Title: NOVEL PROLYLCARBOXYPEPTIDASE INHIBITORS

Abstract: Compounds of structural formula I are inhibitors of prolylcarboxypeptidase (PrCP). The compounds of the present invention are useful for the prevention and treatment of conditions related to the enzymatic activity of PrCP such as abnormal metabolism, including obesity; diabetes; metabolic syndrome; obesity related disorders; and diabetes related disorders.
TITLE OF THE INVENTION
NOVEL PROLYLCARBOXYPEPTIDASE INHIBITORS

FIELD OF THE INVENTION

The present invention relates to compounds which are inhibitors of the prolylcarboxy-
peptidase (PrCP) enzyme and the use of such compounds to control, prevent and/or treat
conditions or diseases mediated by prolylcarboxypeptidase activity. The compounds of the
present invention are useful for the control, prevention and treatment of conditions and diseases
related to abnormal metabolism, including obesity; diabetes; metabolic syndrome, obesity related
disorders and diabetes related disorders.

BACKGROUND OF THE INVENTION

Obesity, which can be defined as a body weight more than 20% above the ideal body
weight, is a major health concern in Western societies. It is estimated that about 97 million
adults in the United States are overweight or obese. Obesity is the result of a positive energy
balance, as a consequence of increased ratio of caloric intake to energy expenditure.
Epidemiological studies have shown that increasing degrees of overweight and obesity are
important predictors of decreased life expectancy. Obesity causes or exacerbates many health
problems, both independently and in association with other diseases. The medical problems
associated with obesity, which can be serious and life-threatening, include hypertension; type 2
diabetes mellitus; elevated plasma insulin concentrations; insulin resistance; dyslipidemias;
hyperlipidemia; endometrial, breast, prostate and colon cancers; osteoarthritis; respiratory
complications, such as obstructive sleep apnea; cholelithiasis; gallstones; arteriosclerosis; heart
disease; abnormal heart rhythms; and heart arrhythmias (Kopelman, P.G., Nature 404, 635-643
(2000)). Obesity is further associated with premature death and with a significant increase in
mortality and morbidity from stroke, myocardial infarction, congestive heart failure, coronary
heart disease, and sudden death.

Pro-opiomelanocortin (POMC) derived peptides are known to affect food intake. Several
lines of evidence support the notion that the G-protein coupled receptors (GPCRs) of the
melanocortin receptor (MC-R) family, several of which are expressed in the brain, are the targets
of POMC derived peptides involved in the control of food intake and metabolism. A specific
single MC-R that may be targeted for the control of obesity has not yet been identified, although
evidence has been presented that MC-4R signalling is important in mediating feed behavior (S.Q.
Giraudo et al., "Feeding effects of hypothalamic injection of melanocortin-4 receptor ligands,"
Brain Research, 80: 302-306 (1998)).

The prohormone pro-opiomelanocortin (POMC) plays a critical role in the regulations
of energy metabolism, and is processed by proteases to produce several peptide hormones,
including alpha-melanocyte-stimulating hormone (α-MSH or α-MSH1-13). α-MSH is a major
regulator of feeding and body weight homeostasis. Studies have shown that $\alpha$-MSH$_{1-13}$ is a critical anorexigenic neuromodulator found in the hypothalamus, which inhibits food intake by binding target neurons expressing melanocortin receptors 3 and 4 (MC3R and MC4R) (see Vaisse et al., J Clin Invest, 106, 253-62 (2000); and Williams et al., Am J Physiol Regul Integr Comp Physiol, 289:R2-R3 (2005). MC-3R is expressed in the brain, gut, and placenta and may be involved in the control of food intake and thermogenesis. MC-4R is uniquely expressed in the brain, and its inactivation was shown to cause obesity (A. Kask, et al, "Selective antagonist for the melanocortin-4 receptor (HS014) increases food intake in free-feeding rats," Biochem Biophys Res Commun, 245: 90-93 (1998)).

The enzyme prolylcarboxypeptidase (PRCP, Lysosomal Pro-X carboxypeptidase, angiotensinase C) is a serine protease that cleaves small, biologically active peptides at carboxyl termini linked to a penultimate proline group, a -MSH is a substrate of PRCP due to its C-terminal amino acid sequence, Pro-Val. Recent studies have shown that PRCP initiates the degradation of $\alpha$-MSH$_{1-13}$ into inactive extracellular a -MSH$_{1-12}$, which is effective in reducing food intake and in regulating neuronal functions via melanocortin receptors. In overnight fasted animals, 2.5 ug of $\alpha$-MSH$_{1-13}$ induced a 40% reduction in food intake relative to control animals, however, overnight fasted animals treated with 2.5 ug of $\alpha$-MSH$_{1-12}$ did not significantly affect food intake compared to the controls. (Wallingford et al., J Clinical Investigation, Vol. 119, No. 8, August 2009).

Further it has been shown that PRCP inhibition by small molecule protease inhibitors administered peripherally or centrally decreased food intake in wild type and genetically obese animals. Specifically, both the intracerebroventricular to rats and systemic administration to obese, leptin deficient mice of t-butyl carbamate-prolyl prolinal (BPP), which is an inhibitor of PRCP, resulted in a suppression of overnight food intake (Wallingford et al., J Clinical Investigation, Vol. 119, No. 8, August 2009).

A recent study also showed that PrCP null mice had elevated hypothalamic levels of $\alpha$-MSH$_{1-13}$ and were leaner compared with wild-type controls when fed regular chow, and were also resistant to high fat diet induced obesity. Specifically, on a high fat diet, PrCP gt/gt mice also showed a significant reduction in body weight and a reduction in food intake (Wallingford et al., J Clinical Investigation, Vol 119, No. 8, August 2009).

These studies suggest that PRCP inhibitors influence food intake and weight maintenance via melanocortin receptors and the control of active $\alpha$-MSH$_{1-13}$ levels, and that targeting PRCP activity with central or peripheral administration of inhibitors can reduce food intake.

WO 2005/115446 discloses the role of prolylcarboxypeptidase inhibitors in weight control, control of body fat and food intake; and specific prolylcarboxypeptidase inhibitors, including t-butyl carbamate (BOC)-prolyl prolinal (BPP), N-benzoylcarbonyl-prolyl-proinal, diisopropyl fluorphosphates, PMSF, antipain, leupeptin, corn trypsin and mercuric chloride,
useful to treat obesity and obesity related disorders. WO 2005/1 15446 also discloses the association of PRCP with hypertension, dyslipidemia, diabetes, stroke, gallbladder disease, cardiovascular disease, osteoarthritis, rheumatoid arthritis, hypercholesterolemia, angina, atherosclerosis, sleep apnea, respiratory problems, and cancer.

US 2008-01 08080 discloses the utility of small molecule compounds with activity against the gene products encoded by PRCP for use in treating obesity.

WO 2007/140896 discloses the association of human PRCP with cardiovascular diseases, hematological diseases, neurological diseases and cancer based upon tissue distribution of PrCP.

The prolylcarboxypeptidase (PRCP) enzyme is disclosed in EP 1498424 and WO 2004/072265.

The present invention is concerned with novel heteroarylpiperidine compounds as inhibitors of prolylcarboxypeptidase which are useful in the treatment and/or prevention of various conditions and diseases mediated by prolylcarboxypeptidase activity including, but not limited to, abnormal metabolism, obesity, diabetes, metabolic syndrome, obesity and diabetes related disorders, such as hypertension, dyslipidemia, stroke, gallbladder disease, cardiovascular disease, osteoarthritis, rheumatoid arthritis, hypercholesterolemia, stable angina, unstable angina, artherosclerosis, sleep apnea, respiratory problems, cancer, stroke, hematological diseases and neurological diseases.

Weight loss drugs that are currently used to treat obesity have limited efficacy and significant side effects. Studies of the weight loss medications orlistat (Davidson, M.H. et al. (1999) JAMA 281:235-42), dexfenfluramine (Guy Grand, B. et al. (1989) Lancet 2:1 142-5), sibutramine (Bray, G. A. et al. (1999) Obes. Res. &:189-98) and phentermine (Douglas, A. et al. (1983) hyl. J. Obes. 7:591-5) have demonstrated a limited weight loss of about 5%-10% of body weight for drug compared to placebo. The side effects of these drugs and anti-obesity agents further limit their use. Dexfenfluramine was withdrawn from the market because of suspected heart valvulopathy; orlistat is limited by gastrointestinal side effects; the use of topiramate is limited by central nervous system effects; and the use of sibutramine is limited by its cardiovascular side effects which have led to reports of deaths and its withdrawal from the market in Italy. Obese patients generally respond well to surgical interventions that modify the gastrointestinal tract and limit food intake. However, one out of fifty bariatric surgery patients dies within the first 30 days post surgery, and 4.6% of bariatric surgery patients die within the first year (J Amer. Med. Assoc., 2005, 294, 1903). Another study indicated that 33% of patients that undergo bariatric surgery have complications that require re-hospitalization within the first 6 months post operation (Medical Care, 2009, 47, 531).

There is a need for a weight loss treatment with enhanced efficacy, increased safety, and fewer undesirable side effects. The instant invention addresses this problem by providing prolylcarboxypeptidase inhibitors useful in the treatment and prevention of obesity, diabetes, and related disorders.
SUMMARY OF THE INVENTION

The present invention relates to compounds of structural formulas I-1 and I-2:

Compounds of formula I are inhibitors of prolylcarboxypeptidase (PrCP) and as such are useful in the treatment, control or prevention of diseases, disorders or conditions responsive to the modulation of the prolylcarboxypeptidase (PrCP) enzyme. In particular, the compounds of formula I act as inhibitors of the prolylcarboxypeptidase (PrCP) enzyme useful in the treatment, control or prevention of diseases, disorders or conditions responsive to the inhibition of prolylcarboxypeptidase (PrCP), such as eating disorders due to excessive food intake, and the resulting obesity and complications associated therewith, including diabetes, obesity related disorders and diabetes related disorders.

The present invention also relates to pharmaceutical compositions comprising the compounds of the present invention and a pharmaceutically acceptable carrier.

The present invention also relates to methods for the treatment, control, or prevention of disorders, diseases, or conditions responsive to inhibition of prolylcarboxypeptidase in a subject in need thereof by administering the compounds and pharmaceutical compositions of the present invention.

The present invention also relates to methods for the treatment, control, or prevention of obesity, Type 2 diabetes, metabolic syndrome, obesity related disorders and diabetes related disorders by administering the compounds and pharmaceutical compositions of the present invention.

DETAILED DESCRIPTION OF THE INVENTION

The present invention is concerned with compounds useful as inhibitors of prolylcarboxypeptidase. Compounds of the present invention are described by structural formulas 1-1 and 1-2:
or a pharmaceutically acceptable salt thereof; wherein

5 "a" is a single bond or absent;

X is independently selected from N and CR.5;

each R is independently selected from the group consisting of:

10 (1) hydrogen,
(2) -C1-6 alkyl,
(3) -C1-6 alkoxy,
(4) -(CH2)mC02Ci-6 alkyl,
(5) -(CH2)m COCi-6 alkyl,
(6) -(CH2)m-C(0)-(CH2)u-halogen,
(7) -(CH2)m-C(0)-(CH2)uC02H,
(8) -(CH2)m-C(0)-(CH2)uC02Ci-6 alkyl,
(9) -(CH2)m-C(0)-(CH2)pN(Rg)2,
(10) -(CH2)m-C(0)-(CH2)p-C3-6cycloalkyl,
(11) -(CH2)m-C(0)-(CH2)p-C2-6cycloalkyl,
(12) -(CH2)m-C(0)-(CH2)p-aryl,
(13) -(CH2)m-C(0)-(CH2)p-heteroaryl,
(14) -(CH2)m-CH(OH)-(CH2)p-C3-6cycloalkyl,
(15) -(CH2)m-CH(OH)-(CH2)p-C2-6cycloalkyl,
(16) -(CH2)m-CH(OH)-(CH2)p-aryl,
(17) -(CH2)m-CH(OH)-(CH2)p-heteroaryl,
(18) -(CH2)m-NRg-Ci-6 alkyl,
(19) -(CH2)m-O-C1-6 alkyl,
(20) -(CH2)m-S-Ci-6 alkyl,
(21) -(CH2)m-S02-Ci-6 alkyl,
(22) -(CH2)m-S02-(CH2)q-aryl,
(23) -(CH2)m C3-7cycloalkyl,
(24) -(CH2)m C2-6cycloheteroalkyl,
(25) -(CH2)m aryl,
(26) -(CH2)m-biphenyl, and
(27) -(CH2)m heteroaryl,
wherein each CH2, alkyl, alkoxy, cycloalkyl, cycloheteroalkyl, aryl, biphenyl and heteroaryl is unsubstituted or substituted with one to three groups independently selected from R:\n
each R2 is independently selected from the group consisting of:
   (1) hydrogen, and
   (2) -Ci-6 alkyl,
wherein each alkyl is unsubstituted or substituted with one to three substituents selected from Rb;

each R3 is independently selected from the group consisting of:
   (1) -Ci-6 alkyl,
   (2) -(CH2)\, C3-7cycloalkyl,
   (3) -(CH2)n C2-6cycloheteroalkyl,
   (4) -(CH2)n aryl, and
   (5) -(CH2)n heteroaryl,
wherein each Cl\, alkyl, cycloalkyl, cycloheteroalkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from Rc;

each R4 is independently selected from the group consisting of:
   (1) -Ci-6 alkyl,
   (2) -C1-6 alkoxy,
   (3) -CF3,
   (4) -(CH2)s C3-7cycloalkyl,
   (5) -(CH2)s C2-6cydoheteroalkyl,
   (6) -(CH2)s aryl, and
   (7) -(CH2)s heteroaryl,
wherein each Cl\, alkyl, alkoxy, cycloalkyl, cycloheteroalkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from Rd;

each R5 is independently selected from the group consisting of:
   (1) hydrogen, and
   (2) -Ci-6 alkyl,
wherein each alkyl is unsubstituted or substituted with one to three substituents selected from Re;

each Ra is independently selected from the group consisting of:
(1) hydrogen,
(2) -OH,
(3) oxo,
(4) -C₁-₆ alkyl,
(5) -OC₁-₆ alkyl,
(6) halogen,
(7) -CF₃,
(8) -OCF₃,
(9) -CN,
(10) -C(O)C₁-₆ alkyl,
(11) -CO₂H,
(12) -CO₂C₁-₆ alkyl,
(13) -(CH₂)ₚ C₃-₇cycloalkyl,
(14) -(CH₂)ₚ C₂-₆cycloalkyl,
(15) -(CH₂)ₚ aryl,
(16) -CH(phenyl)₂,
(17) -(CH₂)ₚ heteroaryl,
(18) -SO₂C₁-₆ alkyl,
(19) -SO₂N(R')₂;
(20) -SO₂-C₃-₇cycloalkyl,
(21) -SO₂-C₂-₆cycloalkyl,
(22) -SO₂-aryl-, and
(23) -SO₂-heteroaryl;

25 each Rₕ is independently selected from the group consisting of:
   (1) halogen,
   (2) -C₁-₆ alkyl,
   (3) -OC₁-₆ alkyl,
   (4) -CO₂H, and
   (5) -CO₂C₁-₆ alkyl,

30 wherein alkyl can be substituted with one to three fluorines;

each Rₖ is independently selected from the group consisting of:
   (1) hydrogen,
   (2) -OH,
   (3) oxo,
   (4) -CI-₆ alkyl,
   (5) -OC₁-₆ alkyl,
(6) halogen,
(7) -CF3,
(8) -OCF3,
(9) -CN,
(10) -CO2H, and
(11) -C2H5 alkyl;

each Rd is independently selected from the group consisting of:

(1) hydrogen,
(2) -OH,
(3) oxo,
(4) -C1 alkyl,
(5) -OC1 alkyl,
(6) halogen,
(7) -CF3,
(8) -OCF3,
(9) -CN,
(10) -CO2H, and
(11) -C2Cl alkyl;

each Re is independently selected from the group consisting of:

(1) halogen,
(2) -C1 alkyl,
(3) -OC1 alkyl,
(4) -CO2H, and
(5) -C2Cl alkyl,

wherein alkyl can be substituted with one to three fluorines;

each Rf is independently selected from the group consisting of:

(1) hydrogen, and
(2) -Cl alkyl;

each Rg is independently selected from the group consisting of:

(1) hydrogen, and
(2) -Cl alkyl;

m is selected from 0, 1, 2 and 3;
n is selected from 0, 1, 2, and 3.
p is selected from 0, 1, 2 and 3;
q is selected from 0, 1, 2 and 3;
r is selected from 0, 1, 2, 3 and 4;
s is selected from 0, 1, 2 and 3;
t is selected from 0, 1, 2 and 3; and
u is selected from 1, 2 and 3.

In one embodiment of the present invention, "a" is a single bond or absent. In another
embodiment of the present invention, "a" is a single bond. In another embodiment of the
present invention, "a" is absent.

In another embodiment of the present invention, X is independently selected from N and
CR5. In another embodiment of the present invention, X is independently selected from N and
CH. In another embodiment of the present invention, X is N. In another embodiment of the
present invention, X is CR5. In another embodiment of the present invention, X is CH.

In another embodiment of the present invention, each R¹ is independently selected from
the group consisting of: hydrogen, -C1-6 alkyl, -C1-6 alkoxy, -(CH2)mC0 2C1-6 alkyl, -(CH2)mCOCi6 alkyl, -(CH2)mC-C(0)-(CH2)u-halogen, -(CH2)mrC(0)-(CH2)uC02H, -(CH2)m-C(0)~(CH2)uC02 Ci-6 alkyl, -(CH2)m-C(0)-(CH2)pN(Rg)2, -(CH2)mrC(0)-(CH2)p-C3-6cycloalkyl, -(CH2)m-C(0)-(CH2)p-C2-6cycloalkylkyl, -(CH2)m-C(0)-(CH2)p-aryl, -(CH2)m-C(0)-(CH2)p-heteroaryl, -(CH2)m-CH(OH)-(CH2)p-C3-6cycloalkyl, -(CH2)m-CH(OH)-(CH2)m-CH(OH)-(CH2)p-C2-6cycloalkylkyl, -(CH2)m-CH(OH)-(CH2)p-aryl, -(CH2)m-CH(OH)-(CH2)p-aryI, -(CH2)m-CH(OH)-(CH2)m-p-heteroaryl, -(CH2)m-NRg-Ci6 alkyl, -(CH2)m-0-Ci6 alkyl, -(CH2)m-S-Ci6 alkyl, -(CH2)m-S02-Ci6 alkyl, -(CH2)m-S02-(CH2)p-aryl, -(CH2)m-C3-7cycloalkyl, -(CH2)m C2-6cycloalkyl, -(CH2)m aryI, -(CH2)m-biphenyl, and -(CH2)m heteroaryl, wherein each
25 CH2, alkyl, alkoxy, cycloalkyl, cyclohexylalkyl, aryl, biphenyl and heteroaryl is unsubstituted or
substituted with one to three groups independently selected from R².

In another embodiment of the present invention, each R¹ is independently selected from
the group consisting of: hydrogen, -Ci-6 alkyl, -(CH2)mCOCi6 alkyl, -(CH2)m-C(0)-(CH2)p-
aryl, -(CH2)mS02-Ci-6 alkyl, (CH2)mS02-(CH2)q-aryl, -(CH2)m C3-7cycloalkyl, -(CH2)m
30 C2-6cycloalkylkyl, -(CH2)m aryl, -(CH2)m-biphenyl, and -(CH2)m heteroaryl, wherein each
CH2, alkyl, cycloalkyl, cyclohexylalkyl, aryl, biphenyl and heteroaryl is unsubstituted or
substituted with one to three groups independently selected from R². In a class of this
embodiment, each R¹ is independently selected from the group consisting of: -C1-6 alkyl,
-COCI6 alkyl, -C(0)-CH2-aryl, -SO2-C1-6 alkyl, -SO2-(CH2)0-1-aryl, -C3-7cycloalkyl, -(CH2)0-2
35 C2-6cycloalkylkyl, -(CH2)0-2 aryl, -(CH2)0-2-biphenyl, and -(CH2)0-2 heteroaryl,
wherein each CH2, alkyl, cycloalkyl, cyclohexylalkyl, aryl, biphenyl and heteroaryl is
unsubstituted or substituted with one to three groups independently selected from R². In another
class of this embodiment, each R¹ is independently selected from the group consisting of:
CH(CH3)2, -CH2CH3, -COCHF2, -CH2-C(0)-phenyl, -C(0)-CH2-phenyl, -C(0)-C1¾-naphthalene, -SO2CH3, -S02-phenyl, -S02-CH2-phenyl, cyclohexyl, -(CH2)2-morpholine, pyrrolidine, azetidine, -phenyl, -(CH2)2-phenyl, -CH2CH(OH)-phenyl, -CH2C(OH)(CH3)-phenyl, -CH2CH(OCH3)-phenyl, -CH2CH(C02CH3)-phenyl, -CH2CH(C02H)-phenyl, -biphenyl, -(CH2)2-pyridine, and pyridine, wherein each CH2, alkyl, cycloalkyl, cycloheteroalkyl, aryl, biphenyl and heteroaryl is unsubstituted or substituted with one to three groups independently selected from R⁰.

In another embodiment of the present invention, each R₁ is independently selected from the group consisting of: -C1-6 alkyl, -COCi-6 alkyl, -C(0)-(CH2)p-aryl, -C3_7cycloalkyl, -CH(CH3)2, -CH2CH3, -COCHF2, -CH2-C(0)-phenyl, -C(0)-CH2-phenyl, -C(0)-C1¾-naphthalene, -SO2CH3, -S02-phenyl, -S02-CH2-phenyl, cyclohexyl, -(CH2)2-morpholine, pyrrolidine, azetidine, -(CH2)2-phenyl, -CH2CH(OH)-phenyl, -(CH2)2-pyridine, and pyridine, wherein each CH2, alkyl, cycloalkyl, cycloheteroalkyl, aryl, and heteroaryl is unsubstituted or substituted with one to three groups independently selected from R⁰. In a class of this embodiment, each R₁ is independently selected from the group consisting of: -(CH2)2-phenyl, -CH2CH(OH)-phenyl, -(CH2)2-pyridine, and pyridine, wherein each CH2, alkyl, cycloalkyl, cycloheteroalkyl, aryl, biphenyl and heteroaryl is unsubstituted or substituted with one to three groups independently selected from R⁰. In another class of this embodiment, each R₁ is independently selected from the group consisting of: -C2-6cycloalkyl, and -(CH2)2-phenyl, wherein each CH2, cycloalkyl and aryl is unsubstituted or substituted with one to three groups independently selected from R⁰. In a class of this embodiment, each R₁ is independently selected from the group consisting of: -(CH2)2-phenyl, and -(CH2)2-pyridine, and pyridine, wherein each CH2, cycloalkyl, and aryl is unsubstituted or substituted with one to three groups independently selected from R⁰. In another class of this embodiment, each R₁ is independently selected from the group consisting of: -C2-6cycloalkyl, and -(CH2)2-phenyl, wherein each CH2, cycloalkyl and phenyl is unsubstituted or substituted with one to three groups independently selected from R⁰. In a class of this embodiment, each R₁ is independently selected from the group consisting of: -C2-6cycloalkyl, and -(CH2)2-phenyl, wherein each CH2, pyrrolidine and phenyl is unsubstituted or substituted with one to three groups independently selected from R⁰. In a subclass of this class, each R₁ is independently selected from the group consisting of: pyrrolidine and -(CH2)2-phenyl, wherein each CH2, pyrrolidine and phenyl is unsubstituted or substituted with one to three groups independently selected from...
Ra. In another subclass of this class, each R is independently selected from the group consisting of: -pyrrolidine, -(CH2)2-phenyl, -CH2CH(OH)phenyl, -CH2C(OH)(CH3)-phenyl, -CH2CH(OCH3)-phenyl, -CH2CH(C02CH3)-phenyl, -CH2CH(C02H)-phenyl, wherein each CH2, pyrrolidine and phenyl is unsubstituted or substituted with one to three groups independently selected from Rα. In another subclass of this class, each R is independently selected from the group consisting of: -pyrrolidine, -(CH2)2-phenyl, -CH2CH(OH)-phenyl, -CH2CH(OCH3)-phenyl, -CH2CH(C02CH3)-phenyl, -CH2CH(C02H)-phenyl, wherein each CH2, pyrrolidine and phenyl is unsubstituted or substituted with one to three groups independently selected from Rα.

In another class of this embodiment, each R is independently selected from the group consisting of: -C2-6 cycloheteroalkyl, wherein cycloheteroalkyl is unsubstituted or substituted with one to three groups independently selected from Rα. In a subclass of this class, R is pyrrolidine, wherein pyrrolidine is unsubstituted or substituted with one to three groups independently selected from Rα.

In another class of this embodiment, R is -(CH2)2 aryl, wherein each CH2 and aryl is unsubstituted or substituted with one to three groups independently selected from Rα. In a subclass of this class, R is -(CH2)2-phenyl, wherein each C1-6 and phenyl is unsubstituted or substituted with one to three groups independently selected from Rα.

In another embodiment of the present invention, each R2 is independently selected from the group consisting of: hydrogen, and -C1-6 alkyl, wherein each alkyl is unsubstituted or substituted with one to three substituents selected from Rβ. In a class of this embodiment of the present invention, each R2 is independently selected from the group consisting of: hydrogen, and -C1-6 alkyl. In another class of this embodiment, each R2 is independently selected from the group consisting of: hydrogen, and -C3-7 cycloalkyl. In another class of this embodiment, R2 is -C1-6 alkyl, wherein each alkyl is unsubstituted or substituted with one to three substituents selected from Rβ. In a subclass of this class, R2 is -CH3.

In another embodiment of the present invention, each R3 is independently selected from the group consisting of: -Cl-6 alkyl, -(CH2)n C3-7 cycloalkyl, -(Cl4)n C2-6 cycloheteroalkyl, -(CH2)n aryl, and -(Cl4)n heteroaryl, wherein each C1-6, alkyl, cycloalkyl, cycloheteroalkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from Rβ. In a class of this embodiment, each R3 is independently selected from the group consisting of: -Cl-6 alkyl, -(C3-7)cycloalkyl, -(CH2)n C2-6 cycloheteroalkyl, -(CH2)n aryl, and -(Cl4)n heteroaryl, wherein each C1-6, aryl, cycloalkyl, cycloheteroalkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from Rβ. In another class of this embodiment, each R3 is independently selected from the group consisting of: -(CH2)n C3-7 cycloalkyl, -(Cl4)n C2-6 cycloheteroalkyl, -(CH2)n aryl, and -(CH2)n.
heteroaryl, wherein each CH₂, cycloalkyl, cycloheteroalkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from R⁰.

In another embodiment of the present invention, each R³ is independently selected from the group consisting of: -C₁₋₆ alkyl, -(CH₂)n aryl, and -(CH₂)n heteroaryl, wherein each C₃⁴, alkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from R⁰. In a class of this embodiment of the present invention, each R³ is independently selected from the group consisting of: -C₁₋₆ alkyl, -(CH₂)₀₋₂ phenyl, and -(C₃⁴)₀₋₂ heteroaryl, wherein each CH₂, alkyl, phenyl and heteroaryl is unsubstituted or substituted with one to three substituents selected from R⁰. In another class of this embodiment, each R³ is independently selected from the group consisting of: -CH₃, -CH₂CH₃, -phenyl, -(CH₂)phenyl, pyridine, pyrimidine, pyrazine, and -(CH₂)₂ imidazole, wherein each CH₂, alkyl, phenyl, pyridine, pyrimidine, pyrazine and imidazole is unsubstituted or substituted with one to three substituents selected from R⁰.

In another embodiment of the present invention, each R³ is independently selected from the group consisting of: -(C₃⁴)n aryI, and -(C₃⁴)n heteroaryl, wherein each CH₂, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from R⁰. In another embodiment of the present invention, each R³ is independently selected from the group consisting of: -(CH₂)₀₋₂ phenyl, and -(CH₂)₀₋₂ heteroaryl, wherein each CH₂, phenyl and heteroaryl is unsubstituted or substituted with one to three substituents selected from R⁰. In a class of this embodiment, each R³ is independently selected from the group consisting of: -(CH₂)₀₋₂ phenyl, -(CH₂)₀₋₂ pyridine, -(CH₂)₀₋₂ pyrimidine, -(CH₂)₀₋₂ pyrazine, -(CH₂)₀₋₂ oxazole and -(CH₂)₀₋₂ imidazole, wherein each CH₂, phenyl, pyridine, pyrimidine, pyrazine, oxazole and imidazole is unsubstituted or substituted with one to three substituents selected from R⁰. In another class of this embodiment, each R³ is independently selected from the group consisting of: -(CH₂)₀₋₂ phenyl, -(CH₂)₀₋₂ pyridine, -(CH₂)₀₋₂ pyrimidine, -(C₈⁴)₀₋₂ pyrazine and -(C₃⁴)₀₋₂ imidazole, wherein each CH₂, phenyl, pyridine, pyrimidine, pyrazine and imidazole is unsubstituted or substituted with one to three substituents selected from R⁰. In another embodiment of the present invention, each R³ is independently selected from the group consisting of: -(C₃⁴)₀₋₂ phenyl, and -(C₃⁴)₀₋₂ heteroaryl, wherein each CH₂, phenyl and heteroaryl is unsubstituted or substituted with one to three substituents selected from R⁰. In another class of this embodiment, each R³ is independently selected from the group consisting of: -phenyl, -(CH₂)phenyl, -pyridine, pyrimidine, pyrazine, -(CH₂)₂ imidazole, wherein each CH₂, phenyl, pyridine, pyrimidine, pyrazine and imidazole is unsubstituted or substituted with one to three substituents selected from R⁰.
each phenyl, pyridine and pyrimidine is unsubstituted or substituted with one to three substituents selected from RP.

In another embodiment of the present invention, each R4 is independently selected from the group consisting of: -C1-6 alkyl, -C1-6 alkoxy, -CF3, -(Cl3)s C3-7cycloalkyl, -(CH2)s C2-6cycloalkyl, -(CH2)s aryl, and -(CH2)s heteroaryl, wherein each C1-6, alkyl, alkoxy, cycloalkyl, cycloalkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from Rd. In a class of this embodiment, each R4 is independently selected from the group consisting of: -C1-6 alkyl, -C1-6 alkoxy, -CF3, -(Cl3)s C3-7cycloalkyl, -(CH2)s C2-6cycloalkyl, -(CH2)s aryl, and -(CH2)s heteroaryl, wherein each CH2, alkyl, alkoxy, cycloalkyl, cycloalkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from Rd. In another class of this embodiment, each R4 is independently selected from the group consisting of: -(CH2)s aryl, and -(CH2)s heteroaryl, wherein each CH2, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from Rd.

In another embodiment of the present invention, each R4 is independently selected from the group consisting of: -(Cl3)s aryl, wherein each Cl3 and aryl is unsubstituted or substituted with one to three substituents selected from Rd. In a class of this embodiment, each R4 is independently selected from the group consisting of: -(CH2)0-1 aryl, wherein each Cl3, and aryl is unsubstituted or substituted with one to three substituents selected from Rd. In another class of this embodiment, R4 is aryl, wherein aryl is unsubstituted or substituted with one to three substituents selected from Rd. In another class of this embodiment, each R4 is -phenyl, wherein phenyl is unsubstituted or substituted with one to three substituents selected from Rd. In another class of this embodiment, each R4 is -phenyl, wherein phenyl is unsubstituted or substituted with one or two substituents selected from Rd. In another class of this embodiment, each R4 is -phenyl, wherein phenyl is unsubstituted or substituted with one substituent selected from Rd.

In another embodiment of the present invention, each R5 is independently selected from the group consisting of: hydrogen, and -C1-6 alkyl, wherein each alkyl is unsubstituted or substituted with one to three substituents selected from Rxs. In a class of this embodiment of the present invention, each R5 is independently selected from the group consisting of: hydrogen, and -C1-6 alkyl. In another class of this embodiment, each R5 is independently selected from the group consisting of: hydrogen, and -CH3. In another class of this embodiment, R5 is hydrogen. In another class of this embodiment, R5 is -C1-6 alkyl, wherein each alkyl is unsubstituted or substituted with one to three substituents selected from Rxs. In a subclass of this class, R5 is -CH3.
In another embodiment of the present invention, each Rₐ is independently selected from the group consisting of: hydrogen, -OH, oxo, -Cl-₆ alkyl, -OC₃-₆ alkyl, halogen, -CF₃, -OCF₃, -CN, -C(0)Cl-₆ alkyl, -CO₂H, -CO₂Cl₆ alkyl, -(CH₂)ₗ C₃-₇cycloalkyl, -(CH₂)ₗ heteroaryl, -(CH₂)ₗ phenyl, -(CH(phenyl))₂, -SO₂Cl₆ alkyl, -Sₐ2N(Rf)₂; -SO₂-C₃₋₇cycloalkyl, -SO₂-C₆cycloheteroalkyl, -Sₐ2-aryl, and -SO₂-heteroaryl.

In another embodiment of the present invention, each Rₐ is independently selected from the group consisting of: hydrogen, -OH, oxo, -Cl-₆ alkyl, -OC₃-₆ alkyl, halogen, -CF₃, -CO₂H, -CO₂Cl₆ alkyl, -(CH₂)₀-₁ ary1, -CH(phenyl)₂, -SO₂Cl₆ alkyl, -SO₂N(Rf)₂, and -SO₂-C₆cycloheteroalkyl.

In another embodiment of the present invention, each Rₐ is independently selected from the group consisting of: -OH, oxo, -Cl-₆ alkyl, -OC₃-₆ alkyl, halogen, -CF₃, -CO₂H, -CO₂Cl₆ alkyl, -(CH₂)₀-₁ phenyl, -CH(phenyl)₂, -SO₂Cl₆ alkyl, -SO₂N(Rf)₂, and -SO₂-C₆cycloheteroalkyl. In a class of this embodiment, each Rₐ is independently selected from the group consisting of: -OH, oxo, -CH₃, -OCH₃, F, Cl, -CF₃, -CO₂H, -CO₂CH₃, phenyl, -CH(phenyl)₂, -SO₂CH₃, -SO₂N(CH₃)₂, and -SO₂-morpholine.

In another embodiment of the present invention, each Rₐ is independently selected from the group consisting of: -OH, oxo, -OC₃-₆ alkyl, -CF₃, -CO₂H, -CO₂Cl₆ alkyl, and -SO₂Cl₆ alkyl. In a class of this embodiment, each Rₐ is independently selected from the group consisting of: -OH, oxo, -OCH₃, -CF₃, -CO₂H, -CO₂CH₃, and -SO₂CH₃.

In another embodiment of the present invention, each RB is independently selected from the group consisting of: halogen, -Cl-₆ alkyl, -OC₃-₆ alkyl, -CO₂H, and -CO₂Cl₆ alkyl, wherein alkyl can be substituted with one to three fluorines. In a class of this embodiment, each RB is independently selected from the group consisting of: halogen, and -Cl-₆ alkyl, wherein alkyl can be substituted with one to three fluorines. In another class of this embodiment, each RB is independently selected from the group consisting of: halogen, and -Cl-₆ alkyl. In another class of this embodiment, RB is -Cl-₆ alkyl. In another class of this embodiment, RB is -Cl-₆ alkyl.

In another embodiment of the present invention, each Rₐ is independently selected from the group consisting of: hydrogen, -OH, oxo, -Cl-₆ alkyl, -OC₃-₆ alkyl, halogen, -CF₃, -OCF₃, -CN, -CO₂H, and -CO₂Cl₆ alkyl. In a class of this embodiment, each Rₐ is independently selected from the group consisting of: hydrogen, -OH, oxo, -Cl-₆ alkyl, -OC₃-₆ alkyl, halogen, -CF₃, -OCF₃, -CN, -CO₂H, and -CO₂Cl₆ alkyl.

In another embodiment of the present invention, each Rₐ is independently selected from the group consisting of: -Cl-₆ alkyl, -OC₃-₆ alkyl, halogen, -CN, -CO₂H, and -CO₂Cl₆ alkyl. In a class of this embodiment, each Rₐ is independently selected from the group consisting of: -Cl-₆ alkyl, -OC₃-₆ alkyl, halogen, and -CO₂Cl₆ alkyl. In a subclass of this class, each Rₐ is independently selected from the group consisting of: -CH₃, -OCH₃, Cl, -CN, -CO₂H, and -CO₂CH₂CH₃.

In another embodiment of the present invention, each Rₐ is independently selected from the group consisting of: -Cl-₆ alkyl, -OC₃-₆ alkyl, halogen, -CN, -CO₂H, and -CO₂Cl₆ alkyl. In a class of this embodiment, each Rₐ is independently selected from the group consisting of: -Cl-₆ alkyl, -OC₃-₆ alkyl, halogen, and -CO₂Cl₆ alkyl. In a subclass of this class, each Rₐ is independently selected from the group consisting of: -CH₃, -OCH₃, Cl, and -CO₂CH₂CH₃.
In another embodiment of the present invention, each Rd is independently selected from the group consisting of: hydrogen, -OH, o xo, -C1-6 alkyl, -OC1-6 alkyl, halogen, -CF3, -OCF3, -CN, -CO2H, and -CO2CI-6alkyl. In a class of this embodiment, each Rd is independently selected from the group consisting of: F, Cl, -CN, -CO2H, and -CO2CH2CH3. In another class of this embodiment, Rd is halogen. In another class of this embodiment, each Rd is independently selected from the group consisting of: F, Cl, and Br. In another class of this embodiment, each Rd is independently selected from the group consisting of: F, and Cl. In another class of this embodiment, Rd is F. In another class of this embodiment, Rd is Cl.

In another embodiment of the present invention, each Re is independently selected from the group consisting of: halogen, -C1-6 alkyl, -OCi-6 alkyl, -CO2H, and -CO2C1-6 alkyl, wherein alkyl can be substituted with one to three fluorines. In a class of this embodiment, each Re is independently selected from the group consisting of: halogen, and -Ci-6 alkyl, wherein alkyl can be substituted with one to three fluorines. In another class of this embodiment, each Re is independently selected from the group consisting of: halogen, and -C1-6 alkyl. In another class of this embodiment, Re is halogen. In another class of this embodiment, Re is -Ci-6 alkyl.

In another embodiment of the present invention, each Rf is independently selected from the group consisting of: hydrogen, and -Ci-6 alkyl. In a class of this embodiment, each Rf is independently selected from the group consisting of: hydrogen, and _CH3. In another class of this embodiment, Rf is hydrogen. In another class of this embodiment, Rf is -C1-6 alkyl. In a subclass of this class, Rf is -CH3.

In another embodiment of the present invention, each Rg is independently selected from the group consisting of: hydrogen, and -Ci-6 alkyl. In a class of this embodiment, each Rg is independently selected from the group consisting of: hydrogen, and -CH3. In another class of this embodiment, Rg is hydrogen. In another class of this embodiment, Rg is -C1-6 alkyl. In a subclass of this class, Rg is -CH3.

In another embodiment of the present invention, m is 0, 1, 2 or 3. In a class of this embodiment, m is 0, 1 or 2. In another class of this embodiment, m is 0 or 1. In another class of this embodiment, m is 1 or 2. In another class of this embodiment, m is 0 or 2. In another class of this embodiment, m is 0. In another class of this embodiment, m is 1. In another class of this embodiment, m is 2. In another class of this embodiment, m is 3.

In another embodiment of the present invention, n is 0, 1, 2 or 3. In a class of this embodiment, n is 0, 1 or 2. In another class of this embodiment, n is 0 or 1. In another class of this embodiment, n is 1 or 2. In another class of this embodiment, n is 0 or 2. In another class of this embodiment, n is 0. In another class of this embodiment, n is 1. In another class of this embodiment, n is 2. In another class of this embodiment, n is 3.

In another embodiment of the present invention, p is 0, 1, 2 or 3. In a class of this embodiment, p is 0, 1 or 2. In another class of this embodiment, p is 0 or 1. In another class of this embodiment, p is 1 or 2. In another class of this embodiment, p is 0 or 2. In another class of
this embodiment, p is 0. In another class of this embodiment, p is 1. In another class of this embodiment, p is 2. In another class of this embodiment, p is 3.

In another embodiment of the present invention, q is 0, 1, 2 or 3. In a class of this embodiment, q is 0, 1 or 2. In another class of this embodiment, q is 0 or 1. In another class of this embodiment, q is 1 or 2. In another class of this embodiment, q is 0 or 2. In another class of this embodiment, q is 0. In another class of this embodiment, q is 1. In another class of this embodiment, q is 2. In another class of this embodiment, q is 3.

In another embodiment of the present invention, r is 0, 1, 2, 3 or 4. In a class of this embodiment, r is 0, 1, 2 or 3. In a class of this embodiment, r is 0, 1 or 2. In another class of this embodiment, r is 0 or 1. In another class of this embodiment, r is 1 or 2. In another class of this embodiment, r is 0 or 2. In another class of this embodiment, r is 0. In another class of this embodiment, r is 1. In another class of this embodiment, r is 2. In another class of this embodiment, r is 3. In another class of this embodiment, r is 4.

In another embodiment of the present invention, s is 0, 1, 2 or 3. In a class of this embodiment, s is 0, 1 or 2. In another class of this embodiment, s is 0 or 1. In another class of this embodiment, s is 1 or 2. In another class of this embodiment, s is 0 or 2. In another class of this embodiment, s is 0. In another class of this embodiment, s is 1. In another class of this embodiment, s is 2. In another class of this embodiment, s is 3.

In another embodiment of the present invention, t is 0, 1, 2 or 3. In a class of this embodiment, t is 0, 1 or 2. In another class of this embodiment, t is 0 or 1. In another class of this embodiment, t is 1 or 2. In another class of this embodiment, t is 0 or 2. In another class of this embodiment, t is 0. In another class of this embodiment, t is 1. In another class of this embodiment, t is 2. In another class of this embodiment, t is 3.

In another embodiment of the present invention, u is 1, 2, or 3. In a class of this embodiment, u is 1 or 2. In another class of this embodiment, u is 1 or 3. In another class of this embodiment, u is 2 or 3. In another class of this embodiment, u is 2. In another class of this embodiment, u is 3.

In another embodiment of the present invention, the invention relates to compounds of structural formula I, wherein:

X is independently selected from N and CH;
each R^1 is independently selected from the group consisting of:

(1) -Ci-6 alkyl,
(2) -COCj-6 alkyl,
(3) -C(0)-(CH2)p-aryl,
(4) -C3-7cycloalkyl,
(5) -(CH2)0-2 C2-6cycloheteroalkyl,
(6) -(CH2)0-2 aryl, and
(7) -(CH2)0-2 heteroaryl,
wherein each CH₂, alkyl, cycloalkyl, cycloheteroalkyl, aryl, and heteroaryl is unsubstituted or substituted with one to three groups independently selected from R¹; 
R² is hydrogen; 
each R³ is independently selected from the group consisting of: 
(1) -Ci-6 alkyl, 
(2) -(CH₂)n aryl, and 
(3) -(CH₄)n heteroaryl, 
wherein each CH₂, alkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from R⁹; and 
each R⁴ is independently selected from the group consisting of: -(CH₂)s aryl, wherein each CH₂ and aryl is unsubstituted or substituted with one to three substituents selected from Rd; 
or a pharmaceutically acceptable salt thereof. 

In another embodiment of the present invention, the invention relates to compounds of structural formula I, wherein: 

"a" is absent; 
X is independently selected from N and CH; 
each R¹ is independently selected from the group consisting of: 
(1) -Ci-6 alkyl, 
(2) -COCi-6 alkyl, 
(3) -C(0)-(CH₂)p-aryl, 
(4) -C3-7cycloalkyl, 
(5) -(CH₂)0-2 C2-6cycloheteroalkyl, 
(6) -(CH₂)0-2 aryl, and 
(7) -(CH₂)0-2 heteroaryl, 
wherein each CH₂, alkyl, cycloalkyl, cycloheteroalkyl, aryl, and heteroaryl is unsubstituted or substituted with one to three groups independently selected from R³; 
R² is hydrogen; 
each R³ is independently selected from the group consisting of: 
(1) -Ci-6 alkyl, 
(2) -(CH₂)n aryl, and 
(3) -(CH₂)n heteroaryl, 
wherein each CH₂, alkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from R⁹; and 
each R⁴ is independently selected from the group consisting of: -(CH₂)s aryl, wherein each CH₂ and aryl is unsubstituted or substituted with one to three substituents selected from Rd; 
or a pharmaceutically acceptable salt thereof. 

In another embodiment of the present invention, the invention relates to compounds of structural formula I, wherein:
"a" is absent;
X is independently selected from N and CH;
each R¹ is independently selected from the group consisting of:
   (1) -(CH₂)₀-² C₂-6cycloalkyl, and
   (2) -(CH₂)₀-² aryl,
wherein each CH₂, cycloalkyl and aryl is unsubstituted or substituted with one to three
groups independently selected from R²;
R² is hydrogen;
each R³ is independently selected from the group consisting of:
   (1) -(CH₂)ⁿ aryl, and
   (2) -(CH₂)ⁿ heteroaryl,
wherein each CH₂, aryl and heteroaryl is unsubstituted or substituted with one to three
substituents selected from R³; and
R⁴ is -phenyl, wherein phenyl is unsubstituted or substituted with one to three substituents selected from
or a pharmaceutically acceptable salt thereof.

In another embodiment of the present invention, the invention relates to compounds of
structural formula I, wherein:
"a" is absent;
X is independently selected from N and CH;
each R¹ is independently selected from the group consisting of:
   (1) pyrrolidine, and
   (2) -(CH₂)₂-phenyl,
wherein each CH₂, pyrrolidine and phenyl is unsubstituted or substituted with one to three
groups independently selected from R²;
R² is hydrogen;
each R³ is independently selected from the group consisting of:
   (1) -(CH₂)₀-² phenyl,
   (2) -(CH₂)₀-² pyridine,
   (3) -(CH₂)₀-² pyrimidine,
   (4) -(CH₂)₀-² pyrazine, and
   (5) -(CH₂)₀-² imidazole,
wherein each CH₂, phenyl, pyridine, pyrimidine, pyrazine and imidazole is unsubstituted or
substituted with one to three substituents selected from R³; and
R⁴ is -phenyl, wherein phenyl is unsubstituted or substituted with one to three substituents
selected from R⁴;
or a pharmaceutically acceptable salt thereof.
In another embodiment of the present invention, the invention relates to compounds of structural formula Ia-1 and Ia-2:

![Structural formula Ia-1](image)

and

![Structural formula Ia-2](image)

In another embodiment of the present invention, the invention relates to compounds of structural formula Ib-1 and Ib-2:

![Structural formula Ib-1](image)

and

![Structural formula Ib-2](image)

In another embodiment of the present invention, the invention relates to compounds of structural formula Ic-1 and Ic-2:

![Structural formula Ic-1](image)

and

![Structural formula Ic-2](image)

In another embodiment of the present invention, the invention relates to compounds of structural formula Id-1 and Id-2:

![Structural formula Id-1](image)

and

![Structural formula Id-2](image)

In another embodiment of the present invention, the invention relates to compounds of structural formula Ie-1 and Ie-2:
In another embodiment of the present invention, the invention relates to compounds of structural formula If-1 and If-2:

\[
\begin{align*}
\text{(If-1)} & \quad (R^2)_1 \quad \text{and} \quad (R^2)_1 \\
\text{(If-2)} & \quad \text{and}
\end{align*}
\]

In another embodiment of the present invention, the invention relates to compounds of structural formula Ig-1 and Ig-2:

\[
\begin{align*}
\text{(Ig-1)} & \quad (R^3)_1 \\
\text{(Ig-2)} & \quad \text{and}
\end{align*}
\]

In another embodiment of the present invention, the invention relates to compounds of structural formula Ih-1 and Ih-2:

\[
\begin{align*}
\text{(Ih-1)} & \quad (R^4)_1 \\
\text{(Ih-2)} & \quad \text{and}
\end{align*}
\]

Illustrative, but nonlimiting, examples of compounds of the present invention that are useful as inhibitors of PrCP are the following:
and pharmaceutically acceptable salts thereof.
As used herein the following definitions are applicable.

"Alkyl", as well as other groups having the prefix "alk", such as alkoxy and alkanoyl, means carbon chains which may be linear or branched, and combinations thereof, unless the carbon chain is defined otherwise. Examples of alkyl groups include methyl, ethyl, propyl, isopropyl, butyl, sec- and tert-butyl, pentyl, hexyl, heptyl, octyl, nonyl, and the like. When no number of carbon atoms is specified, C1-6 is intended.

"Alkenyl" means carbon chains up to 10 carbons, unless otherwise specified, which contain at least one carbon-carbon double bond, and which may be linear or branched or combinations thereof. Examples of alkenyl include vinyl, allyl, isopropenyl, pentenyl, hexenyl, heptenyl, 1-propenyl, 1-butenyl, 2-butenyl, 2-methyl-2-butenyl, and the like.

"Alkynyl" means carbon chains up to 10 carbons, if not otherwise specified, which contain at least one carbon-carbon triple bond, and which may be linear or branched or combinations thereof. Examples of alkynyl include ethynyl, propynyl, butynyl, pentynyl, 3-methyl-1-pentynyl, 2-heptynyl and the like.

The term "alkoxy" refers to straight or branched chain alkoxydes of the number of carbon atoms specified (e.g., C1-6 alkoxy), or any number within this range [i.e., methoxy (MeO-), ethoxy, isopropoxy, etc.].

The term "alkylthio" refers to straight or branched chain alkylsulfides of the number of carbon atoms specified (e.g., C1-g alkylthio), or any number within this range [i.e., methylthio (MeS-), ethylthio, isopropylthio, etc.].

The term "alkylsulfonyl" refers to straight or branched chain alkylsulfones of the number of carbon atoms specified (e.g., C1-6 alkylsulfonyl), or any number within this range [i.e., methylsulfonyl (MeS02-) , ethylsulfonyl, isopropylsulfonyl, etc.].

The term "alkyloxycarbonyl" refers to straight or branched chain esters of a carboxylic acid derivative of the present invention of the number of carbon atoms specified (e.g., C1-6 alkyloxycarbonyl), or any number within this range [i.e., methyloxycarbonyl (MeOCO-), ethyloxycarbonyl, or butyloxycarbonyl].

"Cycloalkyl" means mono- or bicyclic or bridged saturated carbocyclic rings, each having from 3 to 14 carbon atoms. Examples of cycloalkyl include cyclopentyl, cyclobutyl, cyclohexyl, cycloheptyl, cyclooctyl, and decahydronaphthyl, and the like.

"Cycloalkenyl" means nonaromatic, mono- or bicyclic or bridged carbocyclic rings, each having from 3 to 14 carbon atoms and containing at least one double bond. Examples of cycloalkyl include cyclopropenyl, cyclobutenyl, cyclopentenyl, cyclohexenyl, cycloheptenyl, cyclooctenyl, decahydronaphthyl, bicyclo[2.2.1]hept-5-en-2-yl, and the like.

"Cycloheteroalkyl" means nonaromatic, mono- or bicyclic or bridged saturated carbocyclic rings, each having from 2 to 14 carbon atoms and containing 1, 2, 3, 4 or 5 heteroatoms selected from N, NH, O and S. Examples of cycloheteroalkyl include tetrahydrofurananyl, azetidinyl, perhydroazepinyl, dihydrofuranyl, dioxanyl, oxanyl, morpholinyl,
1,4-dithianyl, piperazinyl, piperidinyl, 1,3-dioxolanyl, imidazolidinyl, imidazolinyl, pyrrolinyl, pyrrolidinyl, pyrany1, tetrahydropyranyl, dihydroxypyrany1, oxathiolanyl, dithiolanyl, 1,3-dithianyl, oxathianyl, thiomorpholinyl, dioxidoisothiazolidinyl, azacycloheptyl, diazobicyclo[3.2.1]-octane, and hexahydroindazolyl. The cycloheteroalkyl ring may be substituted on the ring carbons
and/or the ring nitrogens.

"Cycloheteroalkenyl" means nonaromatic mono- or bicyclic or bridged rings each having from 2 to 14 carbon atoms containing at least one double bond and containing 1, 2, 3, 4 or 5 heteroatoms selected from N, NH, O and S. Examples of cycloheteroalkenyl include 1,2,4-oxadiazol-5-one, 1,2,4-thiadiazol-5-one, 1,2,4-triazol-3-one, and 1,2,3,6-tetrahydropyridine, dihydro-1,3,4-oxadiazole, and [1,6]-dihydropyridine and the like. In one embodiment of the present invention, cycloheteroalkenyl is dihydro-1,3,4-oxadiazole. In another embodiment of the present invention, cycloheteroalkenyl is [1,6]-dihydropyridine.

"Aryl" means a monocyclic, bicyclic or tricyclic ring system containing 5-14 carbon atoms, wherein at least one of the rings is aromatic. Aryl thus includes ring systems in which an aromatic ring is fused to a non-aromatic ring, such as a cycloalkyl or cycloalkenyl ring.

Examples of aryl include phenyl, naphthalene, biphenyl, indane and 5,6,7,8-tetrahydronaphthalene, and the like. In one embodiment of the present invention, aryl is phenyl, naphthalene, biphenyl, indane, and 5,6,7,8-tetrahydronaphthalene. In another embodiment of the present invention, aryl is phenyl, naphthalene, indane and 5,6,7,8-tetrahydronaphthalene. In one class of this embodiment, aryl is phenyl and naphthalene. In another class of this embodiment, aryl is phenyl.

"Heteroaryl" means a monocyclic, bicyclic or tricyclic ring system containing 5-14 carbon atoms and containing 1, 2, 3, 4 or 5 heteroatoms selected from N, NH, O and S wherein at least one of the heteroatom containing rings is aromatic. Heteroaryls thus includes heteroaryls fused to other kinds of rings, such as aryls, cycloalkyls and heterocycles that are not aromatic. Heteroaryl includes ring systems in which an aromatic heteroatom containing ring is fused to a non-aromatic ring, such as a cycloalkyl, cycloalkenyl, cyclohexoalkenyl or cyclohexoalkyl ring, and also includes ring systems in which an aryl ring is fused to a non-aromatic heteroatom containing ring, such as a cyclohexoalkyl or cyclohexoalkenyl ring. Examples of heteroaryls include: pyrazole, pyridine, pyrazine, pyrrole, pyrimidine, pyridazine, benzoimidazole, quinoline, isothiazole, isoquinoline, indole, indazole, carbazole, benzotriazole, benzofuran, benzothiazole, benzothiophene, benzoisoxazolone, oxazole, oxadiazole, furan, benzoxazole, isoxazole, indoline, isoindoline, tetrazole, imidazole, oxadiazole (in particular, 1,3,4-oxadiazol-2-yl and 1,2,4-oxadiazol-3-yl), thiazole, thiophene, thiazadiazole, triazole, triazine, tetrazole, thiene, benzothiazole, benzopyrazole, benzothiadiazole, dihydrobenzofuran, indazole, isoindole, dihydrobenzothiophene, indolizine, cinnoline, phthalazine, quinazoline, naphthyridine, carbazole, quinoxaline, purine, isobenzofuran, benzothiene, isoquinoline, dibenzofuran, isothiazole, imidazopyridine, benzodioxole, dihydropyridine, dihydropyrrolopyridine, dihydrobenzoxazine,
benzodioxole, benzodioxine, pyrrolopyridine, triazolopyridine, dihydropyridooxazine, dihydrobenzoxazine, dihydroindole, dihydroisoindole, dihydrobenzoiraidazole, dihydroquinoline, tetrahydroisoquinoline, tetrahydrocyclopentaindole, tetrahydroquinoxaline, and tetrahydropyridine. For heterocycloalkyl and heteroaryl groups, rings and ring systems containing from 3-15 atoms are included, forming 1-3 rings.

"Halogen" includes fluorine, chlorine, bromine and iodine. Chlorine and fluorine are generally preferred. Fluorine is most preferred when the halogens are substituted on an alkyl or alkoxy group (e.g. CF3O and CF3CH2O). In one embodiment of the present invention, halogen is selected from fluorine, chlorine, and bromine.

"Oxo" means the functional group "=O" which is an oxygen atom connected to the molecule via a double bond, such as, for example, (1) "C=(O)", that is a carbonyl group; (2) "S=(O)\textsuperscript{2}" (O\textsuperscript{2})\textsuperscript{2}, that is, a sulfoxide group; and (3) "N=(O)\textsuperscript{2}", that is, an N-oxide group, such as pyridyl-N-oxide.

In choosing compounds of the present invention, one of ordinary skill in the art will recognize that the various substituents, i.e. R\textsuperscript{1}, R\textsuperscript{2}, etc., are to be chosen in conformity with well-known principles of chemical structure connectivity and stability. When any variable (e.g., R\textsuperscript{1}, R\textsuperscript{2}, etc.) occurs more than one time in any constituent or in formula I, its definition on each occurrence is independent of its definition at every other occurrence. Also, combinations of substituents and/or variables are permissible only if such combinations result in stable compounds. A squiggly line across a bond in a substituent variable represents the point of attachment.

Under standard nomenclature used throughout this disclosure, the functional group adjacent to the point of attachment is described first, with our without a bond "\ldots", followed by the terminal portion of the designated side chain. For example, a -C\textsubscript{15}-alkylcarbonylamino C\textsubscript{1-6} alky substituent is equivalent to:

\[
\text{C}_{15}\text{alkyl}! \text{C-NH-C}_{1-6}\text{alkyl}
\]

The term "substituted" shall be deemed to include multiple degrees of substitution by a named substituent. Where multiple substituent moieties are disclosed or claimed, the substituted compound can be independently substituted by one or more of the disclosed or claimed substituent moieties, singly or pluraly. By independently substituted, it is meant that the (two or more) substituents can be the same or different.

The terms "compounds of structural formula I" and "formula I" include the compounds of structural formulas 1-1, 1-2, Ia-1, Ia-2, Ib-1, Ib-2, Ic-1, Ic-2, Id-1, Id-2, Ie-1, Ie-2, If-1, If-2, Ig-1, Ig-2, Ih-1, and Ih-2, and pharmaceutically acceptable salts thereof. The terms "compounds of structural formula 1-1 and/or 1-2" and "formula I-1 and/or 1-2" include the compounds of
structural formulas Ia-1, Ia-2, Ib-1, Ib-2, Ic-1, Ic-2, Id-1, Id-2, Ie-1, Ie-2, If-1, If-2, Ig-1, Ig-2, Ih-1, and Ih-2, and pharmaceutically acceptable salts thereof.

Compounds of structural formula I may be separated into their individual diastereoisomers by, for example, fractional crystallization from a suitable solvent, for example methanol or ethyl acetate or a mixture thereof, or via chiral chromatography using an optically active stationary phase. Absolute stereochemistry may be determined by X-ray crystallography of crystalline products or crystalline intermediates which are derivatized, if necessary, with a reagent containing an asymmetric center of known absolute configuration. Alternatively, any stereoisomer of a compound of the general structural formula I may be obtained by stereospecific synthesis using optically pure starting materials or reagents of known absolute configuration.

Compounds of the Formula I may be separated into diastereomeric pairs of enantiomers by, for example, fractional crystallization from a suitable solvent, for example MeOH or ethyl acetate or a mixture thereof. The pair of enantiomers thus obtained may be separated into individual stereoisomers by conventional means, for example by the use of an optically active amine as a resolving agent or on a chiral HPLC column. Alternatively, any enantiomer of a compound of the general Formula I may be obtained by stereospecific synthesis using optically pure starting materials or reagents of known configuration.

If desired, racemic mixtures of the compounds may be separated so that the individual enantiomers are isolated. Racemic mixtures can be separated into their individual enantiomers by any of a number of conventional methods, which include chiral chromatography, derivatization with a chiral auxiliary followed by separation by chromatography or crystallization, and fractional crystallization of diastereomeric salts. The separation can be carried out by methods well known in the art, such as the coupling of a racemic mixture of compounds to an enantiomerically pure compound to form a diastereomeric mixture, followed by separation of the individual diastereomers by standard methods, such as fractional crystallization or chromatography. The coupling reaction is often the formation of salts using an enantiomerically pure acid or base. The diastereomeric derivatives may then be converted to the pure enantiomers by cleavage of the added chiral residue. The racemic mixture of the compounds can also be separated directly by chromatographic methods utilizing chiral stationary phases, which methods are well known in the art.

Some of the compounds described herein contain olefinic double bonds, and unless specified otherwise, are meant to include both E and Z geometric isomers.

Some of the compounds described herein may exist as tautomers, which have different points of attachment of hydrogen accompanied by one or more double bond shifts. For example, a ketone and its enol form are keto-enol tautomers. The individual tautomers as well as mixtures thereof are encompassed with compounds of the present invention.

In the compounds of structural formula I, the atoms may exhibit their natural isotopic abundances, or one or more of the atoms may be artificially enriched in a particular isotope
having the same atomic number, but an atomic mass or mass number different from the atomic mass or mass number predominately found in nature. The present invention is meant to include all suitable isotopic variations of the compounds of structural formula I. For example, different isotopic forms of hydrogen (H) include protium (\(^1\)H) and deuterium (\(^2\)H). Protium is the predominant hydrogen isotope found in nature. Enriching for deuterium may afford certain therapeutic advantages, such as increasing in vivo half-life or reducing dosage requirements, or may provide a compound useful as a standard for characterization of biological samples. Isotopically-enriched compounds within structural formula I, can be prepared without undue experimentation by conventional techniques well known to those skilled in the art or by processes analogous to those described in the Schemes and Examples herein using appropriate isotopically-enriched reagents and/or intermediates.

It will be understood that, as used herein, references to the compounds of structural formula I are meant to also include the pharmaceutically acceptable salts, and also salts that are not pharmaceutically acceptable when they are used as precursors to the free compounds or their pharmaceutically acceptable salts or in other synthetic manipulations. The compounds of the present invention may be administered in the form of a pharmaceutically acceptable salt.

The term "pharmaceutically acceptable salt" refers to salts prepared from pharmaceutically acceptable non-toxic bases or acids including inorganic or organic bases and inorganic or organic acids. Salts of basic compounds encompassed within the term "pharmaceutically acceptable salt" refer to non-toxic salts of the compounds of this invention which are generally prepared by reacting the free base with a suitable organic or inorganic acid. Representative salts of basic compounds of the present invention include, but are not limited to, the following: acetate, benzenesulfonate, benzoate, bicarbonate, bisulfate, bitartrate, borate, bromide, camysylate, carbonate, chloride, clavulanate, citrate, edetate, edisylate, estolate, esylate, fumarate, gluceptate, gluconate, glutamate, hexylresorcinate, hydrobromide, hydrochloride, hydroxynaphthoate, iodide, isothionate, lactate, lactobionate, laurate, malate, maleate, mandelate, mesylate, methylbromide, methylnitrate, methylsulfate, mucate, napsyate, nitrate, N-methylglucamine ammonium salt, oleate, oxalate, pamoate (embonate), palmitate, pantothenate, phosphate/diphosphate, polygalacturonate, salicylate, stearate, sulfate, subacetate, succinate, tannate, tartrate, teoclate, tosylate, triethiodide and valerate. Furthermore, where the compounds of the invention carry an acidic moiety, suitable pharmaceutically acceptable salts thereof include, but are not limited to, salts derived from inorganic bases including aluminum, ammonium, calcium, copper, ferric, ferrous, lithium, magnesium, manganic, mangamous, potassium, sodium, zinc, and the like. Particularly preferred are the ammonium, calcium, magnesium, potassium, and sodium salts. Salts derived from pharmaceutically acceptable organic non-toxic bases include salts of primary, secondary, and tertiary amines, cyclic amines, and basic ion-exchange resins, such as arginine, betaine, caffeine, choline, N,N-dibenzylethlenediamine, diethylamine, 2-diethylaminoethanol, 2-dimethylaminoethanol,
ethanolamine, ethylenediamine, N-ethylmorpholine, N-ethylpiperidine, glucamine, glucosamine, histidine, isopropylamine, lysine, methylglucamine, morpholine, piperazine, piperidine, polyamine resins, procaine, purines, theobromine, triethylamine, trimethylamine, tripropylamine, tromethamine, and the like.

Also, in the case of a carboxylic acid (-COOH) or alcohol group being present in the compounds of the present invention, pharmaceutically acceptable esters of carboxylic acid derivatives, such as methyl, ethyl, or pivaloyloxymethyl, or acyl derivatives of alcohols, such as acetyl, pivaloyl, benzoyl, and aminoacyl, can be employed. Included are those esters and acyl groups known in the art for modifying the solubility or hydrolysis characteristics for use as sustained-release or prodrug formulations.

Furthermore, some of the crystalline forms for compounds of the present invention may exist as polymorphs and as such are intended to be included in the present invention. In addition, some of the compounds of the instant invention may form solvates with water or common organic solvents. Such solvates are encompassed within the scope of this invention.

The compounds of formula I are effective as inhibitors of prolylcarboxypeptidase (PRCP). The compounds of formula I are therefore useful for the treatment, control and/or prevention of diseases, disorders or conditions responsive to the inhibition of the prolylcarboxypeptidase (PRCP) enzyme, including but not limited to: abnormal metabolism, obesity, diabetes, metabolic syndrome, obesity related disorders, diabetes related disorders, hypertension, dyslipidemia, stroke, gallbladder disease, cardiovascular disease, osteoarthritis, rheumatoid arthritis, hypercholesterolemia, stable angina, unstable angina, atherosclerosis, sleep apnea, respiratory problems, cancer, and stroke.

One aspect of the present invention provides a method for the treatment or prevention of disorders, diseases or conditions responsive to the inhibition of prolylcarboxypeptidase in a subject in need thereof which comprises administering to the subject a therapeutically or prophylactically effective amount of a compound of formula I, or a pharmaceutically acceptable salt thereof.

Another aspect of the present invention provides a method for the treatment or prevention of obesity, diabetes, an obesity related disorder or a diabetes related disorder in a subject in need thereof which comprises administering to said subject a therapeutically or prophylactically effective amount of a prolylcarboxypeptidase inhibitor of formula I. Another aspect of the present invention provides a method for the treatment or prevention of obesity in a subject in need thereof which comprises administering to the subject a therapeutically or prophylactically effective amount of a compound of formula I, or a pharmaceutically acceptable salt thereof.

Another aspect of the present invention provides a method for reducing food intake in a subject in need thereof which comprises administering to the subject a therapeutically or prophylactically effective amount of a compound of formula I, or a pharmaceutically acceptable salt thereof. Another aspect of the present invention provides a method for reducing body fat mass in a
subject in need thereof which comprises administering to the subject a therapeutically or
prophylactically effective amount of a compound of formula I, or a pharmaceutically acceptable
salt thereof. Another aspect of the present invention provides a method for losing weight in a
subject in need thereof which comprises administering to the subject a therapeutically or
prophylactically effective amount of a compound of formula I, or a pharmaceutically acceptable
salt thereof.

Another aspect of the present invention provides a method for the treatment or prevention
of an obesity-related disorder selected from the group consisting of: overeating, binge eating,
hypertension, elevated plasma insulin concentrations, insulin resistance, hyperlipidemia,
endometrial cancer, breast cancer, prostate cancer, colon cancer, kidney cancer, osteoarthritis,
obstructive sleep apnea, heart disease, abnormal heart rhythms and arrhythmias, myocardial
infarction, congestive heart failure, coronary heart disease, sudden death, stroke, polycystic ovary
disease, craniofaryngioma, metabolic syndrome, insulin resistance syndrome, sexual and
reproductive dysfunction, infertility, hypogonadism, hirsutism, obesity-related gastro-esophageal
reflux, Pickwickian syndrome, inflammation, systemic inflammation of the vasculature,
artherosclerosis, hypercholesterolemia, hyperuricaemia, lower back pain, gallbladder disease,
gout, constipation, irritable bowel syndrome, inflammatory bowel syndrome, cardiac
hypertrophy, left ventricular hypertrophy, in a subject in need thereof which comprises
administering to the subject a therapeutically or prophylactically effective amount of a compound
of formula I, or a pharmaceutically acceptable salt thereof.

Another aspect of the present invention provides a method for the treatment or prevention
diabetes, in a subject in need thereof which comprises administering to the subject a
therapeutically or prophylactically effective amount of a compound of formula I, or a
pharmaceutically acceptable salt thereof.

Another aspect of the present invention provides a method for the treatment or prevention
of a diabetes related disorder in a subject in need thereof which comprises administering to the
subject a therapeutically or prophylactically effective amount of a compound of formula I, or a
pharmaceutically acceptable salt thereof.

Another aspect of the present invention provides a method for the treatment or prevention
of a diabetes related disorder selected from the group consisting of: hyperglycemia, low glucose
tolerance, insulin resistance, obesity, lipid disorders, dyslipidemia, hyperlipidemia,
hypertriglyceridemia, hypercholesterolemia, low HDL levels, high LDL levels, atherosclerosis
and its sequelae, vascular restenosis, irritable bowel syndrome, inflammatory bowel disease,
including Crohn's disease and ulcerative colitis, other inflammatory conditions, pancreatitis,
abdominal obesity, neurodegenerative disease, retinopathy, nephropathy, neuropathy, Syndrome
X, and ovarian hyperandrogenism (polycystic ovarian syndrome), in a subject in need thereof
which comprises administering to the subject a therapeutically or prophylactically effective
amount of a compound of formula I, or a pharmaceutically acceptable salt thereof.
The present invention also relates to methods for treating or preventing obesity by
administering a compound of formula I in combination with a therapeutically or prophylactically
effective amount of another agent known to be useful to treat or prevent the condition. The
present invention also relates to methods for treating or preventing diabetes by administering a
compound of formula I in combination with a therapeutically or prophylactically effective amount
of another agent known to be useful to treat or prevent the condition. The present invention also
relates to methods for treating or preventing obesity related disorders by administering a
compound of formula I in combination with a therapeutically or prophylactically effective amount
of another agent known to be useful to treat or prevent the condition.

Yet another aspect of the present invention relates to the use of a therapeutically effective
amount of a compound of formula I, or a pharmaceutically acceptable salt or ester thereof, and a
therapeutically effective amount of at least one agent selected from the group consisting of:
simvastatin, mevastatin, ezetimibe, atorvastatin, sitagliptin, metformin, sibutramine, orlistat,
Qnexa, topiramate, phentermine, losartan, losartan with hydrochlorothiazide, or rimonabant, or a
pharmaceutically acceptable salt or ester or prodrug thereof, for the manufacture of a medicament
useful for the treatment, control, or prevention of obesity, diabetes, a diabetes related disorder, or
an obesity-related disorder in a subject in need of such treatment.

Another aspect of the present invention provides a pharmaceutical composition
comprising a compound of formula I, and a pharmaceutically acceptable carrier.

Yet another aspect of the present invention relates to the use of a compound of formula I
for the manufacture of a medicament useful for the treatment or prevention, or suppression of a
disease mediated by prolylcarboxypeptidase (PRCP) in a subject in need thereof.

Yet another aspect of the present invention relates to the use of a compound of formula I
for the manufacture of a medicament useful for the treatment or prevention, or suppression of a
disease mediated by prolylcarboxypeptidase (PRCP), wherein the disease is selected from the
group consisting of obesity, diabetes, an obesity-related disorder and a diabetes related disorder
in a subject in need thereof.

Yet another aspect of the present invention relates to the use of a therapeutically effective
amount of a compound of formula I, or a pharmaceutically acceptable salt thereof, and a
therapeutically effective amount of an agent selected from the group consisting of an insulin
sensitizer, an insulin mimetic, a sulfonylurea, an a-glucosidase inhibitor, a dipeptidyl peptidase 4
(DPP-4) inhibitor, a glucagon like peptide 1 (GLP-1) agonist, a HMG-CoA reductase inhibitor,
a serotonergic agent, a p3-adrenofeceptor agonist, a neuropeptide Y1 antagonist, a neuropeptide
Y2 agonist, a neuropeptide Y5 antagonist, a pancreatic lipase inhibitor, a cannabinoid CB1
receptor antagonist or inverse agonist, a melanin-concentrating hormone receptor antagonist, a
melanocortin 4 receptor agonist, a bombesin receptor subtype 3 agonist, a ghrelin receptor
agonist, PYY, PYY3-.36, and a NK-1 antagonist, or a pharmaceutically acceptable salt thereof,
for the manufacture of a medicament useful for the treatment, control, or prevention of obesity,
diabetes or an obesity-related disorder in a subject in need of such treatment. Yet another aspect of the present invention relates to the use of a therapeutically effective amount of a compound of formula I, and pharmaceutically acceptable salts and esters thereof, and a therapeutically effective amount of an agent selected from the group consisting of an insulin sensitizing agent, an insulin mimetic, a sulfonylurea, an α-glucosidase inhibitor, a dipeptidyl peptidase IV inhibitor, a glucagon-like peptide 1 agonist, a HMG-CoA reductase inhibitor, a serotonergic agent, a β3-adrenoreceptor agonist, an neuropeptide Y1 antagonist, an neuropeptide Y2 agonist, a neuropeptide Y5 antagonist, a pancreatic lipase inhibitor, a cannabinoid CB1 receptor antagonist or inverse agonist, a melanin-concentrating hormone receptor antagonist, a melanocortin 4 receptor agonist, a bombesin receptor subtype 3 agonist, a ghrelin receptor antagonist, PYY, PYY3-36, and a NK-1 antagonist, or a pharmaceutically acceptable salt thereof, for the manufacture of a medicament for treatment or prevention of obesity, diabetes, an obesity related disorder or a diabetes related disorder which comprises an effective amount of a the compound of formula I, and an effective amount of the agent, together or separately. Yet another aspect of the present invention relates to a product containing a therapeutically effective amount of a compound of formula I, or a pharmaceutically acceptable salt thereof; and and a therapeutically effective amount of an agent selected from the group consisting of an insulin sensitizing agent, an insulin mimetic, a sulfonylurea, an α-glucosidase inhibitor, a HMG-CoA reductase inhibitor, a serotonergic agent, a β3-adrenoreceptor agonist, an neuropeptide Y1 antagonist, an neuropeptide Y2 agonist, a neuropeptide Y5 antagonist, a pancreatic lipase inhibitor, a cannabinoid CB1 receptor antagonist or inverse agonist, a melanocortin 4 receptor agonist, a melanin-concentrating hormone receptor antagonist, a bombesin receptor subtype 3 agonist, a ghrelin receptor antagonist, ρYY, PYY3-36, and a NK-1 antagonist, or a pharmaceutically acceptable salt thereof, as a combined preparation for simultaneous, separate or sequential use in obesity, diabetes, an obesity-related disorder or a diabetes related disorder.

The compounds of formula I can be provided in kit. Such a kit typically contains an active compound in dosage forms for administration. A dosage form contains a sufficient amount of active compound such that a beneficial effect can be obtained when administered to a patient during regular intervals, such as 1, 2, 3, 4, 5 or 6 times a day, during the course of 1 or more days. Preferably, a kit contains instructions indicating the use of the dosage form for weight reduction (e.g., to treat obesity) and the amount of dosage form to be taken over a specified time period.

Compounds of formula I are inhibitors of prolylcarboxypeptidase (FRCP) and as such are useful in the treatment, control or prevention of diseases, disorders or conditions responsive to the modulation of prolylcarboxypeptidase (PRCP). Such diseases, disorders or conditions include, but are not limited to, abnormal metabolism, obesity, diabetes, metabolic syndrome, obesity related disorders, diabetes related disorders, hypertension, dyslipidemia, stroke, gallbladder disease, cardiovascular disease, osteoarthritis, rheumatoid arthritis,
hypercholesterolemia, stable angina, unstable angina, arteriosclerosis, sleep apnea, respiratory
problems, cancer, and stroke. Such diseases, conditions and disorders also include non-obese
overweight conditions and normal weight conditions where weight control or management is
desired in order to prevent an obese or overweight condition from developing, or to maintain a
healthy weight.

The compounds of formula I, and compositions thereof, are useful for the treatment or
prevention of disorders associated with excessive food intake, such as obesity and obesity-related
disorders. The obesity herein may be due to any cause, whether genetic or environmental.

The obesity-related disorders herein are associated with, caused by, or result from obesity.
Examples of obesity-related disorders include overeating, binge eating, bulimia nervosa,
hypertension, type 2 diabetes, elevated plasma insulin concentrations, hyperinsulinemia, insulin
resistance, glucose intolerance, dyslipidemia, hyperlipidemia, endometrial cancer, breast cancer,
prostate cancer, kidney cancer, colon cancer, osteoarthritis, obstructive sleep apnea,
cholelithiasis, cholecystitis, gallstones, gout, gallbladder disease, abnormal heart rhythms and
arrythmias, myocardial infarction, congestive heart failure, coronary heart disease, angina
pectoris, sudden death, stroke, metabolic syndrome, psychological disorders (depression, eating
disorders, distorted bodyweight, and low self esteem), and other pathological conditions showing
reduced metabolic activity or a decrease in resting energy expenditure as a percentage of total fat-
free mass, e.g. children with acute lymphoblastic leukemia. Further examples of obesity-related
disorders are sexual and reproductive dysfunction, such as polycystic ovary disease, infertility,
hypogonadism in males and hirsutism in females, gastrointestinal motility disorders, such as
obesity-related gastroesophageal reflux, respiratory disorders, such as obesity-hypoventilation
syndrome (Pickwickian syndrome), cardiovascular disorders, inflammation, such as systemic
inflammation of the vasculature, arteriosclerosis, hypercholesterolemia, hyperuricaemia, lower
back pain, gallbladder disease, gout, and kidney cancer. Additionally, the present compounds are
useful in the treatment of any condition in which it is desirable to lose weight or to reduce food
intake. Additionally, the present compounds are useful in the treatment of any condition in
which it is desirable to enhance cognition and memory, such as Alzheimer's Disease. The
compositions of the present invention are also useful for reducing the risk of secondary outcomes
of obesity, such as reducing the risk of left ventricular hypertrophy. Therefore, the present
invention provides methods of treatment or prevention of such diseases, conditions and/or
disorders modulated by prolylcarboxypeptidase (PrCP) in an animal which comprises
administering to the animal in need of such treatment a compound of formula I, in particular a
therapeutically or prophylactically effective amount thereof.

The term "inhibitor" as used herein means a composition of matter which when
administered to a mammal, such as a human, inhibits the biological activity attributable to the
level or presence of an endogenous compound in the mammal. Inhibition of PrCP includes, but
is not limited to, inhibiting the biological activity of the PrCP enzyme or molecule.
The term "subject" means a mammal. One embodiment of the term "mammal" is a "human," said human being either male or female. The instant compounds are also useful for treating or preventing obesity and obesity related disorders in cats and dogs. As such, the term "mammal" includes companion animals such as cats and dogs. The term "mammal in need thereof" refers to a mammal who is in need of treatment or prophylaxis as determined by a researcher, veterinarian, medical doctor or other clinician.

The term "composition", as in pharmaceutical composition, is intended to encompass a product comprising the active ingredient(s), and the inert ingredient(s) that make up the carrier, as well as any product which results, directly or indirectly, from combination, complexation or aggregation of any two or more of the ingredients, or from dissociation of one or more of the ingredients, or from other types of reactions or interactions of one or more of the ingredients. Accordingly, the pharmaceutical compositions of the present invention encompass any composition made by admixing a compound of the present invention and a pharmaceutically acceptable carrier.

The term "metabolic syndrome", also known as syndrome X, is defined in the Third Report of the National Cholesterol Education Program Expert Panel on Detection, Evaluation and Treatment of High Blood Cholesterol in Adults (ATP-III). E.S. Ford et al., JAMA, vol. 287 (3), Jan. 16, 2002, pp 356-359. Briefly, a person is defined as having metabolic syndrome if the person has three or more of the following symptoms: abdominal obesity, hypertriglyceridemia, low HDL cholesterol, high blood pressure, and high fasting plasma glucose. The criteria for these are defined in ATP-III.

The term "diabetes," as used herein, includes both insulin-dependent diabetes mellitus (i.e., IDDM, also known as type I diabetes) and non-insulin-dependent diabetes mellitus (i.e., NIDDM, also known as Type II diabetes). Type I diabetes, or insulin-dependent diabetes, is the result of an absolute deficiency of insulin, the hormone which regulates glucose utilization. Type II diabetes, or insulin-independent diabetes (i.e., non-insulin-dependent diabetes mellitus), often occurs in the face of normal, or even elevated levels of insulin and appears to be the result of the inability of tissues to respond appropriately to insulin. Most of the Type II diabetics are also obese. The compositions of the present invention are useful for treating both Type I and Type II diabetes. The compositions are especially effective for treating Type II diabetes. The compounds or combinations of the present invention are also useful for treating and/or preventing gestational diabetes mellitus.

Diabetes is characterized by a fasting plasma glucose level of greater than or equal to 126 mg/dl. A diabetic subject has a fasting plasma glucose level of greater than or equal to 126 mg/dl. Prediabetes is characterized by an impaired fasting plasma glucose (FPG) level of greater than or equal to 110 mg/dl and less than 126 mg/dl; or impaired glucose tolerance; or insulin resistance. A prediabetic subject is a subject with impaired fasting glucose (a fasting plasma glucose (FPG) level of greater than or equal to 110 mg/dl and less than 126 mg/dl); or impaired
glucose tolerance (a 2 hour plasma glucose level of $\geq 140$ mg/dl and $<200$ mg/dl); or insulin resistance, resulting in an increased risk of developing diabetes.

"Diabetes related disorders" are diseases, disorders and conditions that are related to Type 2 diabetes, and therefore may be treated, controlled or in some cases prevented, by treatment with the compounds of this invention: (1) hyperglycemia, (2) low glucose tolerance, (3) insulin resistance, (4) obesity, (5) lipid disorders, (6) dyslipidemia, (7) hyperlipidemia, (8) hypertriglyceridemia, (9) hypercholesterolemia, (10) low HDL levels, (11) high LDL levels, (12) atherosclerosis and its sequela, (13) vascular restenosis, (14) irritable bowel syndrome, (15) inflammatory bowel disease, including Crohn's disease and ulcerative colitis, (16) other inflammatory conditions, (17) pancreatitis, (18) abdominal obesity, (19) neurodegenerative disease, (20) retinopathy, (21) nephropathy, (22) neuropathy, (23) Syndrome X, (24) ovarian hyperandrogenism (polycystic ovarian syndrome), and other disorders where insulin resistance is a component. In Syndrome X, also known as Metabolic Syndrome, obesity is thought to promote insulin resistance, diabetes, dyslipidemia, hypertension, and increased cardiovascular risk. Therefore, inhibitors of prolylcarboxypeptidase (FRCP) may also be useful to treat hypertension associated with this condition.

Treatment of diabetes mellitus refers to the administration of a compound or combination of the present invention to treat diabetes. One outcome of treatment may be decreasing the glucose level in a subject with elevated glucose levels. Another outcome of treatment may be improving glycemic control. Another outcome of treatment may be decreasing insulin levels in a subject with elevated insulin levels. Another outcome of the treatment of diabetes is to reduce an increased plasma glucose concentration. Another outcome of the treatment of diabetes is to reduce an increased insulin concentration. Still another outcome of the treatment of diabetes is to reduce an increased blood triglyceride concentration. Still another outcome of the treatment of diabetes is to increase insulin sensitivity. Still another outcome of the treatment of diabetes may be enhancing glucose tolerance in a subject with glucose intolerance. Still another outcome of the treatment of diabetes is to reduce insulin resistance. Another outcome of the treatment of diabetes is to lower plasma insulin levels. Still another outcome of treatment of diabetes is an improvement in glycemic control, particularly in type 2 diabetes.

Prevention of diabetes mellitus, in particular diabetes associated with obesity, refers to the administration of a compound or combination of the present invention to prevent or treat the onset of diabetes in a subject in need thereof. A subject in need of preventing diabetes in a prediabetic subject.

"Obesity" is a condition in which there is an excess of body fat. The operational definition of obesity is based on the Body Mass Index (BMI), which is calculated as body weight per height in meters squared (kg/m$^2$). "Obesity" refers to a condition whereby an otherwise healthy subject has a Body Mass Index (BMI) greater than or equal to 30 kg/m$^2$, or a condition whereby a subject with at least one co-morbidity has a BMI greater than or equal to 27 kg/m$^2$. 

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An "obese subject" is an otherwise healthy subject with a Body Mass Index (BMI) greater than or equal to 30 kg/m² or a subject with at least one co-morbidity with a BMI greater than or equal to 27 kg/m². A "subject at risk of obesity" is an otherwise healthy subject with a BMI of 25 kg/m² to less than 30 kg/m² or a subject with at least one co-morbidity with a BMI of 25 kg/m² to less than 27 kg/m².

The increased risks associated with obesity occur at a lower Body Mass Index (BMI) in Asians. In Asian countries, including Japan, "obesity" refers to a condition whereby a subject with at least one obesity-induced or obesity-related co-morbidity, that requires weight reduction or that would be improved by weight reduction, has a BMI greater than or equal to 25 kg/m². In Asian countries, including Japan, an "obese subject" refers to a subject with at least one obesity-induced or obesity-related co-morbidity that requires weight reduction or that would be improved by weight reduction, with a BMI greater than or equal to 25 kg/m². In Asia-Pacific, a "subject at risk of obesity" is a subject with a BMI of greater than 23 kg/m² to less than 25 kg/m².

As used herein, the term "obesity" is meant to encompass all of the above definitions of obesity.

Obesity-induced or obesity-related co-morbidities include, but are not limited to, diabetes, non-insulin dependent diabetes mellitus - type II (2), impaired glucose tolerance, impaired fasting glucose, insulin resistance syndrome, dyslipidemia, hypertension, hyperuricacidemia, gout, coronary artery disease, myocardial infarction, angina pectoris, sleep apnea syndrome, Pickwickian syndrome, fatty liver; cerebral infarction, cerebral thrombosis, transient ischemic attack, orthopedic disorders, arthritis deformans, lumbodynia, emmeniopathy, and infertility. In particular, co-morbidities include: hypertension, hyperlipidemia, dyslipidemia, glucose intolerance, cardiovascular disease, sleep apnea, diabetes mellitus, and other obesity-related conditions.

Treatment of obesity and obesity-related disorders refers to the administration of the compounds or combinations of the present invention to reduce or maintain the body weight of an obese subject. One outcome of treatment may be reducing the body weight of an obese subject relative to that subject's body weight immediately before the administration of the compounds or combinations of the present invention. Another outcome of treatment may be preventing body weight regain of body weight previously lost as a result of diet, exercise, or pharmacotherapy. Another outcome of treatment may be decreasing the occurrence of and/or the severity of obesity-related diseases. The treatment may suitably result in a reduction in food or calorie intake by the subject, including a reduction in total food intake, or a reduction of intake of specific components of the diet such as carbohydrates or fats; and/or the inhibition of nutrient absorption; and in weight reduction in subjects in need thereof. The treatment may also result in an alteration of metabolic rate, such as an increase in metabolic rate, rather than or in addition to an inhibition of the reduction of metabolic rate; and/or in minimization of the metabolic resistance that normally results from weight loss.
Prevention of obesity and obesity-related disorders refers to the administration of the compounds or combinations of the present invention to reduce or maintain the body weight of a subject at risk of obesity. One outcome of prevention may be reducing the body weight of a subject at risk of obesity relative to that subject's body weight immediately before the administration of the compounds or combinations of the present invention. Another outcome of prevention may be preventing body weight regain of body weight previously lost as a result of diet, exercise, or pharmacotherapy. Another outcome of prevention may be preventing obesity from occurring if the treatment is administered prior to the onset of obesity in a subject at risk of obesity. Another outcome of prevention may be decreasing the occurrence and/or severity of obesity-related disorders if the treatment is administered prior to the onset of obesity in a subject at risk of obesity. Moreover, if treatment is commenced in already obese subjects, such treatment may prevent the occurrence, progression or severity of obesity-related disorders, such as, but not limited to, arteriosclerosis, Type II diabetes, polycystic ovary disease, cardiovascular diseases, osteoarthritis, hypertension, dyslipidemia, insulin resistance, hypercholesterolemia, hypertriglyceridemia, and cholelithiasis.

The terms "administration of" and or "administering" a compound should be understood to mean providing a compound of the invention or a prodrug of a compound of the invention to a subject in need of treatment. The administration of the compounds of the present invention in order to practice the present methods of therapy is carried out by administering a therapeutically effective amount of the compound to a subject in need of such treatment or prophylaxis. The need for a prophylactic administration according to the methods of the present invention is determined via the use of well known risk factors.

The term "therapeutically effective amount" as used herein means the amount of the active compound that will elicit the biological or medical response in a tissue, system, subject, mammal, or human that is being sought by the researcher, veterinarian, medical doctor or other clinician, which includes alleviation of the symptoms of the disorder being treated. The novel methods of treatment of this invention are for disorders known to those skilled in the art. The term "prophylactically effective amount" as used herein means the amount of the active compound that will elicit the biological or medical response in a tissue, system, subject, mammal, or human that is being sought by the researcher, veterinarian, medical doctor or other clinician, to prevent the onset of the disorder in subjects as risk for obesity or the disorder. The therapeutically or prophylactically effective amount, or dosage, of an individual compound is determined, in the final analysis, by the physician in charge of the case, but depends on factors such as the exact disease to be treated, the severity of the disease and other diseases or conditions from which the patient suffers, the chosen route of administration, other drugs and treatments which the patient may concomitantly require, and other factors in the physician's judgement.

Administration and Dose Ranges

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Any suitable route of administration may be employed for providing a subject or mammal, especially a human with an effective dosage of a compound of the present invention. For example, oral, rectal, topical, parenteral, ocular, pulmonary, nasal, and the like may be employed. Dosage forms include tablets, troches, dispersions, suspensions, solutions, capsules, creams, ointments, aerosols, and the like. Preferably compounds of Formula I are administered orally or topically.

The effective dosage of active ingredient employed may vary depending on the particular compound employed, the mode of administration, the condition being treated and the severity of the condition being treated. Such dosage may be ascertained readily by a person skilled in the art.

Generally satisfactory results are obtained when the compounds of formula I are administered at a daily dosage of from about 0.001 milligram to about 50 milligrams per kilogram of animal body weight, preferably given in a single dose or in divided doses two to six times a day, or in sustained release form. In the case of a 70 kg adult human, the total daily dose will generally be from about 0.07 milligrams to about 3500 milligrams. This dosage regimen may be adjusted to provide the optimal therapeutic response.

In the case where an oral composition is employed, a suitable dosage range is, e.g. from about 0.01 mg to about 1500 mg of a compound of Formula I per day, preferably from about 0.1 mg to about 600 mg per day, more preferably from about 0.1 mg to about 100 mg per day. For oral administration, the compositions are preferably provided in the form of tablets containing from 0.01 to 1,000 mg, preferably 0.01, 0.05, 0.1, 0.5, 1, 2.5, 5, 10, 15, 20, 25, 30, 40, 50, 100, 250, 500, 600, 750, 1000, 1250 or 1500 milligrams of the active ingredient for the symptomatic adjustment of the dosage to the patient to be treated.

For use where a composition for intranasal administration is employed, intranasal formulations for intranasal administration comprising 0.001 - 10% by weight solutions or suspensions of the compounds of formula I in an acceptable intranasal formulation may be used.

For use where a composition for intravenous administration is employed, a suitable dosage range is from about 0.001 mg to about 50 mg, preferably from 0.01 mg to about 50 mg, more preferably 0.1 mg to 10 mg, of a compound of formula I per kg of body weight per day. This dosage regimen may be adjusted to provide the optimal therapeutic response. It may be necessary to use dosages outside these limits in some cases.

For the treatment of diseases of the eye, ophthalmic preparations for ocular administration comprising 0.001-1% by weight solutions or suspensions of the compounds of formula I in an acceptable ophthalmic formulation may be used.

The magnitude of prophylactic or therapeutic dosage of the compounds of the present invention will, of course, vary depending on the particular compound employed, the mode of administration, the condition being treated and the severity of the condition being treated. It will
also vary according to the age, weight and response of the individual patient. Such dosage may be ascertained readily by a person skilled in the art.

**Combination Therapy**

Compounds of Formula I may be used in combination with other drugs that may also be useful in the treatment or amelioration of the diseases or conditions for which compounds of Formula I are useful. Such other drugs may be administered, by a route and in an amount commonly used therefor, contemporaneously or sequentially with a compound of Formula I. In the treatment of patients who have Type 2 diabetes, insulin resistance, obesity, metabolic syndrome, and co-morbidities that accompany these diseases, more than one drug is commonly administered. The compounds of this invention may generally be administered to a patient who is already taking one or more other drugs for these conditions. Often the compounds will be administered to a patient who is already being treated with one or more antidiabetic compound, such as metformin, sulfonylureas, and/or PPAR agonists, when the patient's glycemic levels are not adequately responding to treatment.

When a compound of Formula I is used contemporaneously with one or more other drugs, a pharmaceutical composition in unit dosage form containing such other drugs and the compound of Formula I is preferred. However, the combination therapy also includes therapies in which the compound of Formula I and one or more other drugs are administered on different overlapping schedules. It is also contemplated that when used in combination with one or more other active ingredients, the compound of the present invention and the other active ingredients may be used in lower doses than when each is used singly. Accordingly, the pharmaceutical compositions of the present invention include those that contain one or more other active ingredients, in addition to a compound of Formula I.

Examples of other active ingredients that may be administered in combination with a compound of Formula I, and either administered separately or in the same pharmaceutical composition, include, but are not limited to:

- (a) PPAR gamma agonists and partial agonists, including both glitazones and non-glitazones (e.g. troglitazone, pioglitazone, englitazone, MCC-555, rosiglitazone, balaglitazone, netoglitazone, T-131, LY-300512, LY-818, and compounds disclosed in WO02/08188, WO2004/020408, and WO2004/020409.
- (b) biguanides, such as metformin and phenformin;
- (c) protein tyrosine phosphatase-IB (PTP-1B) inhibitors;
- (d) dipeptidyl peptidase-IV (DPP-4) inhibitors, such as sitagliptin, saxagliptin, vildagliptin, and alogliptin;
- (e) insulin or insulin mimetics;
- (f) sulfonylureas such as tolbutamide, glimepiride, glipizide, and related materials;
- (g) oglucosidase inhibitors (such as acarbose);
agents which improve a patient's lipid profile, such as (i) HMG-CoA reductase inhibitors (lovastatin, simvastatin, rosuvastatin, pravastatin, fluvastatin, atorvastatin, rivastatin, itavastatin, ZD-4522 and other statins), (ii) bile acid sequestrants (cholestryamine, colestipol, and dialkylaminoalkyl derivatives of a cross-linked dextran), (iii) niacin receptor agonists, nicotinyl alcohol, nicotinic acid, or a salt thereof, (iv) PPAR agonists, such as fenofibric acid derivatives (gemfibrozil, clofibrate, fenofibrate and bezafibrate), (v) cholesterol absorption inhibitors, such as ezetimibe, (vi) acyl CoA:cholesterol acyltransferase (ACAT) inhibitors, such as avasimibe, (vii) CETP inhibitors, such as torcetrapib, and (viii) phenolic antioxidants, such as probucol;

(i) PPARα/γ dual agonists, such as muraglitazar, tesaglitazar, farglitazar, arid JT-501;

(0) PPARδ agonists, such as those disclosed in W097/28149;

(k) anti-obesity compounds, such as fenfluramine, dexfenfluramine, phentiramline, subitramine, orlistat, neuropeptide Y Y5 inhibitors, MC4R agonists, cannabinoid receptor 1 (CB-1) antagonists/inverse agonists (e.g., rimonabant and tatanabant), and β3 adrenergic receptor agonists;

(i) ileal bile acid transporter inhibitors;

(m) agents intended for use in inflammatory conditions, such as aspirin, non-steroidal anti-inflammatory drugs, glucocorticoids, azulfidine, and cyclooxygenase-2 (COX-2) selective inhibitors;

(n) glucagon receptor antagonists;

(o) GLP-1;

(P) GIP-1;

(q) GLP-1 analogs and derivatives, such as exendins, (e.g., exenatide and liraglutide);

(r) 11β-hydroxysteroid dehydrogenase-1 (HSD-1) inhibitors;

(s) inhibitors of cholesteryl ester transfer protein (CETP), such as torcetrapib;

(t) SSTR3 antagonists;

(u) other SSTR5 antagonists;

(v) acetyl CoA carboxylase-1 and/or -2 inhibitors;

(w) AMPK activators;

(x) agonists of GPR-119;

(y) glucokinase agonists; and

(z) FGF-21 agonists.

The above combinations include combinations of a compound of the present invention not only with one other active compound, but also with two or more other active compounds. Non-limiting examples include combinations of compounds having Formula I with two or more active compounds selected from biguanides, sulfonylureas, HMG-CoA reductase inhibitors,
other PPAR agonists, PTP-1B inhibitors, DPP-4 inhibitors, and cannabinoid receptor 1 (CB1) inverse agonists/antagonists.

Antiobesity compounds that can be combined with compounds described herein include fenfluramine, dexfenfluramine, phentermine, sibutramine, orlistat, neuropeptide Yi or Y5 antagonists, cannabinoid CB1 receptor antagonists or inverse agonists, melanocortin receptor agonists, in particular, melanocortin-4 receptor agonists, ghrelin antagonists, bombesin receptor agonists, and melanin-concentrating hormone (MCH) receptor antagonists. For a review of antiobesity compounds that can be combined with compounds described herein, see S. Chaki et al., "Recent advances in feeding suppressing agents: potential therapeutic strategy for the treatment of obesity," Expert Opin. Ther. Patents, 11: 1677-1692 (2001); D. Spanswick and K. Lee, "Emerging antiobesity drugs," Expert Opin. Emerging Drugs, 8: 217-237 (2003); and J.A. Fernandez-Lopez, et al, "Pharmacological Approaches for the Treatment of Obesity," Drugs, 62: 915-944 (2002).

Neuropeptide Y5 antagonists that can be combined with compounds described herein include those disclosed in U.S. Patent No. 6,335,345 (1 January 2002) and WO 01/14376 (1 March 2001); and specific compounds identified as GW 59884A; GW 569180A; LY366377; and CGP-71683A.

Cannabinoid CB1 receptor antagonists that can be combined with compounds described herein include those disclosed in PCT Publication WO 03/007887; U.S. Patent No. 5,624,941, such as rimonabant; PCT Publication WO 02/076949, such as SLV-319; U.S. Patent No. 6,028,084; PCT Publication WO 98/41519; PCT Publication WO 00/10968; PCT Publication WO 99/02499; U.S. Patent No. 5,532,237; U.S. Patent No. 5,292,736; PCT Publication WO 03/086288; PCT Publication WO 03/087037; PCT Publication WO 04/048317; PCT Publication WO 04/057660; PCT Publication WO 03/007887; PCT Publication WO 03/063781; PCT Publication WO 03/075660; PCT Publication WO 03/077847; PCT Publication WO 03/082190; PCT Publication WO 03/082191; PCT Publication WO 03/087037; PCT Publication WO 03/086288; PCT Publication WO 04/012671; PCT Publication WO 04/029204; PCT Publication WO 04/040040; PCT Publication WO 01/64632; PCT Publication WO 01/64633; and PCT Publication WO 01/64634.

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WO 02/059095, WO 02/059107, WO 02/059108, WO 02/059117, WO 02/085925, WO 03/004480, WO 03/009850, WO 03/013571, WO 03/01410, WO 03/053927, WO 03/061660, WO 03/066597, WO 03/094918, WO 03/099818, WO 04/037797, WO 04/048345, WO 02/018327, WO 02/080896, WO 02/081443, WO 03/066587, WO 03/066597, WO 03/099818, WO 02/062766, WO 03/000663, WO 03/000666, WO 03/003977, WO 03/040107, WO 03/040117, WO 03/040118, WO 03/013509, WO 03/057671, WO 02/079753, WO 02/092566, WO 03/-093234, WO 03/095474, and WO 03/104761,

Specific compounds of use in combination with a compound of the present invention include: simvastatin, mevastatin, ezetimibe, atorvastatin, sitagliptin, metformin, sibutramine, orlistat, Qnexa, topiramate, naltrexone, bupropion, phentermine, and losartan, losartan with hydrochlorothiazide.


The instant invention also includes administration of a single pharmaceutical dosage formulation which contains both a compound of formula I in combination with a second active ingredient, as well as administration of each active agent in its own separate pharmaceutical dosage formulation. Where separate dosage formulations are used, the individual components of the composition can be administered at essentially the same time, i.e., concurrently, or at separately staggered times, i.e. sequentially prior to or subsequent to the administration of the other component of the composition. The instant invention is therefore to be understood to include all such regimes of simultaneous or alternating treatment, and the terms "administration" and "administering" are to be interpreted accordingly. Administration in these various ways are suitable for the present compositions as long as the beneficial pharmaceutical effect of the combination of the compound of formula I and the second active ingredient is realized by the patient at substantially the same time. Such beneficial effect is preferably achieved when the target blood level concentrations of each active ingredient are maintained at substantially the same time. It is preferred that the combination of the compound of formula I and the second active ingredient be co-administered concurrently on a once-a-day dosing schedule; however, varying dosing schedules, such as the compound of formula I once a day and the second active ingredient once, twice or more times per day or the compound of formula I three times a day and the second active ingredient once, twice or more times per day, is also encompassed herein. A single oral dosage formulation comprised of both a compound of formula I and a second active
ingredient is preferred. A single dosage formulation will provide convenience for the patient, which is an important consideration especially for patients with diabetes or obese patients who may be in need of multiple medications.

The compounds in the combinations of the present invention may be administered separately, therefore the invention also relates to combining separate pharmaceutical compositions into a kit form. The kit, according to this invention, comprises two separate pharmaceutical compositions: a first unit dosage form comprising a prophylactically or therapeutically effective amount of the compound of formula I, or a pharmaceutically acceptable salt or ester thereof, and a pharmaceutically acceptable carrier or diluent in a first unit dosage form, and a second unit dosage form comprising a prophylactically or therapeutically effective amount of the second active ingredient or drug, or a pharmaceutically acceptable salt or ester thereof, and a pharmaceutically acceptable carrier or diluent in a second unit dosage form. In one embodiment, the kit further comprises a container. Such kits are especially suited for the delivery of solid oral forms such as tablets or capsules. Such a kit preferably includes a number of unit dosages. Such kits can include a card having the dosages oriented in the order of their intended use. An example of such a kit is a "blister pack". Blister packs are well known in the packaging industry and are widely used for packaging pharmaceutical unit dosage forms. If desired, a memory aid can be provided, for example in the form of numbers, letters, or other markings or with a calendar insert, designating the days or time in the treatment schedule in which the dosages can be administered.

Pharmaceutical Compositions

Another aspect of the present invention provides pharmaceutical compositions which comprise a compound of formula I, as an active ingredient or a pharmaceutically acceptable salt thereof, and may also contain a pharmaceutically acceptable carrier and optionally other therapeutic ingredients. The term "pharmaceutically acceptable salts" refers to salts prepared from pharmaceutically acceptable non-toxic bases or acids including inorganic bases or acids and organic bases or acids.

The compositions include compositions suitable for oral, rectal, topical, parenteral (including subcutaneous, intramuscular, and intravenous), ocular (ophthalmic), pulmonary (nasal or buccal inhalation), or nasal administration, although the most suitable route in any given case will depend on the nature and severity of the conditions being treated and on the nature of the active ingredient. They may be conveniently presented in unit dosage form and prepared by any of the methods well-known in the art of pharmacy.

In practical use, the compounds of formula I can be combined as the active ingredient in intimate admixture with a pharmaceutical carrier according to conventional pharmaceutical compounding techniques. The carrier may take a wide variety of forms depending on the form of preparation desired for administration, e.g., oral or parenteral (including intravenous). In preparing the compositions for oral dosage form, any of the usual pharmaceutical media may be
employed, such as, for example, water, glycols, oils, alcohols, flavoring agents, preservatives, coloring agents and the like in the case of oral liquid preparations, such as, for example, suspensions, elixirs and solutions; or carriers such as starches, sugars, microcrystalline cellulose, diluents, granulating agents, lubricants, binders, disintegrating agents and the like in the case of oral solid preparations such as, for example, powders, hard and soft capsules and tablets, with the solid oral preparations being preferred over the liquid preparations.

Because of their ease of administration, tablets and capsules represent the typical oral dosage unit form, in which case solid pharmaceutical carriers are typically employed. If desired, tablets may be coated by standard aqueous or nonaqueous techniques. Such compositions and preparations should contain at least 0.1 percent of active compound. The percentage of active compound in these compositions may, of course, be varied and may conveniently be between about 2 percent to about 60 percent of the weight of the unit. The amount of active compound in such therapeutically useful compositions is such that an effective dosage will be obtained. The active compounds can also be administered intranasally as, for example, liquid drops or spray.

The tablets, pills, capsules, and the like may also contain a binder such as gum tragacanth, acacia, corn starch or gelatin; excipients such as dicalcium phosphate; a disintegrating agent such as corn starch, potato starch, alginic acid; a lubricant such as magnesium stearate; and a sweetening agent such as sucrose, lactose or saccharin. When a dosage unit form is a capsule, it may contain, in addition to materials of the above type, a liquid carrier such as a fatty oil.

Various other materials may be present as coatings or to modify the physical form of the dosage unit. For instance, tablets may be coated with shellac, sugar or both. A syrup or elixir may contain, in addition to the active ingredient, sucrose as a sweetening agent, methyl and propylparabens as preservatives, a dye and a flavoring such as cherry or orange flavor.

Compounds of formula I may also be administered parenterally. Solutions or suspensions of these active compounds can be prepared in water suitably mixed with a surfactant such as hydroxy-propylcellulose. Dispersions can also be prepared in glycerol, liquid polyethylene glycols and mixtures thereof in oils. Under ordinary conditions of storage and use, these preparations contain a preservative to prevent the growth of microorganisms.

The pharmaceutical forms suitable for injectable use include sterile aqueous solutions or dispersions and sterile powders for the extemporaneous preparation of sterile injectable solutions or dispersions. In all cases, the form must be sterile and must be fluid to the extent that easy syringability exists. It must be stable under the conditions of manufacture and storage and must be preserved against the contaminating action of microorganisms such as bacteria and fungi. The carrier can be a solvent or dispersion medium containing, for example, water, ethanol, polyl (e.g. glycerol, propylene glycol and liquid polyethylene glycol), suitable mixtures thereof, and vegetable oils.

Preparation of Compounds of the Invention
The compounds of formula I of the present invention can be prepared according to the procedures of the following Schemes and Examples, using appropriate materials and are further exemplified by the following specific examples. Moreover, by utilizing the procedures described herein, one of ordinary skill in the art can readily prepare additional compounds of the present invention claimed herein. The compounds illustrated in the examples are not, however, to be construed as forming the only genus that is considered as the invention. The Examples further illustrate details for the preparation of the compounds of the present invention. Those skilled in the art will readily understand that known variations of the conditions and processes of the following preparative procedures can be used to prepare these compounds. The instant compounds are generally isolated in the form of their pharmaceutically acceptable salts, such as those described previously hereinabove. The free amine bases corresponding to the isolated salts can be generated by neutralization with a suitable base, such as aqueous sodium hydrogencarbonate, sodium carbonate, sodium hydroxide, and potassium hydroxide, and extraction of the liberated amine free base into an organic solvent followed by evaporation. The amine free base isolated in this manner can be further converted into another pharmaceutically acceptable salt by dissolution in an organic solvent followed by addition of the appropriate acid and subsequent evaporation, precipitation, or crystallization. All temperatures are degrees Celsius unless otherwise noted. All temperatures are degrees Celsius unless otherwise noted.

Mass spectra (MS) were measured by electrospray ion-mass spectroscopy (ESMS).

The phrase "standard peptide coupling reaction conditions" means coupling a carboxylic acid with an amine using an acid activating agent such as EDC, DCC, and BOP in an inert solvent such as dichloromethane in the presence of a catalyst such as HOBT. The use of protecting groups for the amine and carboxylic acid functionalities to facilitate the desired reaction and minimize undesired reactions is well documented. Conditions required to remove protecting groups are found in standard textbooks such as Greene, T, and Wuts, P. G. M., Protective Groups in Organic Synthesis, John Wiley & Sons, Inc., New York, NY, 1991. CBZ and BOC are commonly used protecting groups in organic synthesis, and their removal conditions are known to those skilled in the art. For example, CBZ may be removed by catalytic hydrogenation in the presence of a noble metal or its oxide such as palladium on activated carbon in a protic solvent such as methanol or ethanol. In cases where catalytic hydrogenation is contraindicated due to the presence of other potentially reactive functionalities, removal of CBZ groups can also be achieved by treatment with a solution of hydrogen bromide in acetic acid or by treatment with a mixture of TFA and dimethylsulfide. Removal of BOC protecting groups is carried out with a strong acid, such as trifluoroacetic acid, hydrochloric acid, or hydrogen chloride gas, in a solvent such as methylene chloride, methanol, or ethyl acetate.

The LC/MS analyses were performed using a MICROMASS ZQS mass spectrometer coupled to an AGILENT 1100 Series HPLC utilizing an XTerra®MS, C18, 3.5 mm, 2.1 x 20
mm column eluting at 1.5 mL/min with a solvent gradient of 5 to 95% B over 0.75 min, followed by 95-98% B over 0.5 min: solvent A = 0.05% TFA in water; solvent B = 0.05% TFA in acetonitrile. 1H-NMR spectra were obtained on a 500 MHz VARIAN Spectrometer in CDCl₃ or CD₃CN or as indicated and chemical shifts are reported as δ using the solvent peak as reference and coupling constants are reported in hertz (Hz).

Abbreviations Used in the Description of the Preparation of the Compounds of the Present Invention: ACN is acetonitrile; Ac₂O is acetic anhydride; Ac is acetyl; AcOH is acetic acid; aq is aqueous; Boc and BOC is tert-butyloxycarbonyl; (Boc)₂O and (BOC)₂O is di-tert-butyl dicarbonate; Bn is benzyl; BuLi is butyl lithium; brine is saturated aqueous sodium chloride solution; CDI is 1,1'-carbonyldimidazole; Celite™ is diatomaceous earth; CₐH₂ is carbon dioxide; CO is carbon monoxide; D is deuterium; DCM or CH₂CCL₂ is dichloromethane; DBU is 1,8-diazabicyclo[5.4.0]undec-7-ene; DIPEA and DIEA is N,N-diisopropylethylamine; DMAP is 4-AN,N₂-diethylaminopyridine; DME is 1,2-dimethoxyethane; DMF is N,N-dimethylformamide; DMSO is dimethyl sulfoxide; dpff is 1,1'-bis(diphenylphosphino)ferrocene; EDC is 1-(dimethylaminomethyl)-3-ethylcarbodiimide hydrochloride; EtOAc is ethyl acetate; Et (et) is ethyl; EA and EtOAc is ethyl acetate; equiv is equivalent(s); ESI is electrospray ionization; Et₃N is triethylamine; EtOH is ethanol; Et₂O or ether is diethyl ether; EX. NO. or Ex. No. or EX. No. is example number; g is grams; h or hr is hour; HATU is C-(7-azabenzotriazol-1-yl) urea; N,N,N',N'-tetramethyluronium hexafluorophosphate; HCl is hydrochloric acid; HOBt or HOBut is 1-hydroxybenzotriazole; HPLC is high-performance liquid chromatography; in vacuo is rotary evaporation under diminished pressure; iPr is isopropyl; i-PrOH or IPA is isopropyl alcohol; N is normal; LC is liquid chromatography; LC/MS or LC-MS is liquid chromatography/mass spectroscopy; L is liter(s); m-CPBA is 3-chloroperbenzoic acid; mg is milligrams; mL and mL is milliliter; M is molar; mmol is millimole(s); Me is methyl; MeCN or ACN is acetonitrile; MeOH is methanol; min is minute(s); ms or MS is mass spectrum; MTBE is methyl tert-butyl ether; μg is microgram(s); μL is microliter(s); NaOEt is sodium ethoxide; NaOMe is sodium methoxide; NaOAc is sodium acetate; NMR is nuclear magnetic resonance spectroscopy; NO. or No. is number; PE is petroleum ether; Ph is phenyl; RP or rp is reverse phase; Rₚ is retention time; RT and rt is room temperature; sat., sat'd., and sat is saturated; TEA is triethylamine; TFA is trifluoroacetic acid; TFAA is trifluoroacetic anhydride; THF is tetrahydrofuran; TLC is thin layer chromatography; TMS is trimethylsilyl; Tos is 4-toluene sulfonfonyl; TosCl is 4-toluene sulfonyl chloride; OTs is 4-toluene sulfonate; TsOH is p-toluene sulfonic acid; v is volume; v% and v/v are volume percent; and wt% is weight percent.

The following Schemes, Intermediates and Examples are provided to illustrate the invention and are not to be construed as limiting the scope of the invention in any manner. The reaction schemes, intermediates, and examples illustrate the methods employed in the synthesis of the compounds of formula I. All substituents are as defined above unless indicated otherwise.
Several strategies based upon synthetic transformations known in the literature of organic synthesis may be employed for the preparation of the title compounds of general formula I.

The compounds of the present invention may be prepared using procedures analogous to the procedures exemplified in Schemes 1, 2 and 3. In Scheme 1, carboxylic acid A was converted to ketone C via N-methoxy-N-methyl carbamide B. Treatment of ketone C with hydrazine at 75 °C afforded pyrazole D. Removal of the Boc protecting group with an acid, followed by alkylation of the piperidine ring yielded intermediate F. Final compounds G and H were obtained by alkylation or arylation of the pyrazole ring. The ratio of compound G to H varies.

**SCHEME 1**

![Scheme 1](image)

Alternatively, in Scheme 2, alkylation or arylation of the pyrazole ring may be carried out first to give intermediates I and J. Removal of the Boc protecting group and subsequent alkylation or arylation of the piperidine ring gave the same compounds G and H as described in Scheme 1.

**SCHEME 2**
In Scheme 3, triazole ring in intermediate O was prepared via condensation of N-alkyl hydrazine or aryl hydrazine with \( N-(1\text{-thioxoalkyl})\)carbamide N, which was synthesized by the carboxylic acid M and a thiocarbamide. Removal of the Boc protecting group from intermediate O, followed by addition of the \( R^1 \) group on the piperidine ring affords the final product Q.

SCHEME 3

INTERMEDIATE 1
\textbf{Step 1: \textit{tert}-Butyl 4-[methoxy(methyl)carbamoyl]piperidine-1-carboxylate (1.2).} A mixture of 1-(\textit{tert}-butoxycarbonyl)piperidine-4-carboxylic acid (Aldrich, 6.0 g, 26.2 mmol), \textit{N},\textit{O}-dimethylhydroxylamine hydrochloride (3.06 g, 31.4 mmol), DMAP (3.20 g, 26.2 mmol), and triethyl amine (4.74 mL, 34 mmol) in DCM (200 mL) was stirred at room temperature overnight. The mixture was washed with saturated aqueous KHSO\textsubscript{4} and brine. The organic layer was dried (MgSO\textsubscript{4}) and the solvent was removed. The crude material was directly used in the next step.

\textbf{Step 2: \textit{tert}-Butyl 4-[(3-(4-chlorophenyl)-prop-2-ynoyll]piperidine-1-carboxylate (1.3).} To a solution of \textit{t}-chloro-4-ethynylbenzene (2.76 g, 20.2 mmol) in THF (50 mL) at -78 °C was added 2.5 M butyllithium (8.81 ml, 22.03 mmol in hexane) and stirred for 1 h at -78 °C. \textit{tert}-Butyl 4-[methoxy(methyl)carbamoyl]piperidine-1-carboxylate (Compound 1.2, 5 g, 18.36 mmol) was added. The mixture was stirred at -78 °C for 30 min, then warmed up to 0 °C, stirred for another hour at the same temperature, and diluted with water. The organic layer was separated and the aqueous was extracted with ethyl acetate. The combined organic layers were dried over Na\textsubscript{2}SO\textsubscript{4}, and the drying agent was filtered off. The filtrate was concentrated. The residue was purified by silica gel column chromatography using gradient elution (5 to 30%, EtOAc/hexane). LC-MS m/z 292.06 [M -56 + H]+.

\textbf{Step 3: \textit{tert}-Butyl 4-[(3-(4-chlorophenyl)-1 H-pyrazol-5-vllpiperidme-1-carboxylate (1).} To a solution of teri-butyl 4-[3-(4-chlorophenyl)prop-2-ynoyl]piperidine-1-carboxylate (5.28 g, 15.18 mmol) in ethanol (160 mL) was added hydrazine (0.953 mL, 30.4 mmol) and triethylamine (4.23 mL, 30 4 mmol). The mixture was heated at 75 °C for 3 h. The solvent was evaporated
under reduced pressure, and dried under vacuum to give INTERMEDIATE 1. LC-MS m/z 362.19 [M + H]+, 306.14 [M -56 + H]+.

INTERMEDIATE 2

![Chemical Structure](image)

**2-[3-(4-Chlorophenyl)-5-(piperidin-4-yl)-1 H-pyrazol-1-yl]pyrimidine**

Step 1: tert-Butyl 4-B-(4-chlorophenyl)-1-(pyrimidin-2-yl)-1 H-pyrazol-5-yl]pyridine-1-carboxylate (2A). A mixture of INTERMEDIATE 1 (3.0 g, 8.29 mol), 2-bromopyrimidine (2.9 g, 18.24 mol) and Cs2CO3 (10.8 g, 33.2 mol) in toluene (90 mL) in a pressure vessel was degassed with a stream of nitrogen. Cul (1.58 g, 8.29 mol) and 1,10-phenanthroline (2.99 g, 16.58 mol) were added. The mixture was degassed again with nitrogen, capped and heated at 120 °C for 1.5 h. The mixture was allowed to cool to room temperature, filtered through a plug of Ceite™, and washed with EtOAc. The EtOAc was evaporated under reduced pressure. The resulting residue was purified by silica gel column chromatography using gradient elution (10% to 50% ACN-DCM) to give the title compound as an off-white solid. LC-MS m/z 440.14 [M + H]+.

Step 2: 2-[3-(4-Chlorophenyl)-5-(piperidin-4-yl)-1 H-pyrazol-1-yl]pyrimidine. To a solution of tert-butyl 4-[3-(4-chlorophenyl)-1-(pyrimidin-2-yl)-1 H-pyrazol-5-yl]pyridine-1-carboxylate (1.74 g, 6.75 mmol) in dichloromethane (10 mL) was added 4 M HCl (10 mL, 40 mmol in
dioxane). The mixture was stirred at room temperature overnight. Solvent was evaporated under reduced pressure, and the resulting residue was dried under vacuum to give INTERMEDIATE 2. LC-MS m/z 340.14 [M + H]+.

INTERMEDIATES 3 and 4 in Table 1 were prepared according to the methods described for INTERMEDIATES 1 and 2, starting from the appropriate alkynes and aryl halides.

<table>
<thead>
<tr>
<th>Intermediate Number</th>
<th>Chemical name</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>2-[3-(4-fluorophenyl)-5-(piperidin-4-yl)-1 H-pyrazol-1-yl]pyridine</td>
<td><img src="image" alt="Structure" /></td>
</tr>
<tr>
<td>4</td>
<td>2-[3-(4-fluorophenyl)-5-(piperidin-4-yl)-1 H-pyrazol-1-yl]pyrimidine</td>
<td><img src="image" alt="Structure" /></td>
</tr>
</tbody>
</table>

INTERMEDIATE 5

![INTERMEDIATE 5](image)

4-[3-(4-Chlorophenyl)1-phenyl-1H-pyrazol-5-yl]piperidine

INTERMEDIATE 6

![INTERMEDIATE 6](image)

4-[5-(4-Chlorophenyl)1-phenyl-1H-pyrazol-3-ynpiperidine
To a solution of 1-(4-chlorophenyl)-3-(piperidin-4-yl)propane-1,3-dione hydrochloric salt (0.6 g, 1.98 mmol) in ethanol (15 mL) was added phenylhydrazine (0.59 mL, 5.96 mmol) and triethylamine (0.277 mL, 1.98 mmol). The mixture was heated at 75 °C for 5 h. The volatiles were evaporated under reduced pressure, and the residue was dissolved in EtOAc. The EtOAc layer was washed with 1 N NaOH and brine, dried over Na₂SO₄, filtered. The filtrate was concentrated to give a mixture of 4-[3-(4-chlorophenyl)-1-phenyl-1H-pyrazol-5-yl]piperidine (INTERMEDIATE 5) and 4-[5-(4-chlorophenyl)-1-phenyl-1H-pyrazol-3-yl]piperidine (INTERMEDIATE 6). The mixture was separated via silica gel column chromatography using gradient elution (5 to 20%, MeOH/DCM). LC-MS m/z 338.00 [M +1]⁺.

INTERMEDIATE 7

4-[3-(4-Chlorophenyl)-1H-pyrazol-5-yl]piperidine

INTERMEDIATE 7 was obtained by removal of the Boc group of INTERMEDIATE 1 with an acid. To a solution of INTERMEDIATE 1 (2.5 g, 6.92 mmol) in dichloromethane (10 mL) was added 4 M HCl (10 mL, 40 mmol in dioxane). The mixture was stirred at room temperature overnight. Solvent was evaporated under reduced pressure, and the residue was dried under vacuum.

INTERMEDIATE 8

4-[3-(4-Chlorophenyl)-1H-pyrazol-5-yl]-1-f2-phenylethylpiperidine
To a solution of 4-[3-(4-chlorophenyl)-1\textsubscript{H}-pyrazol-5-yl]piperidine (INTERMEDIATE 7, 4 g, 15.28 mmol) in 60 mL DMF was added \text{K}_2\text{CO}_3 (6.3 g, 45.8 mmol), and 2-(bromoethyl)benzene (2.3 mL, 16.8 mmol). The reaction mixture was stirred at ambient temperature overnight, diluted with E\text{tOAc}, and washed three times with water. The organic layer was separated, dried (MgSO\textsubscript{4}), and concentrated under reduced pressure. LC-MS \textit{mlz} = 366.17 [M+1]\textsuperscript{+}.

INTERMEDIATE 9

Step 1: \textit{rr}-Butyl 4-[(4-fluorophenyl)carbonothioyl]carbamoyl]piperidine-1-carboxylate (9.1). \textit{rr}-Butoxycarbonylpiperidine-4-carboxylic acid (0.70 g, 3.05 mmol, Aldrich) and HATU (1.74 g, 4.58 mmol) were suspended in dichloromethane (10.0 ml) at room temperature for 20 minutes, followed by addition of 4-fluorothiobenzamide (0.47 g, 3.05 mmol) in one portion. The...
mixture was stirred at room temperature for 40 minutes, and DIEA (1.18 g, 9.16 mmol) was added. After stirring at room temperature overnight, the solvent was removed under reduced pressure. The residue was passed through a silica gel cartridge using a gradient of 0-50% ethyl acetate in hexanes to give the title compound as a bright yellow solid.

Step 2: tert-Butyl 4-[3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1H-1,2,4-triazol-5-yl]piperidine-1-carboxylate (9.2). tert-Butyl 4-[[4-(fluorophenyl)carbonothioyl]-carbamoyl] piperidine-1-carboxylate (0.37 g, 1.01 mmol) was mixed with 2-hydrazinylprimidine (0.13 g, 1.21 mmol) and sodium acetate (0.099 g, 1.21 mmol) in 3 ml of 1,4-dioxane and acetic acid (1:1). After heating at 80°C for 45 min, the solvents were removed under vacuum. The resulting residue was redissolved in dichloromethane and washed with saturated sodium bicarbonate solution and water. The organic layers were concentrated, and the residue was purified by column chromatography using a gradient of 0-50% ethyl acetate in hexanes. LC-MS (m/z): 369.10 [M -56 + H]+, 447.14 [M + Na + H]+. ¹H NMR (CDCl₃, 500 MHz): 8 8.92 (d, J = 4.8 Hz, 2 H), 8.25 – 8.28 (m, 2 H), 7.38 (t, J = 4.8 Hz, 1 H), 7.15 (t, J = 8.7 Hz, 2 H), 4.24 – 4.27 (br, 2 H), 3.94 - 4.00 (m, 2 H), 2.91 – 2.96 (br, 2 H), 2.00 – 2.10 (br, 4 H), 1.52 (s, 9 H).

Step 3: 2-[3-(4-Fluorophenyl)Va-piperidin-4-yl]-1H-1,2,4-triazol-1-yl]pyrimidine (9). tert-Butyl 4-[3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1H-1,2,4-triazol-5-yl]piperidine-1-carboxylate (0.21 g, 0.50 mmol) was dissolved in 1,4-dioxane (0.5 ml), followed by addition of 4 N HCl solution in 1,4-dioxane (1.26 ml, 5.04 mmol). After stirring at room temperature for 1.5 h, the mixture was concentrated under reduced pressure. Lyophilization from water and acetonitrile afforded INTERMEDIATE 9. LC-MS (m/z): 325.05 [M + H]+.

INTERMEDIATES 10, 11, 12 and 13 in Table 2 were prepared according to the method described for INTERMEDIATE 9, starting from the appropriate carboxylic acids and arylhydrazines.

<table>
<thead>
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<th>Intermediate</th>
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<td></td>
<td>Chemical Structure</td>
<td>Description</td>
</tr>
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</tr>
<tr>
<td>11</td>
<td><img src="image1.png" alt="Chemical Structure" /></td>
<td>2-[(3-(4-fluorophenyl)-5-(piperidin-4-yl)-1H,1,2,4-triazol-1-yl)pyridine</td>
</tr>
<tr>
<td>12</td>
<td><img src="image2.png" alt="Chemical Structure" /></td>
<td>6-[(3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1H,1,2,4-triazol-5-yl)-3-azabicyclo[3.1.0]-hexane</td>
</tr>
<tr>
<td>13</td>
<td><img src="image3.png" alt="Chemical Structure" /></td>
<td>2-[(3-(4-Chlorophenyl)-5-(piperidin-4-yl)-1H,1,2,4-triazol-1-yl)pyrimidine</td>
</tr>
</tbody>
</table>

**EXAMPLE 1**

![Chemical Structure](image4.png)

2-(3-f4-Chlorophenyl)-3-[1-f2-phenylethyl)piperidin-4-yl]-1H-pyrazol-1-yl]pyrimidine.

**EXAMPLE 2**

![Chemical Structure](image5.png)

2-{5-(4-Chlorophenyl)-3-[1-(2-phenylethyl)piperidin-4-yl]-1H-pyrazol-1-yl]pyrimidine. A reaction mixture of INTERMEDIATE 8 (2.64 g, 7.22 mmol), 2-bromopyrimidine (2.5 g, 15.87 mmol) and Cs₂CO₃ (9.40 g, 28.9 mmol) in toluene (80 mL) in a pressure vessel was degassed with a stream of nitrogen. Cul (1.37 g, 7.22 mmol) and 1,10-phenanthroline (2.60 g, 14.43 mmol).
mmol) were added. The resultant mixture was degassed again with nitrogen, and the reaction vessel was capped, and heated at 120 °C for 4 h. The mixture was allowed to cool to room temperature, and filtered through a plug of Celite™. The Celite™ was washed with EtOAc, and the filtrate was evaporated under reduced pressure. The resulting residue containing both EXAMPLE 1 and EXAMPLE 2 was separated using a silica gel column chromatography via gradient elution (10% to 50% ACN-DCM) to give EXAMPLE 1 as the major compound. LC-MS m/z 444.16 [M + H]+. H NMR (500 MHz, CDCl₃) δ 8.85 (d, J = 4.8 Hz, 2 H), 7.91 (d, J = 8.3 Hz, 2 H), 7.42 (d, J = 8.4 Hz, 2 H), 7.38 (d, J = 7.2 Hz, 2 H), 7.30-7.35 (m, 4 H), 6.82 (s, 1 H), 3.99 (t, J = 7.5 Hz, 1 H), 3.81 (d, J = 11.6 Hz, 2 H), 3.36-3.25 (m, 4 H), 2.86 (broad, 2 H), 2.68-2.63 (m, 2 H), 2.38 (d, J = 14 Hz, 2 H). EXAMPLE 2 was isolated as the minor compound. LC-MS m/z 444.16 [M + H]+.

The compounds in Table 3 were prepared according to the methods described in EXAMPLES 1 and 2, and INTERMEDIATES 1, 7 and 8, starting from the appropriate alkynes and halides.

Table 3

<table>
<thead>
<tr>
<th>Ex. No.</th>
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<tr>
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<td>4-[5-(4-Chlorophenyl)-1-phenyl-1 H-pyrazol-3-yl]-1-(2-phenylethyl)-piperidine</td>
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<td>2-{3-(4-chlorophenyl)-5-[1-(2-phenylethyl)piperidin-4-yl]-1 H-pyrazol-1-yl}pyridine</td>
<td><img src="image2" alt="Structure" /></td>
<td>443.14</td>
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<tr>
<td>5</td>
<td>3-{3-(4-chlorophenyl)-5-[1-(2-phenylethyl)piperidin-4-yl]-1 H-pyrazol-1-yl}pyridine</td>
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<tr>
<td>6</td>
<td>3-{3-(4-chlorophenyl)-5-[1-(2-phenylethyl)piperidin-4-yl]-1H-pyrazol-1-yl}benzoic acid</td>
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<tr>
<td>7</td>
<td>4-{3-(4-chlorophenyl)-5-[1-(2-phenylethyl)piperidin-4-yl]-1H-pyrazol-1-yl}benzonitrile</td>
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<td>8</td>
<td>2-{5-(4-chlorophenyl)-3-[1-(2-phenylethyl)piperidin-4-yl]-1H-pyrazol-1-pyrazol-1-yl}</td>
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<td>9</td>
<td>4-[3-(4-chlorophenyl)-1-(4-methoxyphenyl)-1H-pyrazol-5-yl]-1-(2-phenylethyl)piperidine</td>
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<td>10</td>
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<td>11</td>
<td>2-{3-(4-fluorophenyl)-5-[1-(2-phenylethyl)piperidin-4-yl]-1H-pyrazol-1-yl}pyrimidine</td>
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<tr>
<td>12</td>
<td>4-{3-(4-chlorophenyl)-5-[1-(2-phenylethyl)piperidin-4-yl]-1 H-pyrazol-1-yl}pyrimidine</td>
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<tr>
<td>13</td>
<td>4-{5-(4-chlorophenyl)-3-[1-(2-phenylethyl)piperidin-4-yl]-1 Hyl}pyrimidine</td>
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<td>14</td>
<td>4-[3-(4-chlorophenyl)-1-phenyl-1 H-pyrazol-5-yl]-1-(2-phenylethyl)piperidine</td>
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<td>15</td>
<td>Ethyl 3-{3-(4-chlorophenyl)-5-[1-(2-phenylethyl)piperidin-4-yl]-1 H-pyrazol-1-yl}benzoate</td>
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<td>16</td>
<td>4-{1-(4-chloro-3-methylphenyl)-3-(4-chlorophenyl)-1 H-pyrazol-5-yl]-1-(2-phenylethyl)piperidine</td>
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<td>17</td>
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</tr>
</tbody>
</table>
EXAMPLE 21

4-f1-Benzyl-5-f 4.-chlorophenyl]l 1 H -pyrazol-3-yl1 -1 -(2-phenylethyl)piperidine

EXAMPLE 22

4-ri-Benzyl-3-(4-chlorophenyl)\_1 H -pyrazol-5-vn-1-r2-phenylethvDpiperidine
To a solution of INTERMEDIATE 8 (60 mg, 0.164 mmol) in anhydrous DMF (1 mL) was added 60% NaH (8.2 mg, 0.205 mmol). The mixture was stirred at room temperature for 30 min. Benzylbromide (28 mg, 0.02 mmol) was added at room temperature. The mixture was stirred at room temperature for 2 h, diluted with EtOAc, and washed with water three times. The solvent was evaporated under reduced pressure. The residue containing EXAMPLE 21 and EXAMPLE 22 was separated by preparative TLC (5% MeOH/DCM) to give the EXAMPLE 21 as the major compound. LC-MS m/z 456.16 [M + H]^+.

**H NMR** (500 MHz, CDCl_3)  δ 7.38-7.21 (m, 12 H), 7.16-7.08 (m, 2 H), 6.23 (s, 1 H), 5.28 (s, 2 H), 3.19-3.12 (m, 2 H), 2.93-2.87 (m, 2 H), 2.85-2.76 (m, 1 H), 2.74-2.65 (m, 2 H), 2.27-2.18 (m, 2 H), 2.14-2.05 (m, 2 H), 1.69-1.58 (m, 2 H).

EXAMPLE 22 was isolated as the minor isomer. LC-MS m/z 456.16 [M + H]^+.

The compounds in Table 4 were prepared according to the methods described in EXAMPLES 21 and 22, and INTERMEDIATES 1, 7 and 8, starting from the appropriate alkynes and halides.

<table>
<thead>
<tr>
<th>Ex. No.</th>
<th>Name</th>
<th>Structure</th>
<th>LC-MS (m/e +1)</th>
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</thead>
<tbody>
<tr>
<td>23</td>
<td>4-[5-(4-chlorophenyl)-1-ethyl-1H-pyrazol-3-yl]-1-(2-phenylethyl)piperidine</td>
<td><img src="image1" alt="Structure" /></td>
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<tr>
<td>24</td>
<td>4-[5-(4-chlorophenyl)-1-methyl-1H-pyrazol-3-yl]-1-(2-phenylethyl)piperidine</td>
<td><img src="image2" alt="Structure" /></td>
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<tr>
<td>25</td>
<td>4-[3-(4-chlorophenyl)-1-methyl-1H-pyrazol-5-yl]-1-(2-phenylethyl)piperidine</td>
<td><img src="image3" alt="Structure" /></td>
<td>380.08</td>
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<tr>
<td>26</td>
<td>4-[3-(4-chlorophenyl)-1-ethyl-1H-pyrazol-5-yl]-1-(2-phenylethyl)piperidine</td>
<td><img src="image4" alt="Structure" /></td>
<td>394.11</td>
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<td>Example</td>
<td>Chemical Structure</td>
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<tr>
<td>27</td>
<td><img src="image1" alt="Structure 1" /></td>
<td>4-{1-benzyl-3-[1-(2-phenylethyl)piperidin-4-yl]-1H-pyrazol-5-yl}benzonitrile</td>
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<td><img src="image2" alt="Structure 2" /></td>
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<td>29</td>
<td><img src="image3" alt="Structure 3" /></td>
<td>3-{1-benzyl-3-[1-(2-phenylethyl)piperidin-4-yl]-1H-pyrazol-5-yl}benzoic acid</td>
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<td>30</td>
<td><img src="image4" alt="Structure 4" /></td>
<td>Ethyl 3-{1-benzyl-3-[1-(2-phenylethyl)piperidin-4-yl]-1H-pyrazol-5-yl}benzoate</td>
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</table>

**EXAMPLE 31**
Methyl 3-{4-3-(4-chlorophenyl)-1-(pyrimidin-2-yl)-1H-pyrazol-5-yl]piperidin-1-yl}-2-phenylpropanoate. INTERMEDIATE 2 (60 mg, 0.177 mmol) and methyl 3-oxo-2-phenylpropanoate (31.5, 0.177 mmol) in dichloromethane (1 mL) were treated with sodium triacetoxyborohydride (74.8 mg, 35.4 mmol) at room temperature overnight. The reaction was quenched with 1 N NaOH, and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), and concentrated under reduced pressure. The resulting residue was purified with reverse phase HPLC (10 to 100% 0.1% TFA-acetonitrile/0.1% TFA-water) to give the title compound. LC-MS m/z 502.1 [M + H]+.

Example 3

EXAMPLE 3

3-({4-[3-(4-Chlorophenyl)-1-(pyrimidin-2-yl)-1H-pyrazol-5-yl]piperidin-1-yl}-2-phenylpropanoic acid. EXAMPLE 31 (10 mg, 0.019 mmol) was dissolved in a mixture solvent of THF (1 mL) and water (0.5 mL). The mixture was treated with 3 N NaOH (0.5 mL) at 45 °C overnight, acidified with 1 N hydrochloric acid, and extracted with EtOAc. The solvent was evaporated under reduced pressure. The resulting residue was purified with reverse phase HPLC (10 to 100% 0.1% TFA-acetonitrile/0.1% TFA-water) to give the title compound. LC-MS m/z 488.12. H NMR (500 MHz, CD₃OD) δ 8.86 (d, J = 4.8 Hz, 2 H), 7.88 (d, J = 8.4 Hz, 2 H), 7.52-7.30 (m, 8 H), 6.88 (s, 1 H), 4.28-4.20 (m, 1 H), 4.09-3.86 (m, 2 H), 3.84-3.73 (m, 2 H), 3.50-3.38 (m, 2 H), 2.52-2.38 (m, 2 H), 2.16-1.98 (m, 2 H).

The compounds in Table 5 were prepared according to the methods described in EXAMPLES 31 and 32, starting from INTERMEDIATE 4.

Table 5

<table>
<thead>
<tr>
<th>Ex. No.</th>
<th>Name</th>
<th>Structure</th>
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<td>33</td>
<td>Methyl 3-({4-[3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1H-pyrazol-5-yl]piperidin-1-pyrazol-5-yl]piperidin-1</td>
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</table>
EXAMPLE 35

2-\{4-\{3-(4-Chlorophenyl)-1-(pyrimidin-2-yl)\}-1 \text{H}-pyrazol-5-yl\}piperidin-1-yl\}-1-phenylethanol.

To a solution of INTERMEDIATE 2 (500 mg, 1.47 mmol) in acetonitrile (10 mL) was added K₂CO₃ (610 mg, 4.41 mmol) and 2-bromoacetophenone (322 mg, 1.62 mmol). The reaction was stirred at ambient temperature overnight, diluted with water, and extracted with EtOAc. The organic layer was separated, dried (Na₂SO₄), and concentrated under reduced pressure to give the title compound. LC-MS m/z = 458.22 (M⁺+1).

EXAMPLE 36

2-\{4-\{3-(4-Chlorophenyl)-1-(pyrimidin-2-yl)\}-1 \text{H}-pyrazol-5-yl\}piperidin-1-yl\}-1-phenylethanol.

To a solution of the product of EXAMPLE 35 (50 mg, 0.109 mmol) in MeOH (1 mL) was added sodium borohydride (13 mg, 0.33 mmol) at 0 °C. The mixture was stirred at 0 °C for 30 min, diluted with water, and extracted with EtOAc. The combined organic layers were dried over Na₂SO₄, and the solvent was removed under reduced pressure. The residue was purified by preparative TLC (5% MeOH/DCM) to give the title compound. LC-MS m/z = 460.17 (M⁺+1). 

\(^1\text{H} \) NMR (500 MHz, CDCl₃) δ 8.85 (d, J = 4.7 Hz, 2 H), 7.91 (d, J = 8.6 Hz, 2 H), 7.43-7.37 (m, 5 H), 7.32-7.27 (m, 3 H), 6.67 (s, 1 H), 4.83 (dd, J = 10.2 Hz, J = 3.6 Hz, 1 H), 3.81-3.72 (m, 1 H), 3.35 (d, J = 11.4 Hz, 1 H), 3.01 (d, J = 11.8 Hz, 1 H), 2.66-2.46 (m, 3 H), 2.32-2.22 (m, 1 H), 2.21-2.08 (m, 2 H), 1.96-1.80 (m, 2 H).
The compounds in Table 6 were prepared according to the methods described in EXAMPLE 35 and 36, starting from INTERMEDIATE 2, 3 or 4.

<table>
<thead>
<tr>
<th>Ex. No</th>
<th>Name</th>
<th>Structure</th>
<th>LC-MS (m/e +1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>37</td>
<td>2-{4-[3-(4-fluorophenyl)-1-(pyridin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl} - 1-phenylethanol</td>
<td><img src="image" alt="Structure" /> racemic</td>
<td>443.23</td>
</tr>
<tr>
<td>38</td>
<td>2-{4-[3-(4-fluorophenyl)-1-(pyridin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl} - 1-phenylethanone</td>
<td><img src="image" alt="Structure" /></td>
<td>441.21</td>
</tr>
<tr>
<td>39</td>
<td>2-{4-[3-(4-chlorophenyl)-1-(pyrimidin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl} - 1-phenylethanone</td>
<td><img src="image" alt="Structure" /></td>
<td>418.15</td>
</tr>
<tr>
<td>40</td>
<td>2-{4-[3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl} - 1-phenylethanone</td>
<td><img src="image" alt="Structure" /></td>
<td>444.23</td>
</tr>
<tr>
<td>41</td>
<td>2-{4-[3-(4-fluorophenyl)-1-(pyridin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl} - 1-phenylethanol</td>
<td><img src="image" alt="Structure" /> (S or R)</td>
<td>443.23</td>
</tr>
</tbody>
</table>
To a solution of EXAMPLE 37 (5 mg, 0.011 mmol) in DMF (0.1 mL) was added at
0 °C sodium hydride (2.26 g, 0.056 mmol), followed by addition of iodomethane (0.0042 mL,
0.068 mmol). The mixture was stirred at 0 °C for 30 min, diluted with water, and extracted with
EtOAc. The combined organic layers were dried over Na₂SO₄, and the solvent was removed
under reduced pressure. The resulting residue was purified with preparative TLC (5% MeOH/DCM) to give the title compound. LC-MS m/z = 457.26 [M+H]⁺. ³H NMR(500 MHz,
CDCl₃) δ 8.45 (S, 1 H), 8.00 (S, 1 H), 7.90-7.82 (m, 3 H), 7.42-7.41 (m, 5 H), 7.18-7.12 (m, 3
H), 6.59 (s, 1 H), 3.84-3.77 (m, 1 H), 3.28 (s, 3 H), 3.25-3.08 (m, 2 H), 3.01-2.83 (m, 2 H), 2.68-
2.45 (m, 2 H), 2.18-1.82 (m, 5 H).
1-{4-[3-(4-fluorophenyl)-1-(pyridin-2-y1)-1H-pyrazol-5-y1]piperidin-1-yl}-2-phenylpropan-2-ol
To a solution of EXAMPLE 38 (50 mg, 0.114 mmol) in THF (0.1 mL) was added methyl-
5 magnesium bromide (0.1 mL, 0.30 mmol, 3.0 M) at 0 °C. The mixture was stirred at 0 °C for 1 h,
quenched with water, and diluted with EtOAc. The organic layer was separated, and the aqueous
layer was extracted with EtOAc. The combined organic layers were dried over Na2SO4, and the
solvent was removed under reduced pressure. The resulting residue was purified by preparative
TLC (5% MeOH/DCM) to give the title compound. LC-MS m/z = 457.24 [M+1]+.

EXAMPLE 46

2-[3-(4-fluorophenyl)-1-piperylidenyl]-1H-pyrazol-1-yl]pyridine. A mixture of
INTERMEDIATE 3 (40 mg, 0.101 mmol), sodium tert-butoxide (29.2 mg, 0.304), iodobenzene
(41.3 mg, 0.202 mmol), tri-tert-butylphosphine (2.1 mg, 0.01 mmol), and palladium acetate (2.3
mg, 0.01 mmol) in toluene (0.4 mL) in a 4 mL vial was degassed with a stream of nitrogen and
capped. The mixture was stirred at 50 °C for 4 h. The mixture was then allowed to cool to room
and purified by preparative TLC (hexane/EtOAc = 4/1) to give the title compound as an off-
white solid. LC-MS m/z 399.22 [M + H]+; 'H NMR (500 MHz, CDCl3) δ 8.49 (d, J = 3.3 Hz, 1
H), 8.03 (d, J = 8.2 Hz, 1 H), 7.93-7.84 (m, 3 H), 7.34-7.25 (m, 3 H) 7.14 (t, J = 8.6 Hz, 2 H),
7.02 (d, J = 8.4 Hz, 2 H), 7.14 (t, J = 7.4 Hz, 1 H), 6.59 (s, 1 H), 3.95-3.87 (m, 1 H) 3.85-3.79
(m, 2 H) 2.92-2.85 (m, 2 H) 2.26-2.20 (m, 2 H) 1.96-1.85 (m, 2 H).

The compounds in Table 7 were prepared according to the method described in
EXAMPLE 46, starting from INTERMEDIATE 2 and the appropriate aryl halides.

Table 7

<table>
<thead>
<tr>
<th>Ex. No</th>
<th>Name</th>
<th>Structure</th>
<th>LC-MS</th>
</tr>
</thead>
</table>

- 64 -
EXAMPLE 49

2-(2-[(3-(4-Chlorophenyl)-1-phenyl-1H-pyrazol-5-yl)piperidin-1-yl]ethyl)pyridine. To a solution of INTERMEDIATE 5 (30 mg, 0.089 mmol) in dichloromethane (0.4 mL) was added 2-(2-bromoethyl)pyridine (28.4, 0.107 mmol). The mixture was stirred at 60 °C overnight, and then cooled to room temperature. The mixture was diluted with EtOAc, and washed with water. The organic layer was separated, dried and concentrated. The resulting residue was purified by preparative TLC (using 2 M NH₃ in MeOH/MeOH/DCM/0.05 mL/5 mL/9.5 mL) to give the title compound. LC-MS m/z 443.17 [M + H]+.

The compounds in Table 8 were prepared according to the method described in EXAMPLE 49, starting from INTERMEDIATE 5 or 6, and the appropriate alkyl halides.

<table>
<thead>
<tr>
<th>Ex. No</th>
<th>Name</th>
<th>Structure</th>
<th>LC-MS (m/e +1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>47</td>
<td>6-{4-[3-(4-chlorophenyl)-1- (pyrimidin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl}N,N- dimethylpyridine-3- sulfonamide</td>
<td><img src="image" alt="Structure" /></td>
<td>524.14</td>
</tr>
<tr>
<td>48</td>
<td>4-{6-{4-[3-(4-chlorophenyl)-1- (pyrimidin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl}pyridin-3- yl)sulfonyl]morpholine</td>
<td><img src="image" alt="Structure" /></td>
<td>566.15</td>
</tr>
<tr>
<td>No.</td>
<td>Chemical Structure</td>
<td>Molecular Formula</td>
<td>Molecular Weight</td>
</tr>
<tr>
<td>-----</td>
<td>-------------------</td>
<td>-------------------</td>
<td>------------------</td>
</tr>
<tr>
<td>50</td>
<td>4-[(3-(4-chlorophenyl)-1-phenyl-1<em>H</em>-pyrazol-5-yl)-1-[(2-(4-fluorophenyl)ethyl)piperidin-1-yl]ethyl]piperidine</td>
<td>C37H27Cl3N4O</td>
<td>460.13</td>
</tr>
<tr>
<td>51</td>
<td>2-[(2-[(4-[5-(4-chlorophenyl)-1-phenyl-1<em>H</em>-pyrazol-3-yl]piperidin-1-yl)ethyl]pyridine</td>
<td>C36H29Cl3N4</td>
<td>443.18</td>
</tr>
<tr>
<td>52</td>
<td>4-[5-(4-chlorophenyl)-1-phenyl-1<em>H</em>-pyrazol-3-yl]-1-[(2-(4-fluorophenyl)ethyl)piperidine</td>
<td>C37H27Cl3N4</td>
<td>460.15</td>
</tr>
<tr>
<td>53</td>
<td>1-[(2-(4-chlorophenyl)ethyl]-4-[3-(4-chlorophenyl)-1-phenyl-1<em>H</em>-pyrazol-5-yl]piperidine</td>
<td>C36H29Cl3N4</td>
<td>476.2</td>
</tr>
<tr>
<td>54</td>
<td>1-[(2-(4-chlorophenyl)ethyl]-4-[5-(4-chlorophenyl)-1-phenyl-1<em>H</em>-pyrazol-3-yl]piperidine</td>
<td>C36H29Cl3N4</td>
<td>476.25</td>
</tr>
<tr>
<td>55</td>
<td>4-[2-[(4-[3-(4-chlorophenyl)-1-phenyl-1<em>H</em>-pyrazol-5-yl]piperidin-1-yl]ethyl]morpholine</td>
<td>C36H29Cl3N4O</td>
<td>451.37</td>
</tr>
<tr>
<td>56</td>
<td>4-[(2-[(4-[5-(4-chlorophenyl)-1-phenyl-1<em>H</em>-pyrazol-3-yl]piperidin-1-yl]ethyl]morpholine</td>
<td>C36H29Cl3N4O</td>
<td>451.37</td>
</tr>
</tbody>
</table>
EXAMPLE 57

4-[3-(4-Chlorophenyl)-1-phenyl-1-H-pyrazol-5-yl]-1-(propan-2-yl)piperidine. To a solution of INTERMEDIATE 5 (30 mg, 0.089 mmol) in DCM (0.5 mL) were added acetone (0.026 mL, 0.355 mmol), triacetoxyborohydride (75 mg, 0.355 mmol) and acetic acid (10.7 mg, 0.178 mmol). The mixture was stirred at room temperature overnight, quenched with 1 N NaOH, and extracted with EtOAc. The organic layer was washed with water, dried (MgSO₄) and concentrated. The residue was purified by preparative TLC (4%MeOH/DCM) to give the title compound. LC-MS m/z 380.16 [M + H]⁺.

The compounds in Table 9 were prepared according to the method described in EXAMPLE 57, starting from INTERMEDIATE 5 or 6, and the appropriate ketones.

<table>
<thead>
<tr>
<th>Ex. No</th>
<th>Name</th>
<th>Structure</th>
<th>LC-MS (m/e +1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>58</td>
<td>4-[5-(4-chlorophenyl)-1-phenyl-1-H-pyrazol-3-yl]-1-(propan-2-yl)piperidine</td>
<td><img src="image1.png" alt="Structure" /></td>
<td>380.15</td>
</tr>
<tr>
<td>59</td>
<td>4-[3-(4-chlorophenyl)-1-phenyl-1-H-pyrazol-5-yl]-1-[1-(methylsulfonyl)azetidin-3-yl]piperidine</td>
<td><img src="image2.png" alt="Structure" /></td>
<td>471.13</td>
</tr>
<tr>
<td>60</td>
<td>4-[3-(4-chlorophenyl)-1-phenyl-1-H-pyrazol-5-yl]-1-[1-(methylsulfonyl)pyrrolidin-3-yl]piperidine</td>
<td><img src="image3.png" alt="Structure" /></td>
<td>485.01</td>
</tr>
</tbody>
</table>
EXAMPLE 64

5 4-[3-(4-Fluorophenyl)-1-(pyrimidin-2-yl)-1 (trifluoromethyl)phenyl-1 H-pyrazol-5-yl]piperidin-1-yl)-2-L?-(trifluoromethyl)phenylethanone. INTERMEDIATE 4 (20 mg, 0.062 mmol), 2-(trifluoromethyl)phenylacetic acid (12.6 mg, 0.062 mmol), EDC (11.8 mg, 0.062 mmol), 1H-benzotriazol-1-ol (9.5 mg, 0.062 mmol) and triethylamine (8.6 ul, 0.062 mmol) were mixed in DCM (0.4 mL). The mixture was stirred at room temperature for 4 h, diluted with EtOAc and washed with water. The organic layer was separated, dried (MgS0₄), and concentrated. The resulting residue was purified by reverse phase HPLC (10 to 100% ACN/Water, both containing 0.1% TFA) to give the title compound. LC-MS m/z 510.16 [M + H]+.

The compounds in Table 10 were prepared according to the method described in EXAMPLE 64, starting from INTERMEDIATE 2, 3 or 4, and the appropriate carboxylic acids.
<table>
<thead>
<tr>
<th>Ex. No.</th>
<th>Name</th>
<th>Structure</th>
<th>LC-MS (m/e +1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>65</td>
<td>1-{4-[3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl}-2-(naphthalen-1-yl)ethanone</td>
<td><img src="image1.png" alt="Structure Image" /></td>
<td>492.13</td>
</tr>
<tr>
<td>66</td>
<td>1-{4-[3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl}-2-(2-methoxyphenyl)ethanone</td>
<td><img src="image2.png" alt="Structure Image" /></td>
<td>472.09</td>
</tr>
<tr>
<td>67</td>
<td>1-{4-[3-(4-fluorophenyl)-1-(pyridin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl}-2-phenylethanone</td>
<td><img src="image3.png" alt="Structure Image" /></td>
<td>441.19</td>
</tr>
<tr>
<td>68</td>
<td>1-{4-[3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl}-2-[2-(trifluoromethyl)phenyl]ethanone</td>
<td><img src="image4.png" alt="Structure Image" /></td>
<td>510.01</td>
</tr>
<tr>
<td>69</td>
<td>1-{4-[3-(4-chlorophenyl)-1-(pyrimidin-2-yl)-1 H-pyrazol-5-yl]piperidin-1-yl}-2,2-difluoroethanone</td>
<td><img src="image5.png" alt="Structure Image" /></td>
<td>440.20</td>
</tr>
</tbody>
</table>
2-{3-C4-Fluorophenyl)-5-[1-(2-phenylethyl)piperidin-4-yl]l H -l,2,4-triazol-1-yl}pyrimidine.

To a solution of INTERMEDIATE 9 HCl salt (31.0 mg, 0.078 mmol) in DMF (0.5 ml) was added (2-bromoethyl)benzene (21.7 mg, 0.12 mmol), followed by the portionwise addition of potassium carbonate (55.7 mg, 0.40 mmol). After stirring at 40 °C for 4 hours, the reaction mixture was quenched with 4 ml of a mixture of 1,4-dioxane and water (1:1). The solution was purified by RP-HPLC (10 to 100% ACN/Water, both containing 0.1% TFA) to give the title compound as an off-white solid. LC-MS (m/z): 429.19 [M + H]+. 1HNMR (CDCl₃, 500 MHz):

δ 8.92 (d, J = 4.8 Hz, 2 H), 8.25 (dd, J = 7.7 Hz, 5.7 Hz, 2 H), 7.42 (t, J = 4.8 Hz, 1 H), 7.33-7.36 (m, 2 H), 7.24-7.31 (m, 3 H), 7.18 (t, J = 8.6 Hz, 2 H), 4.43 (t, J = 2.8 Hz, 1 H), 3.93-4.00 (m, 1 H), 3.63 (br, 3 H), 3.32-3.36 (m, 2 H), 3.16-3.19 (m, 2 H), 2.62-2.69 (m, 2 H), 2.33 (d, J = 4.2 Hz, 2 H).

The compounds in Table 11 were prepared according to the method described in EXAMPLE 70, starting from INTERMEDIATE 10, or 11.

<table>
<thead>
<tr>
<th>Ex. No.</th>
<th>Name</th>
<th>Structure</th>
<th>LC-MS(m/e +1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>71</td>
<td>2-{3-(4-chlorophenyl)-5-[1-(2-phenylethyl)piperidin-4-yl]l H -l,2,4-triazol-1-yl}pyrimidine</td>
<td><img src="image1" alt="Structure" /></td>
<td>444.96</td>
</tr>
<tr>
<td>72</td>
<td>4-[3-(4-fluorophenyl)-1-phenyl-1 H-1,2,4-triazol-5-yl]-1-(2-phenylethyl)piperidine</td>
<td><img src="image2" alt="Structure" /></td>
<td>427.20</td>
</tr>
</tbody>
</table>
EXAMPLE 74

5 2-[[3-(4-Fluorophenyl)]-1H-1,2,4-triazol-5-yl]piperidin-1-yl]phenylethanoic acid. To a solution of INTERMEDIATE 9 HCl salt (72.4 mg, 0.17 mmol) in DMF (1.0 ml) was added 2-bromoacetoephone (83.0 mg, 0.42 mmol), followed by the portionwise addition of potassium carbonate (138.0 mg, 1.00 mmol). After stirring at ambient temperature overnight, the reaction mixture was quenched with a saturated solution of ammonium chloride, extracted with EtOAc, and washed with water and brine. The organic layer was separated, dried over Na₂SO₄ and concentrated under reduced pressure. The residue was purified using a silica gel cartridge using a gradient of 0-50% ethyl acetate in hexanes, followed by a gradient of 0-10% methanol in dichloromethane to give the title compound as a yellow oil, which solidified on standing. LC-MS (m/z): 443.10 [M + H]+. H NMR (CDCl₃, 500 MHz): δ 8.92 (d, J = 4.8 Hz, 2 H), 8.25-8.27 (m, 2 H), 8.10 (d, J = 8.1 Hz, 2 H), 7.60 - 7.62 (m, 1 H), 7.50 (t, J = 7.6 Hz, 2 H), 7.37 (t, J = 4.8 Hz, 1 H), 7.15 (t, J = 8.6 Hz, 2 H), 3.88 (s, 2 H), 3.80-3.86 (m, 1 H), 3.13-3.15 (br, 2 H), 2.41-2.45 (br, 2 H), 2.07-2.27 (br, 4 H).

EXAMPLE 75
2-{4-r3-(4-Fluorophenyl)-(pyrimidin-2-yl)-1 H-1,2,4-triazol-5-yl)piperidin-1-vU-l-phenylethanol.

To a solution of the product of EXAMPLE 74 (90.0 mg, 0.20 mmol) in methanol (1.0 ml) was added sodium borohydride (15.4 mg, 0.407 mmol). After stirring at ambient temperature for 1 hour, the reaction mixture was diluted with EtOAc, and washed with water and brine. The organic layer was separated, dried over Na2SO4 and concentrated under reduced pressure. The residue was purified using a silica gel cartridge (a gradient of 0-10% methanol in dichloromethane) to give racemic 2-{4-[3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1 H-l,2,4-triazol-5-yl)piperidin-1-yl}-l-phenylethanol as a light yellow solid. LC-MS (m/z): 445.10 [M + H]+. 

1H NMR (CDCl3, 500 MHz): δ 8.90 (d, J = 4.8 Hz, 2 H), 8.23-8.29 (m, 2 H), 7.35-7.37 (m, 4 H), 7.29-7.34 (m, 1 H), 7.13-7.17 (m, 3 H), 4.80 (dd, J = 10.4 Hz, 3.4 Hz, 1 H), 4.28-4.34 (br, 1 H), 3.81-3.87 (br, 1 H), 3.34 (d, J = 11.4 Hz, 1 H), 3.01 (d, J = 1.4 Hz, 1 H), 2.50-2.63 (m, 3 H), 2.02-2.29 (m, 4 H). The racemic 2-{4-[3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1 H-l,2,4-triazol-5-yl)piperidin-1-yl}-l-phenylethanol was resolved using ChiralPak™ AD column using 60% IPA in heptane to give a faster eluting enantiomer and a slower eluting enantiomer.

The compound in Table 12 was prepared according to the method described in EXAMPLE 75, starting from INTERMEDIATE 13.

<table>
<thead>
<tr>
<th>Ex. No.</th>
<th>Name</th>
<th>Structure</th>
<th>LC-MS(m/e +1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>76</td>
<td>2-{4-[3-(4-chlorophenyl)-1-(pyrimidin-2-yl)-1 H-1,2,4-triazol-5-yl)piperidin-1-yl}-1-phenylethanol</td>
<td><img src="example.png" alt="Structure" /></td>
<td>461.13</td>
</tr>
</tbody>
</table>

EXAMPLE 77

1-(4-[3-(4-Fluorophenyl)-1-(pyrimidin-2-yl)-1 H-1,2,4-triazol-5-ynpiper idin-1-yl)-2-(2-methoxyphenyl)pethanone. To a solution of INTERMEDIATE 9 HCl salt (15.0 mg, 0.035 mmol) and triethylamine (22.9 mg, 0.23 mmol) in DMF (0.25 ml) was added (2-methoxyphenyl)acetic acid (22.9 mg, 0.23 mmol) and stirred at ambient temperature for 1 hour. The reaction mixture was diluted with EtOAc, and washed with water and brine. The organic layer was separated, dried over Na2SO4 and concentrated under reduced pressure. The residue was purified using a silica gel cartridge (a gradient of 0-10% methanol in dichloromethane) to give the product as a light yellow solid. LC-MS (m/z): 445.10 [M + H]+.
acid (7.8 mg, 0.045 mmol), followed by addition of HATU (17.2 mg, 0.045 mmol). After stirring at ambient temperature for 1.5 hour, the reaction mixture was quenched with 2 mL of a mixture of 1,4-dioxane and water (1:1). The mixture was purified by RP-HPLC (10 to 100% ACN/Water, both containing 0.1% TFA) to give the title compound as a white powder after lyophilization. LC-MS (m/z): 473.11 [M + H]+.

The compounds in Table 13 were prepared according to the method described in EXAMPLE 77, starting from INTERMEDIATE 9, and the appropriate carboxylic acids.

<table>
<thead>
<tr>
<th>Ex. No.</th>
<th>Name</th>
<th>Structure</th>
<th>LC-MS(m/e +1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>78</td>
<td>1-4-[(3-(4-fluorophenyl)-1-pyrimidin-2-yl)-1H-1,2,4-triazol-5-yl]piperidin-1-yl]-2-[2-(trifluoromethyl)phenyl]ethanone</td>
<td><img src="image" alt="Structure" /></td>
<td>511.10</td>
</tr>
<tr>
<td>79</td>
<td>1-4-[(3-(4-fluorophenyl)-1-pyrimidin-2-yl)-1H-1,2,4-triazol-5-yl]piperidin-1-yl]-2-(naphthalen-1-yl)ethanone</td>
<td><img src="image" alt="Structure" /></td>
<td>493.10</td>
</tr>
</tbody>
</table>

EXAMPLE 80

2-r3-f4-Fluorophenyl]-5-[1-r(2S)-2-methoxy-2-phenylethyl]piperidin-4-yl]-1H-L2,4-triazol-1-yl]pyrimidine. A mixture of (25)-methoxy(phenyl)ethanal (9.0 mg, 0.060 mmol) and INTERMEDIATE 9 HCl salt (20.0 mg, 0.050 mmol) in 1,2-dichloroethane (0.5 ml) was stirred at room temperature for 5 min, followed by addition of sodium triacetoxyborohydride (16.0 mg, 0.076 mmol). After stirring overnight at room temperature, the solvent was removed under
reduced pressure. The residue was purified by RP-HPLC (10 to 100% ACN/Water, both containing 0.1% TFA) to give the title compound. LC-MS (m/z): 459.16 [M + H]+.

The compound in Table 14 was prepared according to the method described in EXAMPLE 80, starting from INTERMEDIATE 9 and the appropriate aldehyde.

Table 14

<table>
<thead>
<tr>
<th>Ex. No.</th>
<th>Name</th>
<th>Structure</th>
<th>LC-MS(m/e +1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>81</td>
<td>2-[3-(4-fluorophenyl)-5-{1-[(2 R)-2-methoxy-2-phenylethyl]piperidin-4-yl}-1 H-1,2,4-triazol-1-yl]pyrimidine</td>
<td><img src="image" alt="Structure" /></td>
<td>459.16</td>
</tr>
</tbody>
</table>

EXAMPLE 82

![Structure](image)

6-[3-(4-Fluorophenyl)-1-(pyrimidin-2-yl)-1 H-1,2,4-triazol-5-yl]-3-(2-phenylethyl)-3-azabicyclor3.1.01hexane. To a solution of INTERMEDIATE 12 HCl salt (30.0 mg, 0.069 mmol) in DMF (0.5 ml) was added (2-bromoethyl)benzene (38.6 mg, 0.21 mmol), followed by addition of potassium carbonate (57.6 mg, 0.42 mmol). After stirring at room temperature overnight, the reaction mixture was quenched with 4 ml of a mixture of 1,4-dioxane and water (1:1). The solution was purified on RP-HPLC (10 to 100% ACN/Water, both containing 0.1% TFA) to give the title compound as a white solid. LC-MS (m/z): 427.08 [M + H]+.

EXAMPLE 83

![Structure](image)
l-{6-[3-(4-Fluorophenyl)-1-(pyrimidin-2-Yr)-1 H1,2,4-triazol-5-yl]-3-azabicyclo[3.1.Q]-hex-3-yl}2-(2-methoxyphenyl)ethanone. To a solution of INTERMEDIATE 12 HCl salt (15.0 mg, 0.038 mmol) and triethylamine (23.0 mg, 0.23 mmol) in DMF (0.25 ml) was added (2-methoxyphenyl)acetic acid (7.7 mg, 0.046 mmol), followed by addition of HATU (17.3 mg, 0.046 mmol). After stirring at ambient temperature for 1.5 h, the reaction mixture was quenched with 2 ml of a mixture of 1,4-dioxane and water (1:1). The mixture was purified on RP-HPLC (10 to 100% ACN/Water, both containing 0.1% TFA) to give the title compound as a white powder after lyophilization. LC-MS (m/z): 471.04 [M + H]+.

The compounds in Table 15 were prepared according to the method described in EXAMPLE 83, starting from INTERMEDIATE 12 and the appropriate aldehydes.

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<thead>
<tr>
<th>Ex. No.</th>
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<tr>
<td>84</td>
<td>1-{6-[3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1 H1,2,4-triazol-5-yl]-3-azabicyclo[3.1.0]hex-3-yl}-2-[2-(trifluoromethyl)phenyl]ethanone</td>
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<td>85</td>
<td>1-{6-[3-(4-fluorophenyl)-1-(pyrimidin-2-yl)-1 H1,2,4-triazol-5-yl]-3-azabicyclo[3.1.0]hex-3-yl}-2-(naphthalen-1-yl)ethanone</td>
<td><img src="image" alt="Structure" /></td>
<td>491.10</td>
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</table>

The utility of the compounds in accordance with the present invention as inhibitors of prolylcarboxypeptidase (PRCP) enzyme activity may be demonstrated by the following assays:

**BIOLOGICAL EXAMPLE 1**

Prolylcarboxypeptidase (PRCP) Enzyme Activity Assay
The potency of compounds of formula I against PrCP was determined by a fluorescence intensity kinetic assay measuring the IC₅₀ values of PrCP inhibitor test compounds. Recombinant human and mouse PrCP enzymes from CHO or HEK expression systems (with comparable results for HEK enzymes) were prepared in-house and used in the assay. The assay was run on a Perkin Elmer Envision 2103 plate reader using a 320 nm excitation filter and a 405 emission filter. The assay was performed using a Hamilton Star liquid handling workstation. The assay employed the internally quenched fluorescent substrate ((1S)-1-carboxy-5-[(2,4-dinitrophenyl)amino]pentyl N-[(7-methoxy-2-oxo-2H-chromen-4-yl)acetyl]-L-alanyl-L-proline prepared in house.

The assay was run in a 384-well microtiter plate at 37°C with a total volume of 50 uL. Final assay concentrations were 0.13 nM human PrCP (CHO) or 0.09 nM mouse PrCP (CHO) enzyme, 15 uM substrate and varying concentrations of inhibitor in buffer containing 10 mM NaOAc, 100 mM NaCl and 19.5 ug/mL BSA at pH 5.5. The assay also contained 2% DMSO used to solubilize the substrate and inhibitor. Inhibitors were prepared in 100% DMSO and serial diluted (in 100% DMSO) to generate 11 point titration curves. Either 39 uL of human or mouse PrCP enzyme was added to the wells of the assay plate, followed by a 1 uL addition of the serially diluted inhibitor and mixed three times using a 30 uL mix volume. The reaction was initiated by the addition of 10 uL substrate and mixed three times using a 30 uL mix volume. The reactions were monitored continuously over 25 min at 37°C to obtain initial velocities. IC₅₀ values were calculated by comparing the resulting rates of reaction of the inhibited and control initial velocities. For the more potent compounds, a modified dilution series at a lower concentration range was used to more accurately determine the potency.

The enzymes were diluted in a mixture of 10 mM NaOAc (pH 5.5)/100 mM NaCl buffer containing 25 ug/mL bovine serum albumin that had been warmed to 37°C in a water bath. The PrCP inhibitor test compounds were plated in 100% DMSO with a highest concentration of 500 uM. There were 12 dilution points for each compound tested including a blank with DMSO only. The test compounds from the source titration plate were transferred into the assay reaction plate at a 1:50 dilution using the Hamilton Star workstation and mixed, resulting in a final concentration for the test compounds in the range of 10,000 to 0.066 nM. Likewise, two control compounds were similarly titrated and included in each assay, with final starting concentrations starting at 10,000 nM and 200 nM, respectively. The reaction was initiated by the addition of 75 uM of the (1S-1-carboxy-5-[(2,4-dinitrophenyl)amino]pentyl N-[(7-methoxy-2-oxo-2H-chromen-4-yl)acetyl]-L-alanyl-L-proline substrate, which was diluted in a mixture of 10 mM NaOAc (pH 5.5)/100 mM NaCl that had been warmed to 37°C in a water bath, dispensed using the Hamilton Star workstation and mixed. The substrate was solubilized in 100% DMSO prior to dilution into the assay. The final assay concentrations in the 50 uL reactions were 15 uM of (1S)-1-carboxy-5-[(2,4-dinitrophenyl)amino]pentyl N-[(7-methoxy-2-oxo-2H-chromen-4-yl)acetyl]-L-alanyl-L-proline prepared in house.
yl)acetyl]-L-alanyl-L-prolinate substrate, 0.13 nM Human PrCP (CHO) or 0.09 nM Mouse PrCP (CHO), and 2% DMSO.

The resulting rates of reaction were used to determine the IC50 value for each PrCP inhibitor compound tested.

The compounds of the present invention, including the compounds of Examples 1 to 85, exhibit a PRCP inhibition constant IC50 of less than 10 µM. Preferred compounds of the present invention were found to exhibit a PRCP inhibition constant IC50 of less than 1 µM. More preferred compounds of the present invention were found to exhibit a PRCP inhibition constant IC50 of less than 0.01 nM.

<table>
<thead>
<tr>
<th>Example Number</th>
<th>Human PrCP IC50 (nM)</th>
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<tr>
<td>1</td>
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<tr>
<td>13</td>
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<td>68</td>
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<tr>
<td>81</td>
<td>2.58</td>
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<tr>
<td>82</td>
<td>28.6</td>
</tr>
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</table>

**EXAMPLES OF PHARMACEUTICAL COMPOSITIONS**

As a specific embodiment of an oral composition of a composition of the present invention, 5 mg of Example 1 is formulated with sufficient finely divided lactose to provide a total amount of 580 to 590 mg to fill a size O hard gelatin capsule.

As another specific embodiment of an oral composition of a compound of the present invention, 2.5 mg of Example 1 is formulated with sufficient finely divided lactose to provide a total amount of 580 to 590 mg to fill a size O hard gelatin capsule.

While the invention has been described and illustrated in reference to certain preferred embodiments thereof, those skilled in the art will appreciate that various changes, modifications and substitutions can be made therein without departing from the spirit and scope of the invention. For example, effective dosages other than the preferred doses as set forth hereinabove may be applicable as a consequence of variations in the responsiveness of the subject or mammal being treated for severity of bone disorders caused by resorption, or for other indications for the
compounds of the invention indicated above. Likewise, the specific pharmacological responses observed may vary according to and depending upon the particular active compound selected or whether there are present pharmaceutical carriers, as well as the type of formulation and mode of administration employed, and such expected variations or differences in the results are contemplated in accordance with the practices of the present invention. It is intended, therefore, that the invention be limited only by the scope of the claims which follow and that such claims be interpreted as broadly as is reasonable.
WHAT IS CLAIMED IS:

1. A compound of structural formula 1-1 or 1-2:

   (I-1)
   
   and

   (I-2)

or a pharmaceutically acceptable salt thereof; wherein

"a" is a single bond or absent;

X is independently selected from N and CR5;

each R is independently selected from the group consisting of:

1. hydrogen,
2. C1-6 alkyl,
3. C1-6 alkoxy,
4. (CH2)m C0 2C1-6 alkyl,
5. (CH2)mCO C1-6 alkyl,
6. (CH2)m-C(0)-(CH2)u halogen,
7. (CH2)m-C(0)-(CH2)u CO C1-6 alkyl,
8. (CH2)m-C(0)-(CH2)u CO2C1-6 alkyl,
9. (CH2)m-C(0)-(CH2)u pN(Rg)2,
10. (CH2)m-C(0)-(CH2)u p-C3-6 cycloalkyl,
11. (CH2)m-C(0)-(CH2)u p-C2-6 cycloheteroalkyl,
12. (CH2)m-C(0)-(CH2)u p-aryl,
13. (CH2)m-C(0)-(CH2)u p-heteroaryl,
14. (CH2)m -CH(OH)-(CH2)u p-C3-6 cycloalkyl,
15. (CH2)m -CH(OH)-(CH2)u p-C2-6 cycloheteroalkyl,
16. (CH2)m -CH(OH)-(CH2)u p-aryl,
17. (CH2)m -CH(OH)-(CH2)u p-heteroaryl,
18. (CH2)m-NRg-C1-6 alkyl,
19. (CH2)m-O-C1-6 alkyl,
20. (CH2)m -C1-6 alkyl,
(21) -(CH2)m-S02-C1-6 alkyl,
(22) -(CH2)m-S02-(CH2)q-aryl,
(23) -(CH2)m C3-7cycloalkyl,
(24) ~(CH2)m C2-6cyclohexylalkyl,
(25) -(CH2)m ary1,
(26) -(CH2)m biphenyl, and
(27) -(CH2)m heteroaryl,
wherein each CH2, alkyl, alkoxy, cycloalkyl, cyclohexylalkyl, aryl, biphenyl and heteroaryl is unsubstituted or substituted with one to three groups independently selected from Ra;

each R2 is independently selected from the group consisting of:
(1) hydrogen, and
(2) -C1-6 alkyl,
wherein each alkyl is unsubstituted or substituted with one to three substituents selected from Rb;

each R3 is independently selected from the group consisting of:
(1) -C1-6 alkyl,
(2) -(CH2)n C3-7cycloalkyl,
(3) -(CH2)n C2-6cyclohexylalkyl,
(4) -(CH2)n ary1, and
(5) -(CH2)n heteroaryl,
wherein each CH2, alkyl, cycloalkyl, cyclohexylalkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from Rc;

each R4 is independently selected from the group consisting of:
(1) -C1-6 alkyl,
(2) -C1-6 alkoxy,
(3) -CF3,
(4) -(CH2)s C3-7cycloalkyl,
(5) -(CH2)s C2-6cyclohexylalkyl,
(6) -(CH2)s ary1, and
(7) -(CH2)s heteroaryl,
wherein each CH2, alkyl, alkoxy, cycloalkyl, cyclohexylalkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from Rd;

each R5 is independently selected from the group consisting of:
(1) hydrogen, and
(2) -C1-6 alkyl,
wherein each alkyl is unsubstituted or substituted with one to three substituents selected from $R_e$;

each $R^a$ is independently selected from the group consisting of:

1. hydrogen,
2. $-\text{OH}$,
3. oxo,
4. $-\text{Ci}_6$ alkyl,
5. $-\text{OCi}_6$ alkyl,
6. halogen,
7. $-\text{CF}_3$,
8. $-\text{OCF}_3$,
9. $-\text{CN}$,
10. $-\text{C}(0)\text{Ci}_6$ alkyl,
11. $-\text{C}_0_2$H,
12. $-\text{C}_0_2\text{C}_1$-6 alkyl,
13. $-\text{(CH}_2)^t \text{C}_3$-7 cycloalkyl,
14. $-\text{(CH}_2)^t \text{C}_2$-6 cycloheteroalkyl,
15. $-\text{(CH}_2)^0$-1 aryl,
16. $-\text{CH(phenyl)}2$,
17. $-\text{(CH}_2)^t$ heteroaryl,
18. $-\text{SO}_2$Cl-6 alkyl,
19. $-\text{SO}_2$N(RF)$_2$;
20. $-\text{SO}_2$-C3-7 cycloalkyl,
21. $-\text{SO}_2$-C2-6 cycloheteroalkyl,
22. $-\text{SO}_2$-aryl, and
23. $-\text{SO}_2$-heteroaryl;

each $R^b$ is independently selected from the group consisting of:

1. halogen,
2. $-\text{Ci}_6$ alkyl,
3. $-\text{OCi}_6$ alkyl,
4. $-\text{C}_0\text{H}_2$ and
5. $-\text{CO}_2\text{Ci}_6$ alkyl,

wherein alkyl can be substituted with one to three fluorines;

each $R^c$ is independently selected from the group consisting of:

1. hydrogen,
2. $-\text{OH}$,
(3) oxo,
(4) -C1-6 alkyl,
(5) -OC1-6 alkyl,
(6) halogen,
(7) -CF3,
(8) -OCF3,
(9) -CN,
(10) -CO2H, and
(11) -CO2C1-6 alkyl;

each Rd is independently selected from the group consisting of:
(1) hydrogen,
(2) -OH,
(3) oxo,
(4) -C1-6 alkyl,
(5) -OC1-6 alkyl,
(6) halogen,
(7) -CF3,
(8) -OCF3,
(9) -CN,
(10) -CO2H, and
(11) -CO2C1-6 alkyl;

each Re is independently selected from the group consisting of:
(1) halogen,
(2) -Cl-6 alkyl,
(3) -OC1-6 alkyl,
(4) -CO2H, and
(5) -CO2Cl-6 alkyl,

wherein alkyl can be substituted with one to three fluorines;

each Rf is independently selected from the group consisting of:
(1) hydrogen, and
(2) -C1-6 alkyl;

each Rg is independently selected from the group consisting of:
(1) hydrogen, and
(2) -Cl-6 alkyl;
m is selected from 0, 1, 2 and 3;
n is selected from 0, 1, 2, and 3;
p is selected from 0, 1, 2 and 3;
q is selected from 0, 1, 2 and 3;
r is selected from 0, 1, 2, 3 and 4;
s is selected from 0, 1, 2 and 3;
t is selected from 0, 1, 2 and 3; and
u is selected from 1, 2 and 3.

2. The compound of Claim 1 wherein R \( \neq \) is hydrogen; or a pharmaceutically acceptable salt thereof.

3. The compound of Claim 1 wherein "a" is a single bond; or a pharmaceutically acceptable salt thereof.

4. The compound of Claim 1 wherein "a" is absent; or a pharmaceutically acceptable salt thereof.

5. The compound of Claim 4 wherein X is independently selected from N and CH; or a pharmaceutically acceptable salt thereof.

6. The compound of Claim 1 wherein each R \( i \) is independently selected from the group consisting of: \(-\text{C}^{1-6}\text{ alkyl}, \) \(-\text{C}^{1-6}\text{ aryl}, \) \(-\text{CH}_2\text{n ary1, and} \) 
wherein each \( \text{C}^{1-6}\text{ alkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from R}^c; \) or a pharmaceutically acceptable salt thereof.

7. The compound of Claim 6 wherein R\( ^4 \) is -phenyl, wherein phenyl is unsubstituted or substituted with one to three substituents selected from R\( ^d; \) or a pharmaceutically acceptable salt thereof.

8. The compound of Claim 5 wherein each R\( ^3 \) is independently selected from the group consisting of:
   (1) \(-\text{C}^{1-6}\text{ alkyl}, \)
   (2) \(-\text{CH}_2\text{n ary1, and} \)
   (3) \(-\text{CH}_2\text{n heteroaryl,} \)
wherein each \( \text{C}^{1-6}\text{ alkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from R}^c; \) or a pharmaceutically acceptable salt thereof.
9. The compound of Claim 8 wherein each \( R_i \) is independently selected from the group consisting of:
   (1) \(-(CH_2)_n\) aryl, and
   (2) \(-(CH_2)_n\) heteroaryl,
wherein each CH\(_2\), aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from \( R_e \); or a pharmaceutically acceptable salt thereof.

10. The compound of Claim 1 wherein each \( R_1 \) is independently selected from the group consisting of:
   (1) \(-C_1-6\) alkyl,
   (2) \(-COCl-6\) alkyl,
   (3) \(-C(0)-(CH_2)_p\)-aryl,
   (4) \(-C_3-7cycloalkyl,\)
   (5) \(-(CH_2)0-2\) C2-6cycloalkyl,
   (6) \(-(CH_2)0-2\) aryl, and
   (7) \(-(CH_2)0-2\) heteroaryl,
wherein each \( CH_4\), alkyl, cycloalkyl, cycloalkylaryl, aryl, and heteroaryl is unsubstituted or substituted with one to three groups independently selected from \( R_a \); or a pharmaceutically acceptable salt thereof.

11. The compound of Claim 1 wherein each \( R_1 \) is independently selected from the group consisting of:
   (1) \(-(CH_2)0-2\) C2-6cycloalkyl, and
   (2) \-(CH_2)0-2\) aryl,
wherein each CH\(_2\), cycloalkylaryl and aryl is unsubstituted or substituted with one to three groups independently selected from \( R_a \); or a pharmaceutically acceptable salt thereof.

12. The compound of Claim 1 wherein

   30 X is independently selected from N and CH;

   each \( R_1 \) is independently selected from the group consisting of:
   (1) \(-C_1-6\) alkyl,
   (2) \(-COCl-6\) alkyl,
   (3) \(-C(0)-(CH_2)_p\)-aryl,
   (4) \(-C_3-7cycloalkyl,\)
   (5) \(-(CH_2)0-2\) C2-6cycloalkyl,
   (6) \-(CH_2)0-2\) aryl, and
(7) \(-(\text{CH}_2)_{0-2}\) heteroaryl,
wherein each CH$_2$, alkyl, cycloalkyl, cycloheteroalkyl, aryl, and heteroaryl is unsubstituted or substituted with one to three groups independently selected from $R^a$;

5 R$_2$ is hydrogen;

each $R^3$ is independently selected from the group consisting of:

- (1) $-\text{C}_1-6$ alkyl,
- (2) \(-(\text{CH}_2)_n\) aryl, and
- (3) \(-(\text{CH}_2)_n\) heteroaryl,

wherein each CH$_3$, alkyl, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from $R^c$; and

each $R^4$ is independently selected from the group consisting of: \(-(\text{CH}_2)_s\) aryl, wherein each CH$_3$, and aryl is unsubstituted or substituted with one to three substituents selected from $R^d$;

or a pharmaceutically acceptable salt thereof.

13. The compound of Claim 1 wherein:

"a" is absent;

X is independently selected from N and CH;

each $R^1$ is independently selected from the group consisting of:

- (1) \(-(\text{CH}_2)_{0-2}\) C$_2$-6cycloheteroalkyl, and
- (2) \(-(\text{CH}_2)_0-2\) aryl,

wherein each CH$_3$, cycloheteroalkyl and aryl is unsubstituted or substituted with one to three groups independently selected from $R^a$;

30 R$_2$ is hydrogen;

each $R^3$ is independently selected from the group consisting of:

- (1) \(-(\text{CH}_2)_n\) aryl, and
- (2) \(-(\text{CH}_2)_n\) heteroaryl,

wherein each CH$_2$, aryl and heteroaryl is unsubstituted or substituted with one to three substituents selected from $R^c$; and
R₄ is -phenyl, wherein phenyl is unsubstituted or substituted with one to three substituents selected from R₆;

or a pharmaceutically acceptable salt thereof.

14. The compound of Claim 1 wherein:

"a" is absent;

X is independently selected from N and CH;

each R₁ is independently selected from the group consisting of:
   (1) pyrrolidine, and
   (2) -(CH₂)₂-phenyl,

wherein each CH₂, pyrrolidine and phenyl is unsubstituted or substituted with one to three groups independently selected from R₄;

R² is hydrogen;

each R³ is independently selected from the group consisting of:
   (1) -(CH₂)₀-2 phenyl,
   (2) -(CH₂)₀-2 pyridine,
   (3) -(CH₂)₀-2 pyrimidine,
   (4) -(Cl₄)₀-2 pyrazine, and
   (5) -(CH₂)₀-2 imidazole,

wherein each CH₂, phenyl, pyridine, pyrimidine, pyrazine and imidazole is unsubstituted or substituted with one to three substituents selected from R₆; and

R₄ is -phenyl, wherein phenyl is unsubstituted or substituted with one to three substituents selected from R₆;

or a pharmaceutically acceptable salt thereof.

15. The compound of Claim 14 selected from the group consisting of:
or a pharmaceutically acceptable salt thereof.

17. A composition comprising a compound according to Claim 1 and a compound selected from simvastatin, ezetimibe, taranabant and sitagliptin; and a pharmaceutically acceptable carrier.

18. Use of a compound of Claim 1 for the treatment in a mammal of a disorder, condition, or disease responsive to inhibition of prolylcarboxypeptidase.

19. The use of Claim 18 wherein the disorder, condition, or disease is selected from the group consisting of: obesity, diabetes, metabolic syndrome, a diabetes related disorder or an obesity related disorder.

20. The use of a compound of Claim 1 for the preparation of a medicament useful for the treatment of a disorder, condition, or disease responsive to inhibition of prolylcarboxypeptidase.

21. The use of Claim 20 wherein the disorder, condition, or disease is selected from the group consisting of: obesity, diabetes, metabolic syndrome, a diabetes related disorder or an obesity related disorder.

22. The use of Claim 21 wherein the disorder, condition, or disease is obesity.

23. A method of treating a disorder, condition or disease responsive to the inhibition of prolylcarboxypeptidase in a patient in need thereof comprising administration of a therapeutically effective amount of a compound according to Claim 1.

24. The method of Claim 23 wherein the disorder, condition, or disease is selected from the group consisting of: obesity, diabetes, metabolic syndrome, a diabetes related disorder or an obesity related disorder.

25. The method of Claim 24 wherein the disorder, condition, or disease is obesity.
INTERNATIONAL SEARCH REPORT

International application No.  PCT/US 11/35501

A. CLASSIFICATION OF SUBJECT MATTER

<table>
<thead>
<tr>
<th>IPC(B)</th>
<th>A01 N 43/56</th>
<th>A61 K 31/41 5 (201.1.01)</th>
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USPC - 514/405; 514/406

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

USPC - 514/405; 514/406

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

USPC - 514/315; 514/317; 514/319; 514/320; 514/321; 514/322; 514/403 (see search terms below)

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

USPTO-WEST - PGBP, USPT, USOC, EPAB, JPAB keywords: MC4-R, agonist, treatment, obesity, administering, patient, effective amount, combination therapy, agonizing, melanocortin receptors, diabetes mellitus, cholesterol, lowering agents, HMG-CoA reductase inhibitors, humans, melanocortin-4 receptor, molecular modeling, agonists, binding pocket, PRCP, INTER

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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</table>

Further documents are listed in the continuation of Box C.

* Special categories of cited documents:

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A - document defining the general state of the art which is not considered to be of particular relevance
E - earlier application or patent but published on or after the international filing date
L - document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
O - document referring to an oral disclosure, use, exhibition or other means
P - document published prior to the international filing date but later than the priority date claimed
```

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T - later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
X - document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
Y - document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
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Date of the actual completion of the international search 08 July 2011 (08.07.2011)

Date of mailing of the international search report 26 JUL 2011

Name and mailing address of the ISA/US

Mail Stop PCT, Attn: ISA/US, Commissioner for Patents P.O. Box 1450, Alexandria, Virginia 22313-1450

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Form PCT/ISA/2 10 (second sheet) (July 2009)