6935 6940 6945 6950 6955 6960 6965 6970 6975 6980 6985 6990 6995 7000 7005 7010 7015 7020 7025 7030 7035 7040 7045 7050 7055 7060 7065 7070 7075 7080 7085 7090 7095 7100 7105 7110 7115 7120 7125 7130 7135 7140 7145 7150 7155 7160 7165 7170 7175 7180 7185 7190 7195 7200 7205 7210 7215 7220 7225 7230 7235 7240 7245 7250 7255 7260 7265 7270 7275 7280 7285 7290 7295 7300 7305 7310 7315 7320 7325 7330 7335 7340 7345 7350 7355 7360 7365 7370 7375 7380 7385 7390 7395 7400 7405 7410 7415 7420 7425 7430 7435 7440 7445 7450 7455 7460 7465 7470 7475 7480 7485 7490 7495 7500 7505 7510 7515 7520 7525 7530 7535 7540 7545 7550 7555 7560 7565 7570 7575 7580 7585 7590 7595 7600 7605 7610 7615 7620 7625 7630 7635 7640 7645 7650 7655 7660 7665 7670 7675 7680 7685 7690 7695 7700 7705 7710 7715 7720 7725 7730 7735 7740 7745 7750 7755 7760 7765 7770 7775 7780 7785 7790 7795 7800 7805 7810 7815 7820 7825 7830 7835 7840 7845 7850 7855 7860 7865 7870 7875 7880 7885 7890 7895 7900 7905 7910 7915 7920 7925 7930 7935 7940 7945 7950 7955 7960 7965 7970 7975 7980 7985 7990 7995 8000 8005 8010 8015 8020 8025 8030 8035 8040 8045 8050 8055 8060 8065 8070 8075 8080 8085 8090 8095 8100 8105 8110 8115 8120 8125 8130 8135 8140 8145 8150 8155 8160 8165 8170 8175 8180 8185 8190 8195 8200 8205 8210 8215 8220 8225 8230 8235 8240 8245 8250 8255 8260 8265 8270 8275 8280 8285 8290 8295 8300 8305 8310 8315 8320 8325 8330 8335 8340 8345 8350 8355 8360 8365 8370 8375 8380 8385 8390 8395 8400 8405 8410 8415 8420 8425 8430 8435 8440 8445 8450 8455 8460 8465 8470 8475 8480 8485 8490 8495 8500 8505 8510 8515 8520 8525 8530 8535 8540 8545 8550 8555 8560 8565 8570 8575 8580 8585 8590 8595 8600 8605 8610 8615 8620 8625 8630 8635 8640 8645 8650 8655 8660 8665 8670 8675 8680 8685 8690 8695 8700 8705 8710 8715 8720 8725 8730 8735 8740 8745 8750 8755 8760 8765 8770 8775 8780 8785 8790 8795 8800 8805 8810 8815 8820 8825 8830 8835 8840 8845 8850 8855 8860 8865 8870 8875 8880 8885 8890 8895 8900 8905 8910 8915 8920 8925 8930 8935 8940 8945 8950 8955 8960 8965 8970 8975 8980 8985 8990 8995 9000 9005 9010 9015 9020 9025 9030 9035 9040 9045 9050 9055 9060 9065 9070 9075 9080 9085 9090 9095 9100 9105 9110 9115 9120 9125 9130 9135 9140 9145 9150 9155 9160 9165 9170 9175 9180 9185 9190 9195 9200 9205 9210 9215 9220 9225 9230 9235 9240 9245 9250 9255 9260 9265 9270 9275 9280 9285 9290 9295 9300 9305 9310 9315 9320 9325 9330 9335 9340 9345 9350 9355 9360 9365 9370 9375 9380 9385 9390 9395 9400 9405 9410 9415 9420 9425 9430 9435 9440 9445 9450 9455 9460 9465 9470 9475 9480 9485 9490 9495 9500 9505 9510 9515 9520 9525 9530 9535 9540 9545 9550 9555 9560 9565 9570 9575 9580 9585 9590 9595 9600 9605 9610 9615 9620 9625 9630 9635 9640 9645 9650 9655 9660 9665 9670 9675 9680 9685 9690 9695 9700 9705 9710 9715 9720 9725 9730 9735 9740 9745 9750 9755 9760 9765 9770 9775 9780 9785 9790 9795 9800 9805 9810 9815 9820 9825 9830 9835 9840 9845 9850 9855 9860 9865 9870 9875 9880 9885 9890 9895 9900 9905 9910 9915 9920 9925 9930 9935 9940 9945 9950 9955 9960 9965 9970 9975 9980 9985 9990 9995 9999

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48 49 50 51 52 53 54 55 56 57 58 59 60 61 62 63 64 65 66 67 68 69 70 71 72 73 74 75 76 77 78 79 80 81 82 83 84 85 86 87 88 89 90 91 92 93 94 95 96 97 98 99 100 101 102 103 104 105 106 107 108 109 110 111 112 113 114 115 116 117 118 119 120 121 122 123 124 125 126 127 128 129 130 131 132 133 134 135 136 137 138 139 140 141 142 143 144 145 146 147 148 149 150 151 152 153 154 155 156 157 158 159 160 161 162 163 164 165 166 167 168 169 170 171 172 173 174 175 176 177 178 179 180 181 182 183 184 185 186 187 188 189 190 191 192 193 194 195 196 197 198 199 200 201 202 203 204 205 206 207 208 209 210 211 212 213 214 215 216 217 218 219 220 221 222 223 224 225 226 227 228 229 230 231 232 233 234 235 236 237 238 239 240 241 242 243 244 245 246 247 248 249

2. Lyse cells by using pipette to up down and spin plates at 2000 rpm at 4°C for 5 min. The supernatant is ready to use.
3. Pipet 50 µl of standard, controls and samples to the wells.
4. Add 50 µl of MBP Assay Buffer to each well.
5. Incubate the well, shaking at 500-700 rpm on orbital microplate shaker for 2 hr at room temperature.
6. Add 100µl of the MBP Antibody-Biotin Conjugate to each well.
7. Incubate the well, shaking at 500-700 rpm on orbital microplate shaker for1 hr at room temperature.
- 10 8. Wash well 5 times with Wash Solution. Blot dry by inverting the plate on absorbent material.
9. Dilute the streptavidin-enzyme conjugate concentrate 1:50 with MBP Elisa Assay buffer. (must be diluted immediately prior to use in the assay).
10. Add 100 µl streptavidin-enzyme conjugate solutions to each well.
- 15 11. Incubate the well, shaking at 500-700 rpm on orbital microplate shaker for 30 min at room temperature.
12. Wash well 5 times with the Wash Solution. Blot dry by inverting the plate on absorbent material.
13. Add 100 µl of TMB Chromogen Solution to each well.
- 20 14. Incubate the well, shaking at 500-700 rpm on orbital microplate shaker for 10-20 min at room temperature. Avoid exposure to direct sunlight.
15. Add 100 µl of the Stopping Solution to each well.

Read the absorbance of the solution in the wells within 30 min, using a microplate reader set to 450 nM

25

In Vivo Proof of Concept Models

Focal Lesions: (used to assess whether compounds protect myelin integrity or accelerate/enhance the rate of remyelination.)

1. Rats 7 weeks of age are given free access to food and water and acclimatized for a minimum of 4 days before use in experiments.
- 30 2. Prior to surgery each animal is weighed. The rat is then anaesthetized with ketamine (100 mg/ml) in combination with xylazine (20 mg/ml) in a ratio of 1.8 : 1. The rats are injected with 0.15ml/180g body weight i.p. of the anaesthetic solution prior to the

surgical procedure. The animal is prepared for surgery using aseptic conditions in accordance with the IACUC guidelines. All surgical instruments will be autoclaved. The hair is clipped between the ears and this region will then be scrubbed with Betadine, flushed with sterile saline and finally wiped with a pre-packaged sterile alcohol swab.

- 5 3. For the surgical procedure, the rat is placed on its ventral surface in a small animal stereotaxic instrument designed to hold the head steady. The incisor bar is always set at -3.9 mm, since this has been shown to achieve a flat-skull position for SD rats.
4. An incision is made in the previously shaven skin overlying the skull between the ears.
- 10 5. A small area of bone (0.75mm in diameter) is drilled at the following coordinates AP – 1.8, ML –3.1 from lambda.
6. The bone is removed and rats are injected with 2 μ l ethidium bromide, lysolecithin, or SIN-1 into the right caudal cerebellar peduncle, DV –7.1 mm, over a 2 min period by means of a Hamilton μ l syringe and needle. Alternatively injections are made into the spinal cord, corpus callosum, or cortex.
- 15 7. The needle is left in position for the subsequent 2 min.
8. After withdrawal of the needle the incision is sutured.
9. Each rat receives an i.m. injection of 0.003mg buprenorphine into a hind leg.
10. The rat is placed in a warming cupboard until it regains consciousness. At which time 20 it is returned to its home cage. Do not allow more than 2 rats per cage, as they will pull each other's suture out.
11. Similar procedures are also done using mice.

Rat Experimental Allergic Encephalomyelitis (Rat EAE) Disease Model:

25 Experimental allergic encephalomyelitis (EAE) is a T-cell-mediated autoimmune disease of the nervous system that develops in susceptible animals following sensitization with either whole spinal cord homogenate or a component (myelin basic protein). The EAE rodent model is an appropriate tool for studying the inflammation of the brain and spinal cord observed in MS patients. In rodents, injection of whole spinal cord or spinal cord components 30 such as myelin basic protein induces an autoimmune response based on the activation of T-lymphocytes. Clinical disease typically becomes manifest around day 8-10 after inoculation, observed as a broad spectrum of behavioral anomalies ranging from mild gait disturbances and tail atony to complete paralysis and death. Weight loss typically occurs. In animals that

survive, spontaneous recovery occurs, accompanied by variable recovery of most motor function. Depending on the species, allergen, and methodology used, animals tested by the EAE model may experience a single (acute EAE) or several (chronic relapsing EAE) attacks. Several treatment paradigms may be used: the drug or treatment of choice may be 5 administered before immunization, during the nonsymptomatic period or during the clinical disease.

Animals:

Female Lewis rats, 160-220g (Charles River)

10 Antigen:

Whole Guinea Pig spinal cord (Harlan Biosciences).

Complete Freund's adjuvant H37 Ra [1mg/ml Mycobacterium Tuberculosis H37 Ra] (Difco).

Additional antigen:

15 Mycobacterium Tuberculosis (Difco).

Bordetella Pertussis [Heat Killed] (Difco).

Antigen preparation: (for approximately 720 animals):

1. Weigh 5 grams of frozen guinea pig spinal cord.
2. Add 5g spinal cord to 5ml 0.9% saline (1g/ml) in a round bottom centrifuge tube
- 20 3. Homogenize on ice with the Tissue-tech until the tissue is completely disrupted (approximately 5 minutes).
4. Add 10 ml Complete Freund's adjuvant H37 Ra supplemented with 200 mg Mycobacterium Tuberculosis (20 mg / ml Complete Freund's adjuvant H37 Ra).
5. Extract homogenate / adjuvant from tube by sucking it into a 10 ml syringe fitted with an 25 18 gauge emulsifying needle.
6. Emulsify between two 30 ml glass syringes until it becomes difficult to continue passing the material through the needle. (Approximately 5 minutes {there must be no separation between the oil phase and the aqueous phase}).
7. Use immediately or keep on ice until needed (not more than 30 min) (do not freeze).

30

Protocol

1. Female Lewis rats (Charles River) are given free access to food and water and should be acclimated a minimum of 3 days before use in experiments.

2. Rats weighing 160 and 220 grams are initially induced with 5% isoflurane (Aerrane, Fort Dodge), 30% O₂, 70% N₂O for 2-5 minutes.
3. The rat is then placed onto a circulating water heating blanket (Gaymar) (dorsal surface up) and into the nose cone for spontaneous respiration of anesthetic gases. The isoflurane is reduced to 2%.
- 5 4. Two subcutaneous injections (0.1 ml each) of either antigen or normal saline are made into ventral surface of the hind paws.
5. The animals are removed from the nose cone, weighed and numbered.
6. The rats are allowed to awake from anesthesia and are placed into individual cages.
- 10 7. The animals are observed daily for signs of EAE induction (see criteria below)

STAGE:0 NORMAL

STAGE 1 Abnormal gate and tail atony

STAGE 2 Mild but definite weakness of one or both hind legs

15 STAGE: 3 Severe weakness of one or both hind legs or mild ataxia

STAGE: 4 Severe paraparesis and minimal hind leg movement

STAGE: 5 No hind leg movement and paraplegia

STAGE: 6 Moribund state with no spontaneous movement and impaired respiration.

Increasing degree of front leg involvement and urinary and fecal incontinence
20 may also occur

STAGE:7 DEATH

Treatment is begun on day 10 after immunization. Since the disease symptoms in this model typically appear 10-11 days after inoculation, this time point may be considered to represent the initial phase of an acute episode of MS. It is judged that this delay of the start of treatment mimics the clinical situation more closely than the traditionally used protocols where drugs are administered at the time of, or even before, inoculation (Teitelbaum D. et al., Proc Natl Acad Sci USA 1999; 96: 3842-3847 and Brod S. A., et al., Ann Neurol 2000; 47: 25 127-131).

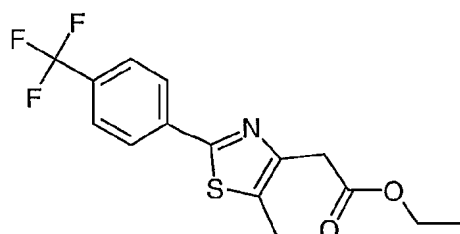
30 This invention is further illustrated by the following examples of compounds used herein which are provided for illustration purposes and in no way limit the scope of the present invention.

Synthetic ExamplesGeneral

Commercial reagents and solvents were used as received. ^1H NMR spectra were recorded on a Varian MercuryPlus-300 (300 MHz) or Varian Unity Inova (400 MHz) spectrometer as indicated. Proton chemical shifts are reported in δ ppm relative to internal tetramethylsilane (0.0 ppm). MS (LC-MS) data is obtained using a Micromass LCT time of flight mass spectrometer with electrospray ionization and 5 min data acquisition time for m/z 100 to 1000. LC (LC-MS) is performed using a Hypersil C18 column (4.6x50mm, 3.5 μ) with mobile phase of 0.1 % TFA in H_2O (A) and 0.1% TFA in ACN (B) and a gradient of 5% to 100% B over 3 min followed by 2 min at 100% B. Alternatively, a Platform LC-MS with electrospray source may be used with a HP1100 LC system running at 2.0 ml/min, 200 $\mu\text{L}/\text{min}$ split to the ESI source with inline HP1100 DAD detection and SEDEX ELS detection. A Luna C18(2) column (30x4.6mm 3 μ) is used with a gradient of 5% to 95% B over 4.5 min with mobile phase of 0.1% formic acid in H_2O and 0.1% formic acid in ACN (B). HPLC purification is performed on a Varian ProStar system using a reversed-phase C18 column with a linear gradient of ACN / H_2O containing 0.1% trifluoroacetic acid. Microwave syntheses were performed using a Personal Chemistry Smithcreator microwave reaction system using 2 or 5 mL reactor vessels.

20

Example 1

25 Intermediate:[5-Methyl-2-(4-trifluoromethyl-phenyl)-thiazol-4-yl]-acetic acid ethyl ester

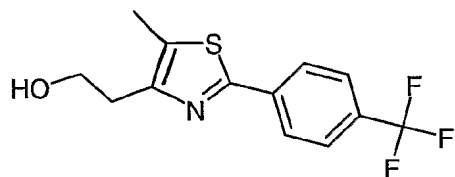
To a solution of 4-trifluoromethyl-benzene-thioamide (1.845g, 9mmol) in ethanol (15 mL, 200 proof) add ethyl-4-bromo-3-oxo-pentanoate (2.07g, 9 mmol). Seal this solution warm the solution to 170°C in a Personal Chemistry™ microwave oven and stir at this temperature for 20 min. Cool the resulting solution to room temperature, concentrate under

reduced pressure and purify the residue by flash chromatography (elute with 30% ethyl acetate / 10% dichloromethane in heptane) and obtain the title compound as a white solid (1.4g).

MS (ESI) m/z 330 (M+H); H1 NMR ($CDCl_3$) δ 1.87 (bs, 1H), 2.49 (s, 3H), 4.86 (s, 2H), 7.67 (d, J = 8Hz, 2H), 8.02 (d, J = 8Hz, 2H).

5

Example 2



Intermediate: 4-(2-hydroxyethyl)-5-methyl-2-(4-trifluoromethyl-phenyl)thiazole

10

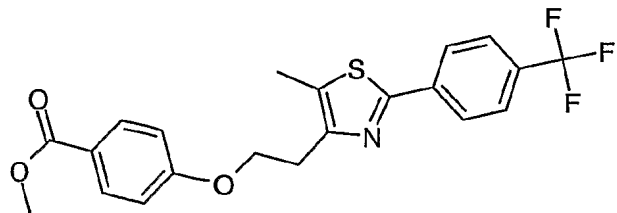
Cool (0°C) a solution of lithium aluminum hydride (5.3 mL, 1M in THF) and add a solution of [5-methyl-2-(4-trifluoromethyl-phenyl)-thiazol-4-yl]-acetic acid ethyl ester (Example 1, 1.4g, 4.25 mmol) in THF (15 mL). On complete addition, remove the cold bath and stir for 2 hrs. Cool this solution to 5°C, and then add water (0.2 mL), dropwise, followed by NaOH solution (0.2 mL, 5M in water) and water (0.2 mL). Dilute the resulting mixture with ethyl acetate and then filter through a pad of celite. Wash the solids with dichloromethane and then concentrate the combined filtrates under reduced pressure. Purify the residue by flash chromatography (elute with 30% ethyl acetate 40% dichloromethane in heptane) to give the title compound as a yellow solid (0.879 g) Use the compound of Example 1 as the starting material to obtain the title compound.

15

MS (ESI) m/z 288 (M+H); H1 NMR ($CDCl_3$) δ 2.44 (s, 3H), 2.91 (t, J = 7Hz, 2H), 3.62 (t, J = 6Hz, 1H), 4.01 (dt, J = 7, 6Hz, 2H), 7.66 (d, J = 8Hz, 2H), 7.96 (d, J = 8Hz, 2H).

20

Example 3



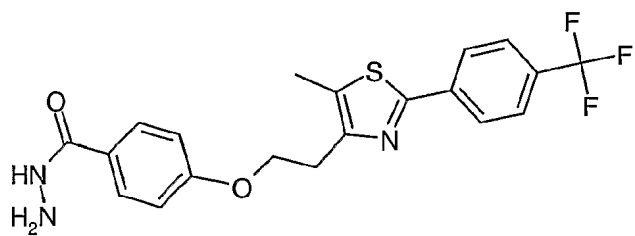
5 Intermediate: 4-[5-Methyl-2-(4-trifluoromethyl-phenyl)-thiazol-4-ylethoxy]-benzoic acid methyl ester.

To a solution of 4-(2-hydroxy-ethyl)-5-methyl-2-(4-trifluoromethyl-phenyl)thiazole (Example 3, 288mg, 1.0 mmol) in THF (3 mL) add 4-hydroxy-benzoic acid methyl ester (167 mg, 1.1 mmol) followed by triphenylphosphine (288mg, 1.1 mmol). To this solution, add, dropwise, diethyl azodicarboxylate (174 μ L, 1.1 mmol). On complete addition, stir the resulting red solution for 20 min. concentrate under reduced pressure and purify the residue by flash chromatography (elute with 15% ethyl acetate / 15% dichloromethane in heptane) to give the title compound as a white solid. (410 mg).

10 MS (ESI) m/z 422 (M+H); H1 NMR (DMSO) δ 2.51 (s, 3H), 3.19 (t, J = 7Hz, 2H), 3.80 (s, 3H), 4.40 (t, J = 7Hz, 2H), 7.05 (d, J = 9Hz, 2H), 7.83 (d, J = 8Hz, 2H), 7.88 (d, J = 8Hz, 2H) 8.05 (d, J = 8Hz, 2H).

Example 4

20



Intermediate: 4-[2-[5-Methyl-2-(4-trifluoromethyl-phenyl)-thiazol-4-yl]-ethoxy]-benzoic acid hydrazide

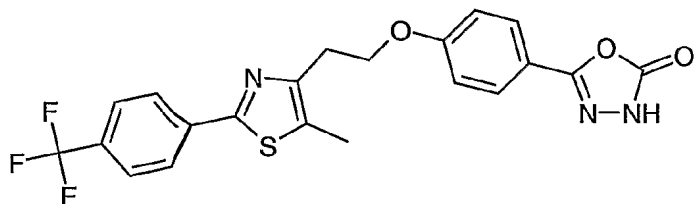
25

To a suspension of 4-[5-Methyl-2-(4-trifluoromethyl-phenyl)-thiazol-4-ylethoxy]-benzoic acid methyl ester (Example 4, 410 mg, 1 mmol) in methanol (3mL) add anhydrous hydrazine (0.32 ml, 10 mmol). Warm the resulting mixture to 60°C and stir at this temperature for 66 hrs. Cool the resulting solution to room temperature, and add 3 drops of water. Filter the precipitate wash with ether to give the title compound (279 mg).

MS (ESI) *m/z* 422 (M+H); H1 NMR (DMSO) δ 2.51 (s, 3H), 3.17 (t, *J* = 7Hz, 2H), 4.36 (t, *J* = 7Hz, 2H), 4.38 (bs, 2H), 6.98 (d, *J* = 9Hz, 2H), 7.77 (d, *J* = 9Hz, 2H), 7.83 (d, *J* = 8Hz, 2H) 8.06 (d, *J* = 8Hz, 2H) 9.58 (bs, 1H).

10

Example 5



15

5-(4-{2-[5-Methyl-2-(4-trifluoromethyl-phenyl)-thiazol-4-yl]-ethoxy}-phenyl)-3H-[1,3,4]oxadiazol-2-one.

20

25

To a suspension of : 4-{2-[5-Methyl-2-(4-trifluoromethyl-phenyl)-thiazol-4-yl]-ethoxy}-benzoic acid hydrazide (Example 4, 276 mg, 0.65 mmol) in dichloromethane (4 mL) add pyridine (104 μL, 1.3 mmol) followed by phenylchloroformate (0.88 μL, 0.71 mmol). Stir the resulting mixture at room temperature until all the starting material is consumed (by TLC analysis). Dilute the mixture with ethyl acetate wash with water then brine dry over MgSO4 and concentrate under reduced pressure. Take the residue up in acetonitrile (5 mL). To this mixture, add DBU (106 μL, 0.71 mmol). Seal the resulting solution; warm it to 170°C in a Personal Chemistry™ microwave oven and stir at this temperature for 120 min. Cool the reaction to room temperature, dilute with ethyl acetate, wash with 1 M HCl solution (or saturated NaH2PO4 solution) dry over MgSO4 and concentrate. Triturate the resulting residue with dichloromethane several times to give the title compound as a tan solid (137 mg). (recrystallized from ethyl acetate in a sealed tube at 140°C).

MS (ESI) m/z 448 (M+H); ^1H NMR (DMSO) δ 2.51 (s, 3H), 3.19 (t, J = 7Hz, 2H), 4.40 (t, J = 7Hz, 2H), 7.10 (d, J = 8Hz, 2H), 7.70 (d, J = 8Hz, 2H), 7.83 (d, J = 8Hz, 2H) 8.05 (d, J = 8Hz, 2H) 12.41 (bs, 1H).

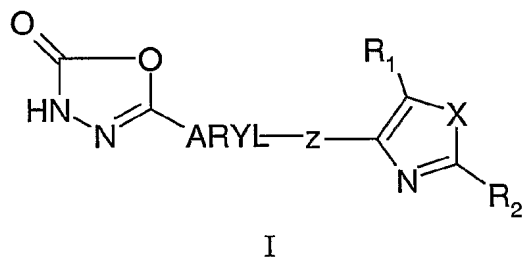
5

10

CLAIMS

What is claimed is:

5 1. A compound of formula I:



10

wherein

ARYL is phenyl or pyridinyl, wherein said phenyl or pyridinyl is optionally substituted with one or more substituents selected from the group consisting of halogen, C₁₋₆alkyl, C₂₋₆alkenyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl; C₁₋₆alkylthio, hydroxy, hydroxyC₁₋₆alkyl, C₁₋₆acyloxy, nitro, cyano, C₁₋₆alkylsulfonyl, amino, C₁₋₆alkylamino and C₁₋₆alkoxycarbonyl;

15

Z is -O(CH₂)_n-, -SO₂(CH₂)_n-, -(CH₂)_n-Y-(CH₂)_n-, -(CH₂)_n-CO-, -O(CH₂)_n-CO-, or -(CH₂)_n-Y-(CH₂)_n-CO- wherein Y is NR₃, O or S and R₃ is selected from the group consisting of H, C₁₋₆alkyl C₃₋₈cycloalkyl, C₁₋₆alkylC₃₋₈cycloalkyl and benzyl and n is independently an integer from 1 to 5;

20

X is NR₃, O or S wherein R₃ is as defined above;

R₁ is H, halogen, C₁₋₆alkyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl; hydroxyC₁₋₆alkyl, nitro, cyano, and C₁₋₆alkylamino; and

25

R₂ is substituted or unsubstituted phenyl, pyridinyl or thienyl wherein the substituents are selected from the group consisting of halogen, C₁₋₆alkyl, C₂₋₆alkenyl, C₁₋₆alkoxy, C₁₋₆perfluoroalkyl, C₁₋₆alkylthio, hydroxy, hydroxyC₁₋₆alkyl, C₁₋₄acyloxy, nitro, cyano, C₁₋₆alkylsulfonyl, amino, C₁₋₆alkylamino and C₁₋₆alkoxycarbonyl;

with the proviso that when Z is -O(CH₂)_n- or -SO₂(CH₂)_n-, and ARYL is phenyl then R₂ is other than phenyl;

30

or a stereoisomer, a tautomer or a solvate thereof or a pharmaceutically acceptable salt thereof.

2. The compound according to claim 1 wherein ARYL is phenyl; and
X is O or S

3. The compound according to claim 2 wherein X is O.

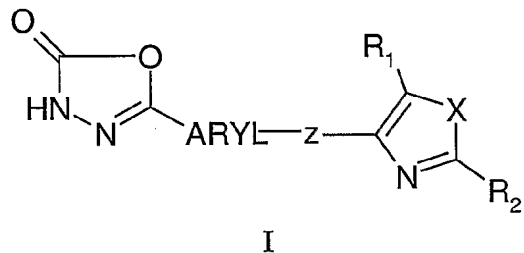
5

4. A compound which is 5-(4-{2-[5-methyl-2-(4-trifluoromethyl-phenyl)-thiazol-4-yl]-ethoxy}-phenyl)-3H-[1,3,4]oxadiazol-2-one.

5. A pharmaceutical composition comprising an effective amount of a compound according to
10 claim 1 and a pharmaceutical acceptable carrier.

6. A method of treating a disease in a mammal wherein the disease is capable of being,
modulated by PPARdelta ligand binding activity, which comprises administering to said
mammal having said disease a therapeutically effective amount of a compound of formula I:

15



wherein

20 ARYL is phenyl or pyridinyl, wherein said phenyl or pyridinyl is optionally substituted with
one or more substituents selected from the group consisting of halogen, C₁-6alkyl, C₂-
6alkenyl, C₁-6alkoxy, C₁-6perfluoroalkyl; C₁-6alkylthio, hydroxy, hydroxyC₁-6alkyl, C₁-
4acyloxy, nitro, cyano, C₁-6alkylsulfonyl, amino, C₁-6alkylamino and C₁-
6alkoxycarbonyl;

25 Z is -O(CH₂)_n-, -SO₂(CH₂)_n-, -(CH₂)_n-Y-(CH₂)_n-, -(CH₂)_n-CO-, -O(CH₂)_n-CO-
or -(CH₂)_n-Y-(CH₂)_n-CO- wherein Y is NR₃, O or S and R₃ is selected from the group
consisting of H, C₁-6alkyl C₃-8cycloalkyl, C₁-6alkylC₃-8cycloalkyl and benzyl and n is
independently an integer from 1 to 5;

X is NR₃, O or S wherein R₃ is as defined above;

30 R₁ is H, halogen, C₁-6alkyl, C₁-6alkoxy, C₁-6perfluoroalkyl; hydroxyC₁-6alkyl, nitro, cyano,
and C₁-6alkylamino; and

R_2 is substituted or unsubstituted phenyl, pyridinyl or thienyl wherein the substituents are selected from the group consisting of halogen, C_{1-6} alkyl, C_{2-6} alkenyl, C_{1-6} alkoxy, C_{1-6} perfluoroalkyl, C_{1-6} alkylthio, hydroxy, hydroxy C_{1-6} alkyl, C_{1-4} acyloxy, nitro, cyano, C_{1-6} alkylsulfonyl, amino, C_{1-6} alkylamino and C_{1-6} alkoxycarbonyl; or a stereoisomer, a tautomer or a solvate thereof or a pharmaceutically acceptable salt thereof.

5 7. The method according to claim 6 wherein ARYL is phenyl.

8. The method according to claim 6 wherein ARYL is phenyl; and

10 R_2 is phenyl.

9. The method according to claim 6 wherein ARYL is phenyl;

Z is $-O(CH_2)_n-$; and

R_2 is phenyl.

15

10. The method according to claim 6 wherein ARYL is phenyl;

Z is $-O(CH_2)_n-$;

X is O or S; and

R_2 is phenyl.

20

11. The method according to claim 6 wherein ARYL is phenyl;

Z is $-O(CH_2)_n-$;

X is O or S; and

R_1 is C_{1-6} alkyl; and

25 R_2 is phenyl.

12. The method according to claim 11 wherein X is O.

13. The method according to claim 12 wherein X is S.

30

14. The method according to claim 6 wherein said disease is a demyelinating disease selected from the group consisting of multiple sclerosis, Charcot-Marie-Tooth disease, Pelizaeus-Merzbacher disease, encephalomyelitis, neuromyelitis optica,

adrenoleukodystrophy, Guillain-Barre syndrome and disorders in which myelin forming glial cells are damaged including spinal cord injuries, neuropathies and nerve injury.

15. The method according to claim 14 wherein the demyelinating disease is multiple sclerosis.

5

16. The method according to claim 6 wherein said disease is selected from the group consisting of obesity, hypertriglyceridemia, hyperlipidemia, hypoalphalipoproteinemia, hypercholesterolemia, dyslipidemia, Syndrome X, Type II diabetes mellitus and complications thereof selected from the group consisting of neuropathy, nephropathy, retinopathy and cataracts, hyperinsulinemia, impaired glucose tolerance, insulin resistance, atherosclerosis, hypertension, coronary heart disease, peripheral vascular disease or congestive heart failure.

10

15

20

25