(54) Titre : SELS ORGANIQUES D'ANTAGONISTES DE L'ANGIOTENSINE II AUGMENTANT LE TAUX DE MONOXYDE D'AZOTE ENDOGENE, COMPOSITIONS ET PROCEDES D'UTILISATION CORRESPONDANTS

(54) Title: ORGANIC NITRIC OXIDE ENHANCING SALTS OF ANGIOTENSIN II ANTAGONISTS, COMPOSITIONS AND METHODS OF USE

(57) Abrégé/Abstract:
The invention describes compositions and kits comprising at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, optionally, at least one nitric oxide enhancing compound and/or at least one therapeutic agent. The invention also provides methods for (a) treating cardiovascular diseases; (b) treating renovascular diseases; (c) treating diabetes; (d) treating diseases resulting from oxidative stress; (e) treating endothelial dysfunctions; (f) treating diseases caused by endothelial dysfunctions; (g) treating cirrhosis; (h) treating pre-eclampsia; (i) treating osteoporosis; (k) treating nephropathy; (l) treating peripheral vascular diseases; (m) treating portal hypertension; (n) treating ophthalmic disorders; (o) treating metabolic syndrome; and (p) treating hyperlipidemia. The organic nitric oxide enhancing compounds that form salts with the angiotensin II antagonists are organic nitrates, organic nitrates, nitrosothiols, thionitrites, thionitrates, NONOates, heterocyclic nitric oxide donors and/or nitroxides. The heterocyclic nitric oxide donors are furoxans, sydnonimines, oxatriazole-5-ones and/or oxatriazole-5-imines.
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(72) Inventors; and

CAI, Xiong [US/US]; 26 Concord Avenue, Belmont, MA 02478 (US).
LIN, Chia-En [US/US]; 98 Prescott Road, Concord, MA 01742 (US).
RANATUNGE, Ramini, R. [US/US]; 11 Bates Road, Lexington, MA 02421 (US).
STEVENSON, Cheri, A. [US/US]; 81 Old Yankee Road, Haverhill, MA 01832 (US).
WEY, Shiw-Jyi [CN/US]; 10 CHARNSTAFFE LANE, Unit 302, Billerica, MA 01821 (US).


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(57) Abstract: The invention describes compositions and kits comprising at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, optionally, at least one nitric oxide enhancing compound and/or at least one therapeutic agent. The invention also provides methods for (a) treating cardiovascular diseases; (b) treating renovascular diseases; (c) treating diabetes; (d) treating diseases resulting from oxidative stress; (e) treating endothelial dysfunctions; (f) treating diseases caused by endothelial dysfunctions; (g) treating cirrhosis; (h) treating pre-ecchymas; (i) treating osteoporosis; (k) treating nephropathy; (l) treating peripheral vascular diseases; (m) treating portal hypertension; (n) treating ophthalmic disorders; (o) treating metabolic syndrome; and (p) treating hyperlipidemia. The organic nitric oxide enhancing compounds that form salts with the angiotensin II antagonists are organic nitrates, organic nitrites, nitrosothiols, thionitrites, thionitrates, NONOates, heterocyclic nitric oxide donors and/or nitroxides. The heterocyclic nitric oxide donors are furoxans, sydnonimines, oxazatriazole-5-ones and/or oxazatriazole-5-imines.
ORGANIC NITRIC OXIDE ENHANCING SALTS OF ANGIOTENSIN II
ANTAGONISTS, COMPOSITIONS AND METHODS OF USE

RELATED APPLICATIONS
This application claims priority under 35 USC § 119 to U.S. Application No. 60/659,401 filed March 9, 2005 and U.S. Application No. 60/750,773 filed December 15, 2005.

FIELD OF THE INVENTION
The invention describes compositions and kits comprising at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, optionally, at least one nitric oxide enhancing compound and/or at least one therapeutic agent. The invention also provides methods for (a) treating cardiovascular diseases; (b) treating renovascular diseases; (c) treating diabetes; (d) treating diseases resulting from oxidative stress; (e) treating endothelial dysfunctions; (f) treating diseases caused by endothelial dysfunctions; (g) treating cirrhosis; (h) treating pre-eclampsia; (j) treating osteoporosis; (k) treating nephropathy; (l) treating peripheral vascular diseases; (m) treating portal hypertension; (n) treating ophthalmic disorders; (o) treating metabolic syndrome; and (p) treating hyperlipidemia. The organic nitric oxide enhancing compounds that form salts with the angiotensin II antagonists are organic nitrates, organic nitrites, nitrosothiols, thionitrites, thionitrates, NONOates, heterocyclic nitric oxide donors and/or nitroxides. The heterocyclic nitric oxide donors are furoxans, sydnonimines, oxatriazole-5-ones and/or oxatriazole-5-imines.

BACKGROUND OF THE INVENTION
The decline in cardiovascular morbidity and mortality in the United States over the past three decades has been the result of significant advances in research on cardiovascular disease mechanisms and therapeutic strategies. The incidence and prevalence of myocardial infarction and death from myocardial infarction, as well as that from cerebrovascular accident, have decreased significantly over this period largely owing to advances in prevention, early diagnosis, and treatment of these very common diseases.

The compounds administered for the treatment of diuresis, cardiovascular diseases, and diseases resulting from oxidative and/or endothelial dysfunctions often result in toxic, chronic and/or debilitating side effects. Cardiovascular compounds such as ACE inhibitors, beta-adrenergic blockers, antithrombotic and vasodilator compounds or anti-hyperlipidemic compounds, show, for example, respiratory toxicity resulting in asthma and/or bronchitis. Hence there is a need in the art for compounds that have improved efficacy, lower toxicity
and that can be used at low dosages. The invention is directed to these, as well as other, important ends.

**SUMMARY OF THE INVENTION**

The invention provides novel organic nitric oxide enhancing salts of angiotensin II antagonists. The angiotensin II antagonists must contain one or more of the following functionalities: a carboxylic acid group (−COOH), a hydroxyl group (−OH), a thiol group (−SH) and/or a primary or secondary amine group (−NH). The organic nitric oxide enhancing compounds that form salts with the angiotensin II antagonists are organic nitrates, organic nitrites, nitrosothiols, thionitrites, thionitrates, NONOates, heterocyclic nitric oxide donors and/or nitroxides. The heterocyclic nitric oxide donors are furoxans, sydnonimines, oxatriazole-5-ones and/or oxatriazole-5-imines. The invention also provides compositions comprising the novel compounds described herein in a pharmaceutically acceptable carrier.

The invention is also based on the discovery that administering at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, optionally, at least one nitric oxide enhancing compound improves the properties of the angiotensin II antagonist. Nitric oxide enhancing compounds include, for example, S-nitrosothiols, nitrites, nitrates, N-oxo-N-nitrosamines, furoxans, sydnonimines, SPM 3672, SPM 4757, SPM 5185, SPM 5186 and analogues thereof, substrates of the various isozymes of nitric oxide synthase, and nitroxides. Thus, another embodiment of the invention provides compositions comprising at least one organic nitric oxide enhancing salt of an angiotensin II antagonist and at least one nitric oxide enhancing compound. The invention also provides for such compositions in a pharmaceutically acceptable carrier.

The invention provides compositions comprising at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, optionally, at least one nitric oxide enhancing compound and/or at least one therapeutic agent, including, but not limited to, aldosterone antagonists, α-adrenergic receptor agonists, α-adrenergic receptor antagonists, angiotensin II antagonists, angiotensin-converting enzyme (ACE) inhibitors, antidiabetic compounds, anti-hyperlipidemic compounds, antimicrobial compounds, antioxidants, antithrombotic and vasodilator compounds, β-adrenergic antagonists, calcium channel blockers, carbonic anhydrase inhibitors, digitalis, diuretics, endothelin antagonists, hydralazine compounds, H₂ receptor antagonists, neutral endopeptidase inhibitors, nonsteroidal antiinflammatory compounds (NSAIDs), phosphodiesterase inhibitors, potassium channel blockers, platelet reducing agents, prostaglandins, proton pump inhibitors, renin inhibitors, selective cyclooxygenase-2 (COX-2) inhibitors, steroids, and combinations
of two or more thereof. In one embodiment the at least one therapeutic agent is selected from the group consisting of an aldosterone antagonist, an angiotensin II antagonist, an angiotensin-converting enzyme (ACE) inhibitor, a β-adrenergic antagonist, a calcium channel blocker, a diuretic, a hydralazine compound and a renin inhibitor. The invention also provides for such compositions in a pharmaceutically acceptable carrier.

Another embodiment of the invention provides compositions comprising an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and at least one therapeutic agent selected from the group consisting of an aldosterone antagonist, an angiotensin II antagonist, an angiotensin-converting enzyme (ACE) inhibitor, a β-adrenergic antagonist, a calcium channel blocker, a diuretic, a hydralazine compound and a renin inhibitor. The invention also provides for such compositions in a pharmaceutically acceptable carrier.

The invention provides methods for (a) treating cardiovascular diseases; (b) treating renovascular diseases; (c) treating diabetes; (d) treating diseases resulting from oxidative stress; (e) treating endothelial dysfunctions; (f) treating diseases caused by endothelial dysfunctions; (g) treating cirrhosis; (h) treating pre-eclampsia; (i) treating osteoporosis; (k) treating nephropathy; (l) treating peripheral vascular diseases; (m) treating portal hypertension; (n) treating ophthalmic disorders; (o) treating metabolic syndrome; and (p) treating hyperlipidemia in a patient in need thereof comprising administering to the patient an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, optionally, at least one therapeutic agent, such as, for example, aldosterone antagonists, α-adrenergic receptor agonists, α-adrenergic receptor antagonists, angiotensin II antagonists, angiotensin-converting enzyme (ACE) inhibitors, antidiabetic compounds, anti-hyperlipidemic compounds, antimicrobial compounds, antioxidants, antithrombotic and vasodilator compounds, β-adrenergic antagonists, calcium channel blockers, carbonic anhydrase inhibitors, digitalis, diuretics, endothelin antagonists, hydralazine compounds, H2 receptor antagonists, neutral endopeptidase inhibitors, nonsteroidal antiinflammatory compounds (NSAIDs), phosphodiesterase inhibitors, potassium channel blockers, platelet reducing agents, prostaglandins, proton pump inhibitors, renin inhibitors, selective cyclooxygenase-2 (COX-2) inhibitors, steroids, and combinations of two or more thereof. The methods can optionally further comprise the administration of at least one nitric oxide enhancing compound. In this embodiment of the invention, the methods can involve (i) administering the organic nitric oxide enhancing salt of the angiotensin II antagonists, (ii) administering the organic nitric oxide enhancing salt of the angiotensin II antagonists, and
nitric oxide enhancing compounds, (iii) administering the organic nitric oxide enhancing salt of the angiotensin II antagonists and therapeutic agents, or (iv) administering the organic nitric oxide enhancing salt of the angiotensin II antagonists, nitric oxide enhancing compounds, and therapeutic agents. In one embodiment the at least one therapeutic agent is selected from the group consisting of an aldosterone antagonist, an angiotensin II antagonist, an angiotensin-converting enzyme (ACE) inhibitor, a β-adrenergic antagonist, a calcium channel blocker, a diuretic, a hydralazine compound and a renin inhibitor. The organic nitric oxide enhancing salt of the angiotensin II antagonists, nitric oxide enhancing compounds, and/or therapeutic agents can be administered separately or as components of the same composition in one or more pharmaceutically acceptable carriers.

Another embodiment of the invention provides kits comprising at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, optionally, at least one nitric oxide enhancing compound. The kit can further comprise at least one therapeutic agent, such as, for example, aldosterone antagonists, α-adrenergic receptor agonists, α-adrenergic receptor antagonists, angiotensin II antagonists, angiotensin-converting enzyme (ACE) inhibitors, antidiabetic compounds, anti-hyperlipidemic compounds, antimicrobial compounds, antioxidants, antithrombotic and vasodilator compounds, β-adrenergic antagonists, calcium channel blockers, carbonic anhydrase inhibitors, digitalis, diuretics, endothelin antagonists, hydralazine compounds, H₂ receptor antagonists, neutral endopeptidase inhibitors, nonsteroidal antiinflammatory compounds (NSAIDs), phosphodiesterase inhibitors, potassium channel blockers, platelet reducing agents, prostaglandins, proton pump inhibitors, renin inhibitors, selective cyclooxygenase-2 (COX-2) inhibitors, steroids, and combinations of two or more thereof. The organic nitric oxide enhancing salt of the angiotensin II antagonist, the nitric oxide enhancing compound and/or therapeutic agent, can be separate components in the kit or can be in the form of a composition in one or more pharmaceutically acceptable carriers.

These and other aspects of the invention are described in detail herein.

DETAILED DESCRIPTION OF THE INVENTION

As used throughout the disclosure, the following terms, unless otherwise indicated, shall be understood to have the following meanings.

"Cardiovascular disease or disorder" refers to any cardiovascular disease or disorder known in the art, including, but not limited to, heart failure, restenosis, hypertension (e.g. pulmonary hypertension, systolic hypertension, labile hypertension, idiopathic hypertension, low-renin hypertension, salt-sensitive hypertension, low-renin, salt-sensitive hypertension,
thromboembolic pulmonary hypertension; pregnancy-induced hypertension; renovascular hypertension; hypertension-dependent end-stage renal disease, hypertension associated with cardiovascular surgical procedures, hypertension with left ventricular hypertrophy, and the like), diastolic dysfunction, coronary artery disease, myocardial infarctions, cerebral infarctions, arterial stiffness, atherosclerosis, atherogenesis, cerebrovascular disease, angina, (including chronic, stable, unstable and variant (Prinzmetal) angina pectoris), aneurysm, ischemic heart disease, cerebral ischemia, myocardial ischemia, thrombosis, platelet aggregation, platelet adhesion, smooth muscle cell proliferation, vascular or non-vascular complications associated with the use of medical devices, wounds associated with the use of medical devices, vascular or non-vascular wall damage, peripheral vascular disease, neointimal hyperplasia following percutaneous transluminal coronary angiograph, vascular grafting, coronary artery bypass surgery, thromboembolic events, post-angioplasty restenosis, coronary plaque inflammation, hypercholesterolemia, embolism, stroke, shock, arrhythmia, atrial fibrillation or atrial flutter, thrombotic occlusion and reclusion cerebrovascular incidents, left ventricular dysfunction and hypertrophy, and the like.

"Heart failure" includes, but is not limited to congestive heart failure, compensated heart failure, decompensated heart failure, and the like.

"Thromboembolic events" include, but are not limited to, ischemic stroke, transient ischemic stroke, myocardial infarction, angina pectoris, thrombosis (for example, restenosis, arterial thrombosis, coronary thrombosis, heart valve thrombosis, coronary stenosis, stent thrombosis, graft thrombosis, and first and subsequent thrombotic stroke, and the like), thromboembolism (for example, pulmonary thromboembolism, cerebral thromboembolism, and the like), thrombophlebitis, thrombocytopenia, bleeding disorders, trombotic occlusion and reocclusion and acute vascular events. Patients who are at risk of developing thromboembolic events, may include those with a familial history of, or genetically predisposed to, thromboembolic disorders, who have had ischemic stroke, transient ischemic stroke, myocardial infarction, and those with unstable angina pectoris or chronic stable angina pectoris and patients with altered prostacyclin/thromboxane A2 homeostasis or higher than normal thromboxane A2 levels leading to increase risk for thromboembolism, including patients with diabetes and rheumatoid arthritis.

"Diseases resulting from oxidative stress" refers to any disease that involves the generation of free radicals or radical compounds, such as, for example, atherogenesis, atheromatosis, arteriosclerosis, atherosclerosis, vascular hypertrophy associated with hypertension, hyperlipoproteinaemia, normal vascular degeneration through aging,
parathyroidal reactive hyperplasia, renal disease (e.g., acute or chronic), neoplastic diseases, inflammatory diseases, neurological and acute bronchopulmonary disease, tumorigenesis, ischemia-reperfusion syndrome, arthritis, sepsis, cognitive dysfunction, endotoxic shock, endotoxin-induced organ failure, and the like.

"Renovascular diseases" refers to any disease or dysfunction of the renal system including, but not limited to, renal failure (e.g., acute or chronic), renal insufficiency, nephrotic edema, acute glomerulonephritis, oliguric renal failure, renal deterioration associated with severe hypertension, unilateral perichymal renal disease, polycystic kidney disease, chronic pyelonephritis, renal diseases associated with renal insufficiency, complications associated with dialysis or renal transplantation, renovascular hypertension, nephropathy, glomerulonephritis, scleroderma, glomerular sclerosis, renal artery stenosis, AIDS-associated nephropathy, immune-mediated renal disease, atheroembolic renal disease, pre-renal azotemia, and the like.

"Endothelial dysfunction" refers to the impaired ability in any physiological processes carried out by the endothelium, in particular, production of nitric oxide regardless of cause. It may be evaluated by, such as, for example, invasive techniques, such as, for example, coronary artery reactivity to acetylcholine or methacholine, and the like, or by noninvasive techniques, such as, for example, blood flow measurements, brachial artery flow dilation using cuff occlusion of the arm above or below the elbow, brachial artery ultrasonography, imaging techniques, measurement of circulating biomarkers, such as, asymmetric dimethylarginine (ADMA), and the like. For the latter measurement the endothelial-dependent flow-mediated dilation will be lower in patients diagnosed with an endothelial dysfunction.

"Methods for treating endothelial dysfunction" include, but are not limited to, treatment prior to the onset/diagnosis of a disease that is caused by or could result from endothelial dysfunction, such as, for example, atherosclerosis, hypertension, diabetes, heart failure, and the like.

"Methods for treating diseases caused by endothelial dysfunction" include, but are not limited to, the treatment of any disease resulting from the dysfunction of the endothelium, such as, for example, arteriosclerosis, heart failure, hypertension, cardiovascular diseases, cerebrovascular diseases, renovascular diseases, mesenteric vascular diseases, pulmonary vascular diseases, ocular vascular diseases, peripheral vascular diseases, peripheral ischemic diseases, and the like.

"Ophthalmic disorders" include, but are not limited to, ophthalmic infections,
cataracts, glaucoma, elevated intraocular pressure, ocular pain (e.g., following corneal surgery), dry eye disorder, ocular hypertension, ocular bleeding, retinal diseases or disorders, presbyopia, macular degeneration, choroidal neovascularization (CNV), retinopathies, such as for example, diabetic retinopathy, vitreoretinopathy, and the like, retinitis, such as for example, cytomegalovirus (CMV) retinitis, uveitis, macular edema, neuropathies and the like.

"Metabolic syndrome" also known as "insulin-resistance syndrome" or "syndrome X" refers to a condition characterized by an increased amount of adipose tissue inside the abdominal cavity, insulin resistance with increased risk of developing senile diabetes, i.e. diabetes type II, high levels of blood fats and high blood pressure.

"Therapeutic agent" includes any therapeutic agent that can be used to treat or prevent the diseases described herein. "Therapeutic agents" include, for example, aldosterone antagonists, α-adrenergic receptor agonists, α-adrenergic receptor antagonists, angiotensin II antagonists, angiotensin-converting enzyme (ACE) inhibitors, antidiabetic compounds, anti-hyperlipidemic compounds, antimicrobial compounds, antioxidants, antithrombotic and vasodilator compounds, β-adrenergic antagonists, calcium channel blockers, carbonic anhydrase inhibitors, digitalis, diuretics, endothelin antagonists, hydralazine compounds, H₂ receptor antagonists, neutral endopeptidase inhibitors, nonsteroidal antiinflammatory compounds (NSAIDs), phosphodiesterase inhibitors, potassium channel blockers, platelet reducing agents, prostaglandins, proton pump inhibitors, renin inhibitors, selective cyclooxygenase-2 (COX-2) inhibitors, steroids, and the like. Therapeutic agent includes the pharmaceutically acceptable salts thereof, pro-drugs, and pharmaceutical derivatives thereof including, but not limited to, the corresponding nitrosated and/or nitrosoylated and/or heterocyclic nitric oxide donor derivatives and/or nitrooxide derivative. Although nitric oxide enhancing compounds have therapeutic activity, the term "therapeutic agent" does not include the nitric oxide enhancing compounds described herein, since nitric oxide enhancing compounds are separately defined.

"Prodrug" refers to a compound that is made more active in vivo.

"Antioxidant" refers to and includes any compound that can react and quench a free radical.

"Angiotensin converting enzyme (ACE) inhibitor" refers to compounds that inhibit an enzyme which catalyzes the conversion of angiotensin I to angiotensin II. ACE inhibitors include, but are not limited to, amino acids and derivatives thereof, peptides, including di- and tri-peptides, and antibodies to ACE which intervene in the renin-angiotensin system by inhibiting the activity of ACE thereby reducing or eliminating the formation of the pressor
substance angiotensin II.

"Angiotensin II antagonists" refers to compounds which interfere with the function, synthesis or catabolism of angiotensin II. Angiotensin II antagonists include peptide compounds and non-peptide compounds, including, but not limited to, angiotensin II antagonists, angiotensin II receptor antagonists, agents that activate the catabolism of angiotensin II, and agents that prevent the synthesis of angiotensin I from angiotensin II. The renin-angiotensin system is involved in the regulation of hemodynamics and water and electrolyte balance. Factors that lower blood volume, renal perfusion pressure, or the concentration of sodium in plasma tend to activate the system, while factors that increase these parameters tend to suppress its function.

"Anti-hyperlipidemic compounds" refers to any compound or agent that has the effect of beneficially modifying serum cholesterol levels such as, for example, lowering serum low density lipoprotein (LDL) cholesterol levels, or inhibiting oxidation of LDL cholesterol, whereas high density lipoprotein (HDL) serum cholesterol levels may be lowered, remain the same, or be increased. Preferably, the anti-hyperlipidemic compound brings the serum levels of LDL cholesterol and HDL cholesterol (and, more preferably, triglyceride levels) to normal or nearly normal levels.

"Diuretic compound" refers to and includes any compound or agent that increases the amount of urine excreted by a patient.

"Neutral endopeptidase inhibitors" refers to and includes compounds that are antagonists of the renin angiotensin aldosterone system including compounds that are dual inhibitors of neutral endopeptidases and angiotensin converting (ACE) enzymes.

"Renin inhibitors" refers to compounds which interfere with the activity of renin.

"Phosphodiesterase inhibitor" or "PDE inhibitor" refers to any compound that inhibits the enzyme phosphodiesterase. The term refers to selective or non-selective inhibitors of cyclic guanosine 3',5'-monophosphate phosphodiesterases (cGMP-PDE) and cyclic adenosine 3',5'-monophosphate phosphodiesterases (cAMP-PDE).

"Platelet reducing agents" refers to compounds that prevent the formation of a blood thrombus via any number of potential mechanisms. Platelet reducing agents include, but are not limited to, fibrinolytic agents, anti-coagulant agents and any inhibitors of platelet function. Inhibitors of platelet function include agents that impair the ability of mature platelets to perform their normal physiological roles (i.e., their normal function, such as, for example, adhesion to cellular and non-cellular entities, aggregation, release of factors such as growth factors) and the like.
"Proton pump inhibitor" refers to any compound that reversibly or irreversibly blocks gastric acid secretion by inhibiting the H⁺/K⁺-ATPase enzyme system at the secretory surface of the gastric parietal cell.

"NSAID" refers to a nonsteroidal anti-inflammatory compound or a nonsteroidal anti-inflammatory drug. NSAIDs inhibit cyclooxygenase, the enzyme responsible for the biosyntheses of the prostaglandins and certain autacoid inhibitors, including inhibitors of the various isozymes of cyclooxygenase (including but not limited to cyclooxygenase-1 and -2), and as inhibitors of both cyclooxygenase and lipoxygenase.

"Cyclooxygenase-2 (COX-2) selective inhibitor" refers to a compound that selectively inhibits the cyclooxygenase-2 enzyme over the cyclooxygenase-1 enzyme. In one embodiment, the compound has a cyclooxygenase-2 IC₅₀ of less than about 2 µM and a cyclooxygenase-1 IC₅₀ of greater than about 5 µM, in the human whole blood COX-2 assay (as described in Brideau et al., Inflamm Res., 45: 68-74 (1996)) and also has a selectivity ratio of cyclooxygenase-2 inhibition over cyclooxygenase-1 inhibition of at least 10, and preferably of at least 40. In another embodiment, the compound has a cyclooxygenase-1 IC₅₀ of greater than about 1 µM, and preferably of greater than 20 µM. The compound can also inhibit the enzyme, lipoxygenase. Such selectivity may indicate an ability to reduce the incidence of common NSAID-induced side effects.

"Patient" refers to animals, preferably mammals, most preferably humans, and includes males and females, and children and adults.

"Transdermal" refers to the delivery of a compound by passage through the skin and into the blood stream.

"Transmucosal" refers to delivery of a compound by passage of the compound through the mucosal tissue and into the blood stream.

"Penetration enhancement" or "permeation enhancement" refers to an increase in the permeability of the skin or mucosal tissue to a selected pharmacologically active compound such that the rate at which the compound permeates through the skin or mucosal tissue is increased.

"Carriers" or "vehicles" refers to carrier materials suitable for compound administration and include any such material known in the art such as, for example, any liquid, gel, solvent, liquid diluent, solubilizer, or the like, which is non-toxic and which does not interact with any components of the composition in a deleterious manner.

"Sustained release" refers to the release of an active compound and/or composition such that the blood levels of the active compound are maintained within a desirable
therapeutic range over a period of time. The sustained release formulation can be prepared using any conventional method known to one skilled in the art to obtain the desired release characteristics.

"Nitric oxide enhancing" refers to compounds and functional groups which, under physiological conditions can increase endogenous nitric oxide. Nitric oxide enhancing compounds include, but are not limited to, nitric oxide releasing compounds, nitric oxide donating compounds, nitric oxide donors, radical scavenging compounds and/or reactive oxygen species scavenger compounds. In one embodiment the radical scavenging compound contains a nitrooxide group.

"Nitrooxide group" refers to compounds that have the ability to mimic superoxide dimutase and catalase and act as radical scavengers, or react with superoxide or other reactive oxygen species via a stable aminoxyl radical i.e. N-oxide.

"Nitric oxide adduct" or "NO adduct" refers to compounds and functional groups which, under physiological conditions, can donate, release and/or directly or indirectly transfer any of the three redox forms of nitrogen monoxide (NO⁺, NO⁻, NO•), such that the biological activity of the nitrooxide species is expressed at the intended site of action.

"Nitric oxide releasing" or "nitric oxide donating" refers to methods of donating, releasing and/or directly or indirectly transferring any of the three redox forms of nitrogen monoxide (NO⁺, NO⁻, NO•), such that the biological activity of the nitrogen monoxide species is expressed at the intended site of action.

"Nitric oxide donor" or "NO donor" refers to compounds that donate, release and/or directly or indirectly transfer a nitrogen monoxide species, and/or stimulate the endogenous production of nitric oxide or endothelium-derived relaxing factor (EDRF) in vivo and/or elevate endogenous levels of nitric oxide or EDRF in vivo and/or are oxidized to produce nitric oxide and/or are substrates for nitric oxide synthase and/or cytochrome P450. Nitric oxide donors also include compounds that are precursors of L-arginine, inhibitors of the enzyme arginase and nitric oxide mediators.

"Heterocyclic nitric oxide donor" refers to a trisubstituted 5-membered ring comprising two or three nitrogen atoms and at least one oxygen atom. The heterocyclic nitric oxide donor is capable of donating and/or releasing a nitrogen monoxide species upon decomposition of the heterocyclic ring. EXEMPLARY heterocyclic nitric oxide donors include oxatriazol-5-ones, oxatriazol-5-imines, sydnonimines, furoxans, and the like.

"Organic nitric oxide enhancing salt" refers to any organic compound that contains a nitric oxide enhancing group and is capable of donating or transferring a biologically active
form of nitrogen monoxide (i.e., nitric oxide or one of its redox congers, such as, for example, nitrosonium cation, nitroxy anion, nitrosyl hydride, and the like) or is capable of increasing the levels of endogeneous nitric oxide and also capable of ionically associating with a compound through at least one acidic group or basic group. Exemplary organic nitric oxide enhancing salts include N-[4-(hydroxymethyl)-1,2,5-oxadiazol-3-yl]carbonylglycine N-oxide (ACS registry number 158590-81-9), 3-[[5-oxido-4-(phenylsulfonyl)-1,2,5-oxadiazol-3-yl]oxy]methylpyridine (ACS registry number 174187-57-6), N,N-dimethyl-2-[[5-oxido-4-(phenylsulfonyl)-1,2,5-oxadiazol-3-yl]oxy]-ethanamine (ACS registry number 186408-97-9), 2,2',2"-nitrilotriethanol trinitrate (ACS registry number 7077-34-1), N,N-bis(2-hydroxyethyl)-nicotinamide dinitrate (ACS registry number 1157-74-0), [1-hydroxy-4-[[4-[(nitrooxy)methyl]benzoyl]amino] butylidene]bis- phosphonic acid (ACS registry number 636585-86-9), 4-(nitrooxy)-, (S)-(2-sulfoethyl) butanethioate (ACS registry number 586351-09-9), 3-(Nitrilox)-2,2-bis[(nitrilox) methyl] propionic acid (ACS registry number 67406-79-5), (S)-[2-[4-(2-hydroxyethyl)-1-piperidinyl]-1,1-dimethylethyl] thionitrite (ACS registry number 364590-39-6), (S)-[1,1-dimethyl-2-[(3-pyridinylcarbonyl)amino]ethyl] thionitrite (ACS registry number 307492-58-6), 2-(acetylamino)-2-carboxy-1,1-dimethylethyl thionitrite (ACS registry number 67776-06-1), and the like.

"Alkyl" refers to a lower alkyl group, a substituted lower alkyl group, a haloalkyl group, a hydroxyalkyl group, an alkenyl group, a substituted alkenyl group, an alkynyl group, a bridged cycloalkyl group, a cycloalkyl group or a heterocyclic ring, as defined herein. An alkyl group may also comprise one or more radical species, such as, for example a cycloalkylalkyl group or a heterocyclicalkyl group.

"Lower alkyl" refers to branched or straight chain acyclic alkyl group comprising one to about ten carbon atoms (preferably one to about eight carbon atoms, more preferably one to about six carbon atoms). Exemplary lower alkyl groups include methyl, ethyl, n-propyl, isopropyl, n-butyl, isobutyl, sec-butyl, t-butyl, pentyl, neopentyl, iso-amyl, hexyl, octyl, and the like.

"Substituted lower alkyl" refers to a lower alkyl group, as defined herein, wherein one or more of the hydrogen atoms have been replaced with one or more $\text{R}^{100}$ groups, wherein each $\text{R}^{100}$ is independently a hydroxy, an ester, an amidyl, an oxo, a carboxyl, a carboxamido, a halo, a cyano, a nitrate, a nitrite, a thionitrile, a thionitrite or an amino group, as defined herein.
"Haloalkyl" refers to a lower alkyl group, an alkenyl group, an alkynyl group, a bridged cycloalkyl group, a cycloalkyl group or a heterocyclic ring, as defined herein, to which is appended one or more halogens, as defined herein. Exemplary haloalkyl groups include trifluoromethyl, chloromethyl, 2-bromobutyl, 1-bromo-2-chloro-pentyl, and the like.

"Alkenyl" refers to a branched or straight chain C₂-C₁₀ hydrocarbon (preferably a C₂-C₈ hydrocarbon, more preferably a C₂-C₆ hydrocarbon) that can comprise one or more carbon-carbon double bonds. Exemplary alkenyl groups include propyl-1-yl, isobutenyl, penten-1-yl, 2,2-methylbuten-1-yl, 3-methylbuten-1-yl, hexan-1-yl, hepten-1-yl, octen-1-yl, and the like.

"Lower alkenyl" refers to a branched or straight chain C₂-C₄ hydrocarbon that can comprise one or two carbon-carbon double bonds.

"Substituted alkenyl" refers to a branched or straight chain C₂-C₁₀ hydrocarbon (preferably a C₂-C₈ hydrocarbon, more preferably a C₂-C₆ hydrocarbon) which can comprise one or more carbon-carbon double bonds, wherein one or more of the hydrogen atoms have been replaced with one or more R¹⁰⁰ groups, wherein each R¹⁰⁰ is independently a hydroxy, an oxo, a carboxyl, a carboxamido, a halo, a cyano or an amino group, as defined herein.

"Alkynyl" refers to an unsaturated acyclic C₂-C₁₀ hydrocarbon (preferably a C₂-C₈ hydrocarbon, more preferably a C₂-C₆ hydrocarbon) that can comprise one or more carbon-carbon triple bonds. Exemplary alkynyl groups include ethynyl, propynyl, butyn-1-yl, butyn-2-yl, pentyn-1-yl, pentyn-2-yl, 3-methylbutyn-1-yl, hexyn-1-yl, hexyn-2-yl, hexyn-3-yl, 3,3-dimethylbutyn-1-yl, and the like.

"Bridged cycloalkyl" refers to two or more cycloalkyl groups, heterocyclic groups, or a combination thereof fused via adjacent or non-adjacent atoms. Bridged cycloalkyl groups can be unsubstituted or substituted with one, two or three substituents independently selected from alkyl, alkoxy, amino, alkylamino, dialkylamino, hydroxy, halo, carboxyl, alkylcarboxylic acid, aryl, amide, ester, alkylcarboxylic ester, carboxamido, alkylcarboxamido, oxo and nitro. Exemplary bridged cycloalkyl groups include adamantyl, decahydrocyclopentyl, quinuclidyl, 2,6-dioxabicyclo(3.3.0)octane, 7-oxabicyclo(2.2.1)heptyl, 8-azabicyclo(3.2.1)oct-2-enyl and the like.

"Cycloalkyl" refers to a saturated or unsaturated cyclic hydrocarbon comprising from about 3 to about 10 carbon atoms. Cycloalkyl groups can be unsubstituted or substituted with one, two or three substituents independently selected from alkyl, alkoxy, amino, alkylamino, dialkylamino, arylamino, diarylamino, alkylarylamino, aryl, amide, ester, hydroxy, halo, carboxyl, alkylcarboxylic acid, alkylcarboxylic ester, carboxamido, alkylcarboxamido, oxo,
alkylsulfinyl, and nitro. Exemplary cycloalkyl groups include cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cyclohexenyl, cyclohepta-1,3-dienyl, and the like.

"Heterocyclic ring or group" refers to a saturated or unsaturated cyclic hydrocarbon group having about 2 to about 10 carbon atoms (preferably about 4 to about 6 carbon atoms) where 1 to about 4 carbon atoms are replaced by one or more nitrogen, oxygen and/or sulfur atoms. Sulfur may be in the thio, sulfinyl or sulfonyl oxidation state. The heterocyclic ring or group can be fused to an aromatic hydrocarbon group. Heterocyclic groups can be unsubstituted or substituted with one, two or three substituents independently selected from alkyl, alkoxy, amino, alkylthio, aryloxy, arylthio, arylalkyl, hydroxy, oxo, thial, halo, carboxyl, carboxylic ester, alkylcarboxylic acid, alkylcarboxylic ester, aryl, arylcarboxylic acid, arylcarboxylic ester, amido, ester, alkylcarboxyl, arylcarboxyl, alkylsulfinyl, carboxamido, alkylcarboxamido, arylcarboxamido, sulfonic acid, sulfonic ester, sulfonamide nitrate and nitro. Exemplary heterocyclic groups include pyrrolyl, furyl, thienvl, 3-pyrrolinyl, 4,5,6-trihydro-2H-pyranyl, pyridinyl, 1,4-dihdropyridinyl, pyrazolyl, triazolyl, pyrimidinyl, pyridazinyl, oxazolyl, thiazolyl, imidazolyl, indolyl, thiophenyl, furanyl, tetrahydrofuranyl, tetrazolyl, pyrrolinyl, pyrrolinyl, oxazolinyl, 1,3-dioxolany, imidazolinyln, imidazolinyl, pyrazolindinyl, pyrazolindinyl, isoxazolyl, isothiazolyl, 1,2,3-oxadiazolyl, 1,2,3-triazolyl, 1,3,4-thiadiazolyl, 2H-pyran, 4H-pyran, piperidinyl, 1,4-dioxany, morpholinyl, 1,4-dithianyl, thiomorpholinyl, pyrazinyl, pipersazinyl, 1,3,5-triazinyl, 1,3,5-trithianyl, benzo(b)thiophenyl, benzimidazolyl, benzothiazolnyl, quinolinyl, 2,6-dioxabicyclo(3.3.0)octane, and the like.

"Heterocyclic compounds" refer to mono- and polycyclic compounds comprising at least one aryl or heterocyclic ring.

"Aryl" refers to a monocyclic, bicyclic, carbocyclic or heterocyclic ring system comprising one or two aromatic rings. Exemplary aryl groups include phenyl, pyridyl, naphtyl, quinoyl, tetrahydrornaphthyl, furanyl, indanyl, indenyl, indoly, and the like. Aryl groups (including bicyclic aryl groups) can be unsubstituted or substituted with one, two or three substituents independently selected from alkyl, alkoxy, alkylthio, amino, alkylamino, dialkylamino, arylamino, diarylamino, alkylarylamino, halo, cyano, alkylsulfinyl, hydroxy, carboxyl, carboxylic ester, alkylcarboxylic acid, alkylcarboxylic ester, aryl, arylcarboxylic acid, arylcarboxylic ester, alkylcarboxyl, arylcarboxyl, amido, ester, carboxamido, alkylcarboxamido, carbonyl, sulfonic acid, sulfonic ester, sulfonamido and nitro. Exemplary substituted aryl groups include tetrafluorophenyl, pentafluorophenyl, sulfonamide, alkylsulfonyl, arylsulfonyl, and the like.
"Cycloalkenyl" refers to an unsaturated cyclic C2-C10 hydrocarbon (preferably a C2-C8 hydrocarbon, more preferably a C2-C6 hydrocarbon) which can comprise one or more carbon-carbon double bonds.

"Alkylaryl" refers to an alkyl group, as defined herein, to which is appended an aryl group, as defined herein. Exemplary alkylaryl groups include benzyl, phenylethyl, hydroxybenzyl, fluorobenzyl, fluorophenylethyl, and the like.

"Arylalkyl" refers to an aryl radical, as defined herein, attached to an alkyl radical, as defined herein. Exemplary arylalkyl groups include benzyl, phenylethyl, 4-hydroxybenzyl, 3-fluorobenzyl, 2-fluorophenylethyl, and the like.

"Arylalkenyl" refers to an aryl radical, as defined herein, attached to an alkenyl radical, as defined herein. Exemplary arylalkenyl groups include styryl, propenylphenyl, and the like.

"Cycloalkylalkyl" refers to a cycloalkyl radical, as defined herein, attached to an alkyl radical, as defined herein.

"Cycloalkylalkoxy" refers to a cycloalkyl radical, as defined herein, attached to an alkoxy radical, as defined herein.

"Cycloalkylalkythio" refers to a cycloalkyl radical, as defined herein, attached to an alkylthio radical, as defined herein.

"Heterocyclicalkyl" refers to a heterocyclic ring radical, as defined herein, attached to an alkyl radical, as defined herein.

"Arylheterocyclic ring" refers to a bi- or tricyclic ring comprised of an aryl ring, as defined herein, appended via two adjacent carbon atoms of the aryl ring to a heterocyclic ring, as defined herein. Exemplary arylheterocyclic rings include dihydroindole, 1,2,3,4-tetrahydroquinoline, and the like.

"Alkylheterocyclic ring" refers to a heterocyclic ring radical, as defined herein, attached to an alkyl radical, as defined herein. Exemplary alkylheterocyclic rings include 2-pyridylmethyl, 1-methylpiperidin-2-one-3-methyl, and the like.

"Alkoxy" refers to R50O-, wherein R50 is an alkyl group, as defined herein (preferably a lower alkyl group or a haloalkyl group, as defined herein). Exemplary alkoxy groups include methoxy, ethoxy, t-butoxy, cyclopentyloxy, trifluoromethoxy, and the like.

"Aryloxy" refers to R55O-, wherein R55 is an aryl group, as defined herein. Exemplary arylkoxy groups include naphthoxy, quinolxyo, isoquinolizinyloxy, and the like.

"Alkylthio" refers to R50S-, wherein R50 is an alkyl group, as defined herein.
"Lower alkylthio" refers to a lower alkyl group, as defined herein, appended to a thio group, as defined herein.

"Arylalkoxy" or "alkoxyaryl" refers to an alkoxy group, as defined herein, to which is appended an aryl group, as defined herein. Exemplary arylalkoxy groups include benzylxoxy, phenylethoxy, chlorophenylethoxy, and the like.

"Arylalklythio" refers to an alkylthio group, as defined herein, to which is appended an aryl group, as defined herein. Exemplary arylalklythio groups include benzylthio, phenylethylthio, chlorophenylethylthio, and the like.

"Arylalklythioalkyl" refers to an arylalklythio group, as defined herein, to which is appended an alkyl group, as defined herein. Exemplary arylalklythioalkyl groups include benzylthiomiethyl, phenylethylthiomiethyl, chlorophenylethylthioethyl, and the like.

"Alkylthioalkyl" refers to an alkylthio group, as defined herein, to which is appended an alkyl group, as defined herein. Exemplary alkylthioalkyl groups include allylthiomethyl, ethylthiomethyl, trifluoroethylthiomethyl, and the like.

"Alkoxyalkyl" refers to an alkoxy group, as defined herein, appended to an alkyl group, as defined herein. Exemplary alkoxyalkyl groups include methoxymethyl, methoxyethyl, isopropanoxymethyl, and the like.

"Alkoxyhaloalkyl" refers to an alkoxy group, as defined herein, appended to a haloalkyl group, as defined herein. Exemplary alkoxyhaloalkyl groups include 4-methoxy-2-chlorobutyl and the like.

"Cycloalkoxy" refers to R54O-, wherein R54 is a cycloalkyl group or a bridged cycloalkyl group, as defined herein. Exemplary cycloalkoxy groups include cyclopropoxy, cyclopentxyloxy, cyclohexyloxy, and the like.

"Cycloalkylthio" refers to R54S-, wherein R54 is a cycloalkyl group or a bridged cycloalkyl group, as defined herein. Exemplary cycloalkylthio groups include cyclopropylthio, cyclopentylthio, cyclohexylthio, and the like.

"Haloalkoxy" refers to an alkoxy group, as defined herein, in which one or more of the hydrogen atoms on the alkoxy group are substituted with halogens, as defined herein. Exemplary haloalkoxy groups include 1,1,1-trichloroethoxy, 2-bromobutoxy, and the like.

"Hydroxy" refers to -OH.

"Oxy" refers to -O-

"Oxo" refers to =O.

"Oxylate" refers to -O` R77+ wherein R77 is an organic or inorganic cation.

"Thiol" refers to -SH.
"Thio" refers to –S–.

"Oxime" refers to =N-OR₈₁ wherein R₈₁ is a hydrogen, an alkyl group, an aryl group, an alkylsulfonyl group, an arylsulfonyl group, a carboxylic ester, an alkylcarbonyl group, an arylcarbonyl group, a carboxamido group, an alkoxyalkyl group or an alkoxyaryl group.

"Hydrazone” refers to =N-N(R₈₁)(R’₈₁) wherein R’₈₁ is independently selected from R₈₁, and R₈₁ is as defined herein.

"Hydrazino” refers to H₂N-N(H)–.

"Organic cation" refers to a positively charged organic ion. Exemplary organic cations include alkyl substituted ammonium cations, and the like.

"Inorganic cation" refers to a positively charged metal ion. Exemplary inorganic cations include Group I metal cations such as for example, sodium, potassium, magnesium, calcium, and the like.

"Hydroxyalkyl” refers to a hydroxy group, as defined herein, appended to an alkyl group, as defined herein.

"Nitrate" refers to -O-NO₂ i.e. oxidized nitrogen.

"Nitrite" refers to -O-NO i.e. oxidized nitrogen.

"Thionitrate" refers to -S-NO₂.

"Thionitrite" and "nitrosothiol" refer to -S-NO.

"Nitro" refers to the group -NO₂ and "nitrosated" refers to compounds that have been substituted therewith.

"Nitroso" refers to the group -NO and "nitrosylated" refers to compounds that have been substituted therewith.

"Nitrile" and "cyano" refer to -CN.

"Halogen" or "halo" refers to iodine (I), bromine (Br), chlorine (Cl), and/or fluorine (F).

"Imine” refers to –C(=N-R₅₁)- wherein R₅₁ is a hydrogen atom, an alkyl group, an aryl group or an arylheterocyclic ring, as defined herein

“Amine” refers to any organic compound that contains at least one basic nitrogen atom.

"Amino" refers to -NH₂, an alkylamino group, a dialkylamino group, an arylamino group, a diarylamino group, an alkylarylamino group or a heterocyclic ring, as defined herein.

"Alkylamino" refers to R₅₀NH₂, wherein R₅₀ is an alkyl group, as defined herein. Exemplary alkylamino groups include methylamino, ethylamino, butylamino, cyclohexylamino, and the like.
"Arylamino" refers to $R_{35}NH\text{-}$, wherein $R_{35}$ is an aryl group, as defined herein.

"Dialkylamino" refers to $R_{52}R_{53}N\text{-}$, wherein $R_{52}$ and $R_{53}$ are each independently an alkyl group, as defined herein. Exemplary dialkylamino groups include dimethylamino, diethylamino, methyl propargylamino, and the like.

"Diarylamo" refers to $R_{55}R_{60}N\text{-}$, wherein $R_{55}$ and $R_{60}$ are each independently an aryl group, as defined herein.

"Alkylarylamino" or "arylalkylamino" refers to $R_{52}R_{55}N\text{-}$, wherein $R_{52}$ is an alkyl group, as defined herein, and $R_{55}$ is an aryl group, as defined herein.

"Alkylarylamino" refers to $R_{52}R_{79}N\text{-}$, wherein $R_{52}$ is an alkyl group, as defined herein, and $R_{79}$ is an arylalkyl group, as defined herein.

"Alkylcycloalkylamino" refers to $R_{52}R_{80}N\text{-}$, wherein $R_{52}$ is an alkyl group, as defined herein, and $R_{80}$ is a cycloalkyl group, as defined herein.

"Aminoalkyl" refers to an amino group, an alkylamino group, a dialkylamino group, an arylamino group, a diarylaminogroup, an alkylarylamino group or a heterocyclic ring, as defined herein, to which is appended an alkyl group, as defined herein. Exemplary aminoalkyl groups include dimethylaminopropyl, diphenylaminocyclopentyl, methylaminomethyl, and the like.

"Aminoaryl" refers to an aryl group to which is appended an alkylamino group, an arylamino group or an arylalkylamino group. Exemplary aminoaryl groups include anilino, N-methylanilino, N-benzylanilino, and the like.

"Sulfonyl" refers to $-S(O)\text{-}$.

"Methanthial" refers to $-C(S)\text{-}$.

"Thial" refers to $=S$.

"Sulfonyl" refers to $-S(O)_2$.

"Sulfonic acid" refers to $-S(O)_2OR_{76}$, wherein $R_{76}$ is a hydrogen, an organic cation or an inorganic cation, as defined herein.

"Alkylsulfonic acid" refers to a sulfonic acid group, as defined herein, appended to an alkyl group, as defined herein.

"Arylsulfonic acid" refers to a sulfonic acid group, as defined herein, appended to an aryl group, as defined herein

"Sulfonic ester" refers to $-S(O)_2OR_{58}$, wherein $R_{58}$ is an alkyl group, an aryl group, or an aryl heterocyclic ring, as defined herein.

"Sulfonamido" refers to $-S(O)_2-N(R_{51})(R_{57})$, wherein $R_{51}$ and $R_{57}$ are each independently a hydrogen atom, an alkyl group, an aryl group or an arylheterocyclic ring, as
defined herein, or R₅₁ and R₅₇ when taken together are a heterocyclic ring, a cycloalkyl group or a bridged cycloalkyl group, as defined herein.

"Alkylsulfonamido" refers to a sulfonamido group, as defined herein, appended to an alkyl group, as defined herein.

"Arylsulfonamido" refers to a sulfonamido group, as defined herein, appended to an aryl group, as defined herein.

"Alkylthio" refers to RₛₒS-, wherein Rₛₒ is an alkyl group, as defined herein (preferably a lower alkyl group, as defined herein).

"Arylthio" refers to Rₛ₅S-, wherein Rₛ₅ is an aryl group, as defined herein.

"Arylalkylthio" refers to an aryl group, as defined herein, appended to an alkylthio group, as defined herein.

"Alkylsulfinyl" refers to Rₛₒ-S(O)-, wherein Rₛₒ is an alkyl group, as defined herein.

"Alkylsulfonyl" refers to Rₛₒ-S(O)₂-, wherein Rₛₒ is an alkyl group, as defined herein.

"Alkylsulfonyloxy" refers to Rₛₒ-S(O)₂-O-, wherein Rₛₒ is an alkyl group, as defined herein.

"Arylsulfinyl" refers to Rₛ₅-S(O)-, wherein Rₛ₅ is an aryl group, as defined herein.

"Arylsulfonyl" refers to Rₛ₅-S(O)₂-, wherein Rₛ₅ is an aryl group, as defined herein.

"Arylsulfonyloxy" refers to Rₛ₅-S(O)₂-O-, wherein Rₛ₅ is an aryl group, as defined herein.

"Amidyl" refers to Rₛ₁C(O)N(Rₛ₇)- wherein Rₛ₁ and Rₛ₇ are each independently a hydrogen atom, an alkyl group, an aryl group or an arylheterocyclic ring, as defined herein.

"Ester" refers to Rₛ₁C(O)Rₛ₂- wherein Rₛ₁ is a hydrogen atom, an alkyl group, an aryl group or an arylheterocyclic ring, as defined herein and Rₛ₂ is oxygen or sulfur.

"Carbamoyl" refers to -O-C(O)NRₛ₁(Rₛ₇), wherein Rₛ₁ and Rₛ₇ are each independently a hydrogen atom, an alkyl group, an aryl group or an arylheterocyclic ring, as defined herein, or Rₛ₁ and Rₛ₇ taken together are a heterocyclic ring, a cycloalkyl group or a bridged cycloalkyl group, as defined herein.

"Carboxyl" refers to -C(O)ORₛ₆, wherein Rₛ₆ is a hydrogen, an organic cation or an inorganic cation, as defined herein.

"Carbonyl" refers to -C(O)-.

"Alkylcarbonyl" refers to Rₛ₂-C(O)⁻, wherein Rₛ₂ is an alkyl group, as defined herein.

"Arylcarbonyl" refers to Rₛ₅-C(O)⁻, wherein Rₛ₅ is an aryl group, as defined herein.

"Arylalkylcarbonyl" refers to Rₛ₅-Rₛ₂-C(O)⁻, wherein Rₛ₅ is an aryl group, as defined herein, and Rₛ₂ is an alkyl group, as defined herein.
“Alkylarylcarbonyl” refers to $R_{52}$-$R_{55}$-$C(O)$-, wherein $R_{55}$ is an aryl group, as defined herein, and $R_{52}$ is an alkyl group, as defined herein.

"Heterocyclicalkylcarbonyl" refer to $R_{78}$-$C(O)$- wherein $R_{78}$ is a heterocyclicalkyl group, as defined herein.

"Carboxylic ester" refers to -$(C(O))OR_{58}$, wherein $R_{58}$ is an alkyl group, an aryl group or an aryl heterocyclic ring, as defined herein.

"Alkylcarboxylic acid" and "alkylcarboxyl" refer to an alkyl group, as defined herein, appended to a carboxyl group, as defined herein.

"Alkylcarboxylic ester" refers to an alkyl group, as defined herein, appended to a carboxylic ester group, as defined herein.

"Alkyl ester" refers to an alkyl group, as defined herein, appended to an ester group, as defined herein.

"Arylcarboxylic acid" refers to an aryl group, as defined herein, appended to a carboxyl group, as defined herein.

"Arylcarboxylic ester" and "arylcarboxyl" refer to an aryl group, as defined herein, appended to a carboxylic ester group, as defined herein.

"Aryl ester" refers to an aryl group, as defined herein, appended to an ester group, as defined herein.

"Carboxamido" refers to -$(C(O))N(R_{51})(R_{57})$, wherein $R_{51}$ and $R_{57}$ are each independently a hydrogen atom, an alkyl group, an aryl group or an arylheterocyclic ring, as defined herein, or $R_{51}$ and $R_{57}$ when taken together are a heterocyclic ring, a cycloalkyl group or a bridged cycloalkyl group, as defined herein.

"Alkylcarboxamido" refers to an alkyl group, as defined herein, appended to a carboxamido group, as defined herein.

"Arylcarboxamido" refers to an aryl group, as defined herein, appended to a carboxamido group, as defined herein.

"Urea" refers to -$(N(R_{59})-C(O))N(R_{51})(R_{57})$ wherein $R_{51}$, $R_{57}$, and $R_{59}$ are each independently a hydrogen atom, an alkyl group, an aryl group or an arylheterocyclic ring, as defined herein, or $R_{51}$ and $R_{57}$ taken together are a heterocyclic ring, a cycloalkyl group or a bridged cycloalkyl group, as defined herein.

"Phosphoryl" refers to -$P(R_{70})(R_{71})(R_{72})$, wherein $R_{70}$ is a lone pair of electrons, thial or oxo, and $R_{71}$ and $R_{72}$ are each independently a covalent bond, a hydrogen, a lower alkyl, an alkoxy, an alkylamino, a hydroxy, an oxy or an aryl, as defined herein.

"Phosphoric acid" refers to -$P(O)(OR_{51})$OH wherein $R_{51}$ is a hydrogen atom, an alkyl
group, an aryl group or an arylheterocyclic ring, as defined herein.

"Phosphinic acid" refers to \(-\text{P}(\text{O})(\text{R}_{51})\text{OH}\) wherein \(\text{R}_{51}\) is a hydrogen atom, an alkyl group, an aryl group or an arylheterocyclic ring, as defined herein.

"Silyl" refers to \(-\text{Si}(\text{R}_{73})(\text{R}_{74})(\text{R}_{75})\), wherein \(\text{R}_{73}, \text{R}_{74}\) and \(\text{R}_{75}\) are each independently a covalent bond, a lower alkyl, an alkoxy, an aryl or an arylalkoxy, as defined herein.

"Organic acid" refers to compound having at least one carbon atom and one or more functional groups capable of releasing a proton to a basic group. The organic acid preferably contains a carboxyl, a sulfonic acid or a phosphoric acid moiety. Exemplary organic acids include acetic acid, benzoic acid, citric acid, camphorsulfonic acid, methanesulfonic acid, taurocholic acid, chlordronic acid, glyphosphate, medronic acid, and the like.

"Inorganic acid" refers to a compound that does not contain at least one carbon atom and is capable of releasing a proton to a basic group. Exemplary inorganic acids include hydrochloric acid, sulfuric acid, nitric acid, phosphoric acid, and the like.

"Organic base" refers to a carbon containing compound having one or more functional groups capable of accepting a proton from an acid group. The organic base preferably contains an amine group. Exemplary organic bases include triethylamine, benzyl-diethylamine, dimethylethyl amine, imidazole, pyridine, piperydine, and the like.

The contemplated angiotensin II antagonists are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, (1996); Merck Index on CD-ROM, 13th Edition; STN Express, file phar and
file registry, the disclosures of each of which are incorporated by reference herein in their entirety.

In one embodiment the angiotensin II antagonists must contain one or more of the following functionalities: a carboxylic acid group (-COOH), or an acidic nitrogen group (-NH). The angiotensin II antagonists form salts with at least one organic nitric oxide enhancing compound that is ionically associated with the angiotensin II antagonist through one or more acid groups. The organic nitric oxide enhancing compounds that form salts with the angiotensin II antagonists are organic nitrates, organic nitrites, nitrosothiols, thionitrites, thionitrates, NONOates, heterocyclic nitric oxide donors and/or nitrooxides that must contain a basic functionality, such as, for example an amidine group (-C(=NH)-NH₂), a guanidine group (-N(H)C(O)-NH₂) and/or a primary or secondary amine group (-NH), and the like. The heterocyclic nitric oxide donors are furoxanis, sydnonimines, oxatriazole-5-ones and/or oxatriazole-5-imines.

In another embodiment, the invention describes angiotensin II antagonists of Formula (I):

![Chemical structure]

wherein:

X₃ is:

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(5) \[
\begin{array}{c}
\text{N-S} \\
\text{Z}
\end{array}
\]

(6) \[-\text{N}(\text{H}^* \text{Z})-\text{S(O)}_2-\text{CF}_3;\]

(7) \[
\begin{array}{c}
\text{S} \\
\text{N} \\
\text{O} \\
\text{H} \\
\text{C}_3 \text{C}_2 \text{H}_4 \text{H}_5
\end{array}
\]

(8) \[-\text{N}(\text{D}_1)-\text{C(O)}-\text{N(\text{D}_1)}-\text{CH}_2-\text{CH}_2-\text{CH}_3;\]

(9) \[-\text{R}_{37};\]

(10) \[-\text{C(O)}-\text{CH}_2-\text{NH(\text{D}_4)};\]

(11) \[-\text{S(O)}_2-\text{N(\text{H}^* \text{Z})-C(O)}-\text{C}_6\text{H}_5;\]

(12) \[-\text{S(O)}_2-\text{N(\text{H}^* \text{Z})-C(O)}-\text{N}\text{D}_4^- \text{CH}_2-\text{CH}_2-\text{CH}_3; \text{ or}\]

(13) \[-\text{S(O)}_2-\text{N(\text{D}_4)}-\text{OD}_1;\]

\(Z_3\) is a carbon, -CH or a nitrogen atom;

\(R_{10}\) is a fluorine or a hydrogen atom;

\(Y_3\) is:

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$Z_4$ is C-$R_{29}$ or a nitrogen;

$R_{37}$ is $-\text{C(O)U}_3\text{D}_1$ or $-\text{COOH}\cdot Z$;

$R_{11}$ is:

(1) $-\text{CH}_2\cdot \text{OD}_1$;

(2) $-R_{37}$;
(3) -C(O)-O-CH(CH₃)-O-C(O)-OR₁₃; or
(4) -CH₂-N(D₁)-C(O)-OR₁₃;

R₁₂ is a chlorine, -SCH₃ or a haloalkyl;
R₁₃ is a lower alkyl or K;
R₁₄ is a lower alkyl or a cycloalkyl;
R₁₅ is:

1. hydrogen;
2. a lower alkyl;
3. 

(4)

(5) -R₃⁷;

R₁₆ is a hydrogen, a lower alkyl, an alkoxy, -OD₁, a cyano, -R₃⁷, NH(D₁) or an alkylcarbonyl;
R₁₇ is an aryl or a cycloalkyl;
R₁₈ at each occurrence is independently selected from a lower alkyl, an alkoxyalkyl, an alkylcarboxylic acid, an alkyl-R₃⁷, an hydroxyalkyl, an arylalkoxy, an arylalkyl or an aryl;
R₁₉ is a hydrogen or -R₃⁷;
R₂₀ is a hydrogen, a lower alkyl or -R₃⁷;

R₂₁ is:

(1)

(2)

R₂₂ is a hydrogen, -R₃⁷ or
R_{32} is a hydrogen, an alkyl or an aryl;  
R_{33} is \(-(\text{CH}_2)_2\)-OD$_1$ or  
\[ \text{R}_{34} \text{ is a hydrogen, a lower alkyl, a lower haloalkyl, an aryl or an arylalkyl;} \]  
\[ \text{R}_{27} \text{ is a lower alkyl, an aryl an arylalkyl or } -(\text{CH}_2)_k\text{-R}_{37}; \]  
\[ \text{R}_{28} \text{ is } -\text{OD}_1, -\text{S(O)}_2\text{-N(D}_1\text{)H, } -\text{N(D}_1\text{)H, } -\text{R}_{37} \text{ or } \text{CH}_2\text{-OD}_1; \]  
\[ \text{R}_{29} \text{ is a hydrogen, a lower alkyl or } -\text{R}_{37}; \]  
\[ \text{R}_{30} \text{ is a lower alkyl or a haloalkyl;} \]  
\[ \text{R}_{31} \text{ is:} \]  
\[ \text{R}_{45} \text{ is a hydrogen or a lower alkyl;} \]  
\[ \text{R}_{46} \text{ is an alkoxy, an amino group or } -\text{N(R}_{13}\text{)(R}_{13}); \]  
\[ \text{R}_{42} \text{ is a lower alkyl or } -(\text{C(R}_a\text{)R}_p)_{k}\text{-V}_4; \]  
\[ \text{R}_{43} \text{ and } \text{R}_{44} \text{ taken together are:} \]  
\[ \text{\(o_1\) is an integer from 0 to 3;} \]  
\[ \text{k is an integer from 1 to 3;} \]
D₁ is a hydrogen, -H· Z or K;
D₄ is a hydrogen or K;
Z is an organic base or -N(R₃₈)(R₃₉)(R₄₀);
R₃₈, R₃₉ and R₄₀ are each independently selected from K or R₆, or R₃₈ and R₃₉ taken together with the nitrogen to which they are attached are a heterocyclic ring, with the proviso that when the heterocyclic ring is an aromatic ring it can be substituted at any position by L and R₃₉ is not present;
L is -(W₃)a-E_b-(C(R₆)(R₇))p₁-E_c-(C(R₆)(R₇))q₁-(W₃)d-E_k-(W₃)i-E_j-(W₃)g-(C(R₆)(R₇))z-V₄;
K is -(W₃)a-E_b-(C(R₆)(R₇))p₁-E_c-(C(R₆)(R₇))q₁-(W₃)d-E_k-(W₃)i-E_j-(W₃)g-(C(R₆)(R₇))z-V₄;
a, b, c, d, g, i and j are each independently an integer from 0 to 3;
p₁, x, y and z are each independently an integer from 0 to 10;
V₄ is V₃, R₆, -U₃-V₅ or V₆;
V₃ is:
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R_{24} \text{ is } -\text{C}_6\text{H}_4\text{R}_{27}, -\text{CN}, -\text{S(O)}_2-\text{C}_6\text{H}_4\text{R}_{27}, -\text{C(O)}-\text{N(R}_a\text{)(R}_i\text{)}, -\text{NO}_2 \text{ or } -\text{C(O)}-\text{OR}_{25};

R_{25} \text{ is an aryl group, a lower alkyl group, a haloalkyl group, a hydroxyalkyl group or an arylalkyl group;}

R_{26} \text{ is } -\text{C(O)}- \text{ or } -\text{S(O)}_2-;

R_{27} \text{ is a hydrogen, } -\text{CN}, -\text{S(O)}_2-\text{R}_{25}, -\text{C(O)}-\text{N(R}_a\text{)(R}_i\text{)}, -\text{NO}_2 \text{ or } -\text{C(O)}-\text{OR}_{25};

T' \text{ is oxygen, sulfur or } \text{NR}_6;

R_6 \text{ is a hydrogen, a lower alkyl group, an aryl group;}

V_6 \text{ is:}

Z_5 \text{ is } -\text{CH}_2 \text{ or oxygen;
Z₃ is -CH or nitrogen;

W₃ at each occurrence is independently -C(O)-, -C(S)-, -T₃-, -(C(R₆)(R₇))ₙ-, -N(R₄)R₅, an alkyl group, an aryl group, a heterocyclic ring, an arylheterocyclic ring, -(CH₂CH₂O)ₚ⁻ or a heterocyclic nitric oxide donor;

E at each occurrence is independently -T₃-, an alkyl group, an aryl group, -(C(R₆)(R₇))ₙ-, a heterocyclic ring, an arylheterocyclic ring, -(CH₂CH₂O)ₚ⁻ or Y₃;

Y₃ is:

T is a -S(O)ₚ⁻; a carbonyl or a covalent bond;

o is an integer from 0 to 2;

Rₗ and Rₘ are independently selected from an alkyl group, an aryl group, or Rₗ and Rₘ taken together with the nitrogen atom to which they are attached are a heterocyclic ring;

T₃ at each occurrence is independently a covalent bond, a carbonyl, an oxygen, -S(O)ₚ⁻ or -N(R₄)R₅;

h is an integer form 1 to 10;
$q_1$ is an integer from 1 to 5;

$R_3$ and $R_4$ are each independently a hydrogen, an alkyl, a cycloalkoxy, a halogen, a hydroxy, an hydroxyalkyl, an alkoxylalkyl, an arylheterocyclic ring, an alkylaryl, an alkylcycloalkyl, an alkylheterocyclic ring, a cycloalkylalkyl, a cycloalkylthio, an arylalklythio, an arylalklythioalkyl, an alkylthioalkyl, a cycloalkenyl, an heterocyclicalkyl, an alkoxy, a haloalkoxy, an amino, an alkylamino, a dialkylamino, an arylamino, a diarylamino, an alkylarylaminio, an alkoxylhaloalkyl, a sulfonic acid, a sulfonic ester, an alkylsulfonic acid, an arylsulfonic acid, an arylalkoxy, an alkylthio, an arythio, a cyano, an aminoalkyl, an aminoaryl, an aryl, an arylalkyl, an alkylaryl, a carboxamido, an alkylcarboxamido, an arylcarboxamido, an amidyl, a carboxyl, a carbamoyl, an alkylcarboxylic acid, an arylcarboxylic acid, an alkylcarbonyl, an arylcarbonyl, an ester, a carboxylic ester, an alkylcarboxylic ester, an arylcarboxylic ester, a sulfonylamido, an alkylsulfonamido, an arylsulfonamido, an alkylsulfonyl, an alkylsulfonyloxy, an arylsulfonil, arylsulphonyloxy, a sulfonic ester, an alkyl ester, an aryl ester, a urea, a phosphoryl, a nitro, $-U_3-V_5$, $V_6$, $-(C(R_0)(R_0))_{k_1}-U_3-V_5$, $-(C(R_0)(R_0))_{k_1}-U_3-V_5$, $-(C(R_0)(R_0))_{k_1}-U_3-V_5$, $-(C(R_0)(R_0))_{k_1}-U_3-C(O)-V_6$, or $R_3$ and $R_4$ taken together with the carbons to which they are attached form a carbonyl, a methanthial, a heterocyclic ring, a cycloalkyl group, an aryl group, an oxime, an imine, a hydrzone, a bridged cycloalkyl group.

R$_5$ and R$_p$ are each independently a hydrogen, an alkyl, a cycloalkoxy, a halogen, a hydroxy, an hydroxyalkyl, an alkoxylalkyl, an arylheterocyclic ring, an alkylaryl, an alkylcycloalkyl, an alkylheterocyclic ring, a cycloalkylalkyl, a cycloalkylthio, an arylalklythio, an arylalklythioalkyl, an alkylthioalkyl, a cycloalkenyl, an heterocyclicalkyl, an alkoxy, a haloalkoxy, an amino, an alkylamino, a dialkylamino, an arylamino, a diarylamino, an alkylarylaminio, an alkoxylhaloalkyl, a sulfonic acid, a sulfonic ester, an alkylsulfonic acid, an arylsulfonic acid, an arylalkoxy, an alkylthio, an arythio, a cyano an aminoalkyl, an aminoaryl, an aryl, an arylalkyl, an alkylaryl, a carboxamido, an alkylcarboxamido, an
arylcarboxamido, an amidyl, a carboxyl, a carbamoyl, an alkylcarboxylic acid, an alkylcarboxylic acid, an alkylcarbonyl, an arylcarbonyl, an ester, a carboxylic ester, an alkylcarboxylic ester, an arylcarboxylic ester, a sulfonamido, an alkylsulfonamido, an arylsulfonamido, an alkylsulfonil, an alkylsulfonyloxy, an arylsulfonil, arylsulphonyloxy, a sulfonic ester, an alkyl ester, an aryl ester, a urea, a phosphoryl, a nitro, -U₃-V₅, V₆, or R₀ and Rₚ taken together with the carbons to which they are attached form a carbonyl, a methanthial, a heterocyclic ring, a cycloalkyl group, an aryl group, an oxime, an imine, a hydrazone a bridged cycloalkyl group,

\[ \text{(1)} \]

\[
\begin{align*}
\text{H}_3\text{C} & \text{ } \text{CH}_3 \\
\text{Z}_5 & \text{ } \text{N} \text{ } \text{O} \\
\text{H}_3\text{C} & \text{ } \text{CH}_3
\end{align*}
\]

or

\[ \text{(2)} \]

\[
\begin{align*}
\text{H}_3\text{C} & \text{ } \text{CH}_3 \\
\text{Z}_5 & \text{ } \text{N} \text{ } \text{O} \\
\text{H}_3\text{C} & \text{ } \text{CH}_3
\end{align*}
\]

U₃ is an oxygen, sulfur or -N(R₉)R₉;
V₅ is -NO or -NO₂ (i.e. an oxidized nitrogen);
k₁ is an integer from 1 to 3;
R₉ is a lone pair of electrons, a hydrogen or an alkyl group;
R₉ is a hydrogen, an alkyl, an aryl, an alkylcarboxylic acid, an arylcarboxylic acid, an alkylcarboxylic ester, an arylcarboxylic ester, an alkylcarboxamido, an arylcarboxamido, an alkylaryl, an alkylsulfanyl, an alkylsulfonil, an alkylsulfonyloxy, an arylsulfonil, an alkylsulfonamido, a carboxamido, a carboxylic ester, an aminoalkyl, an aminoaryl, -(U₃-V₅)(R₀)(R₉), a bond to an adjacent atom creating a double bond to that atom or -(N₂O₂⁻)•M₁⁺, wherein M₁⁺ is an organic or inorganic cation; and
with the proviso that the compound of Formula (I) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group and/or acidic nitrogen group.

In cases where multiple designations of variables which reside in sequence are chosen as a "covalent bond" or the integer chosen is 0, the intent is to denote a single covalent bond connecting one radical to another. For example, E₀ would denote a covalent bond, while E₂ denotes (E-E) and (C(R₄)(R₄))₂ denotes -C(R₄)(R₄)C(R₄)(R₄)-.

In another embodiment, the invention describes compounds of Formula (II):

34
wherein

R_{37} is as defined herein; and

with the proviso that the compound of Formula (II) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group.

In another embodiment, the invention describes compounds of Formula (III):

wherein:

X_{3} and Y_{3} are as defined herein; and

with the proviso that the compound of Formula (III) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group and/or acidic nitrogen group.

In another embodiment, the invention describes compounds of Formula (IV):
wherein:

$X_3$ and $Y_3$ are as defined herein; and

with the proviso that the compound of Formula (IV) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group and/or acidic nitrogen group.

In another embodiment, the invention describes compounds of Formula (V):

wherein:

$X_3$ and $Y_3$ are as defined herein; and

with the proviso that the compounds of Formula (V) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group and/or acidic nitrogen group.

In another embodiment, the invention describes compounds of Formula (VI):

wherein:

$X_3$ and $Y_3$ are as defined herein; and
with the proviso that the compounds of Formula (VI) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group and/or acidic nitrogen group.

In another embodiment, the invention describes angiotensin II antagonist compounds of Formula (VII):

![Structure VI](image)

(VII)

wherein:
- $R_{47}$ is a lower alkyl or $-(C(R_{g})R_{h})_{k}-V_{4}$;
- $X_{3}$, $Y_{3}$, $R_{g}$, $R_{h}$, $V_{4}$, $K$ and $k$ are as defined herein; and
- with the proviso that the compounds of Formula (VII) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group and/or acidic nitrogen group.

In another embodiment, the invention describes compounds of Formula (VIII):

![Structure VIII](image)

(VIII)

wherein:
- $X_{3}$ and $Y_{3}$ are as defined herein; and
- with the proviso that the compounds of Formula (VIII) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group and/or acidic nitrogen group.

In other embodiments of the invention the compound of Formula (I) is an organic nitric oxide enhancing salt of abitesartan, an organic nitric oxide enhancing salt of candesartan, an organic nitric oxide enhancing salt of CV-11974, an organic nitric oxide enhancing salt of elisartan analogue, an organic nitric oxide enhancing salt of embusartan, an organic nitric oxide enhancing salt of enoltasosartan, an organic nitric oxide enhancing salt of

In other embodiments of the invention, the organic nitric oxide enhancing salt of the angiotensin II antagonists of Formula (I) is an organic nitric oxide enhancing salt of abitesartan of Formula (IX), an organic nitric oxide enhancing salt of CV-11974 of Formula (X), an organic nitric oxide enhancing salt of elisartan of Formula (XI), an organic nitric oxide enhancing salt of embusartan of Formula (XII), an organic nitric oxide enhancing salt of enoltasosartan of Formula (XIII), an organic nitric oxide enhancing salt of fonsartan of Formula (XIV), an organic nitric oxide enhancing salt of forasartan of Formula (XV), an organic nitric oxide enhancing salt of glycylllosartan of Formula (XVI), an organic nitric oxide enhancing salt of irbesartan of Formula (XVII), an organic nitric oxide enhancing salt of losartan of Formula (XVIII), an organic nitric oxide enhancing salt of olmesartan of Formula (XIX), an organic nitric oxide enhancing salt of milfasartan of Formula (XX), an organic nitric oxide enhancing salt of ripisartan of Formula (XXII), an organic nitric oxide enhancing salt of tasosartan of Formula (XXIII), an organic nitric oxide enhancing salt of telmisartan of Formula (XXIV), an organic nitric oxide enhancing salt of valsartan of Formula (XXV), a nitric oxide enhancing analogue related to EMD 66684 of Formula (XXVI); a nitric oxide enhancing EXP 3134 of Formula (XXVII); a nitric oxide enhancing MK-996 of Formula (XXVIII); the organic nitric oxide enhancing salt of angiotensin II antagonists of Formula (II) is an organic nitric oxide enhancing salt of eprosartan of Formula (XXIX); and the organic nitric oxide enhancing salt of angiotensin II antagonist of Formula (III) is an organic nitric oxide enhancing salt of saprisartan of Formula (XXX), or an organic nitric oxide enhancing salt of zolasartan of Formula (XXXI),

wherein the compound of Formula (IX) is:
and the compound of Formula (X) is:

and the compound of Formula (XI) is:

and the compound of Formula (XII) is:
and the compound of Formula (XIII) is:

(XII)

and the compound of Formula (XIV) is:

(XIII)

and the compound of Formula (XIV) is:

(XIII)

and the compound of Formula (XV) is:

(XIV)

and the compound of Formula (XVI) is:

(XV)
and the compound of Formula (XVII) is:

and the compound of Formula (XVIII) is:

and the compound of Formula (XIX) is:

and the compound of Formula (XX) is:
and the compound of Formula (XXI) is:

and the compound of Formula (XXII) is:

and the compound of Formula (XXIII) is:
and the compound of Formula (XXIV) is:

and the compound of Formula (XXV) is:

and the compound of Formula (XXVI) is:
and the compound of Formula (XXVII) is:

(XXVII)

and the compound of Formula (XXVIII) is:

(XXVIII)

and the compound of Formula (XXIX) is:

(XXIX)

and the compound of Formula (XXX) is:
and the compound of Formula (XXXI) is:

wherein
\( R_{35} \) is \(-\text{C(O)U}_{3}\text{D}_{1} \) or \(-\text{COOH•Z}_{1} \);
\( R_{36} \) is a hydrogen or \(-\text{H•Z}_{1} \);
nBu is the lower alkyl group \( \text{CH}_{3} \text{-CH}_{2} \text{-CH}_{2} \text{-CH}_{2} \cdot \);
nPr is the lower alkyl group \( \text{CH}_{3} \text{-CH}_{2} \text{-CH}_{2} \cdot \);
iPr is the lower alkyl group \( \text{(CH}_{3} \text{)}_{2} \text{-CH} \cdot \);
OEt is the alkoxy group \(-\text{OCH}_{2} \text{-CH}_{3} \);
\( Z_{1} \) is:

\[
\begin{align*}
(1) & \quad \text{[Diagram of chemical structure]} \\
(2) & \quad \text{[Diagram of chemical structure]} 
\end{align*}
\]
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U₃ and D₁ are as defined herein; and

with the proviso that the compounds of Formula (IX) to Formula (XXXI) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., • or ■) to at least one carboxylic acid group and/or acidic nitrogen group in the compounds of Formula (IX) to Formula (XXXI).

In another embodiment, the organic nitric oxide enhancing salt of Formula (I) is:
L-valine, N-(1-oxopentyl)-N-[[2'-(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl)methyl]-; salt with 3-pyridinecarboxamide, N-[3-(nitrooxy)propyl]- (1:2);
L-valine, N-(1-oxopentyl)-N-[[2'-(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl)methyl]-; salt with 1-propanol, 3-amino-, nitrate (ester) (1:2);
L-valine, N-(1-oxopentyl)-N-[[2'-(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl)methyl]-; salt with 1,3-propanediol, 2-amino-2-[(nitrooxy)methyl]-, dinitrate (ester);
L-valine, N-(1-oxopentyl)-N-[[2'-(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl)methyl]-; salt with 2-propanediol, 3-amino-, dinitrate (ester), (2R)-.

In other embodiments, the organic nitric oxide enhancing compounds that form salts are organic nitrates, organic nitrites, nitroethers, thionitrites, thionitrates, NONOates, heterocyclic nitric oxide donors and nitroxides.

In one embodiment, the organic nitric oxide enhancing salts of angiotensin II antagonist does not contain at least one nitrate ion mole per mole of the angiotensin II antagonist.

Compounds of the invention that have one or more asymmetric carbon atoms may exist as the optically pure enantiomers, pure diastereomers, mixtures of enantiomers, mixtures of diastereomers, racemic mixtures of enantiomers, diastereomeric racemates or mixtures of diastereomeric racemates. It is to be understood that the invention anticipates and includes within its scope all such isomers and mixtures thereof.

Another embodiment of the invention provides the organic nitric oxide enhancing salts of the metabolites of the angiotensin II antagonists. These metabolites include, but are not limited to, degradation products, hydrolysis products, and the like, of the angiotensin II antagonists.
Another embodiment of the invention provides processes for making the novel salts of the invention. The reactions are performed in solvents appropriate to the reagents and materials used are suitable for the transformations being effected. It is understood by one skilled in the art of organic synthesis that the functionality present in the molecule must be consistent with the chemical transformation proposed. This will, on occasion, necessitate judgment by the routinier as to the order of synthetic steps, protecting groups required, and deprotection conditions. Substituents on the starting materials may be incompatible with some of the reaction conditions required in some of the methods described, but alternative methods and substituents compatible with the reaction conditions will be readily apparent to one skilled in the art. The use of sulfur and oxygen protecting groups is well known for protecting thiol and alcohol groups against undesirable reactions during a synthetic procedure and many such protecting groups are known and described by, for example, Greene and Wuts, *Protective Groups in Organic Synthesis*, Third Edition, John Wiley & Sons, New York (1999).

The chemical reactions described herein are generally disclosed in terms of their broadest application to the preparation of the compounds of this invention. Occasionally, the reactions may not be applicable as described to each compound included within the disclosed scope. The compounds for which this occurs will be readily recognized by one skilled in the art. In all such cases, either the reactions can be successfully performed by conventional modifications known to one skilled in the art, e.g., by appropriate protection of interfering groups, by changing to alternative conventional reagents, by routine modification of reaction conditions, and the like, or other reactions disclosed herein or otherwise conventional, will be applicable to the preparation of the corresponding compounds of this invention. In all preparative methods, all starting materials are known or readily prepared from known starting materials.

The salts of the invention are formulated according to well known techniques in the prior art, see for example, Remington's Pharmaceutical Sciences.

The angiotensin II antagonists are either commercially available or can be prepared according to the methods described are described more fully in the literature, such as in Goodman and Gilman, *The Pharmacological Basis of Therapeutics* (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, 13th Edition; and on STN Express, file phar and file registry.

The novel organic nitric oxide enhancing compounds can be synthesized by one skilled in the art using conventional methods. Known methods for linking a nitric oxide
enhancing group to compounds, such as, for example, linking nitrates, thionitrates, nitrites, thionitrites, (i.e. nitrosated and/or nitrosylated compounds), NONOates, heterocyclic nitric oxide donors, and the like are described in the literature. For example, heterocyclic nitric oxide donor compounds are described in WO 99/64417, WO 94/01422; EP 0 574 726 A1, EP 0 683 159 A1; and in J. Med. Chem., 47: 2688-2693 (2004); J. Med. Chem., 47: 1840-1846 (2004); J. Med. Chem., 46: 3762-3765 (2003); J. Med. Chem., 46: 747-754 (2003); Chem Rev., 102: 1091-1134 (2002); J. Med. Chem., 42: 1941-1950 (1999); J. Med. Chem., 41: 5393-5401 (1998); J. Med. Chem., 38: 4944-4949 (1995); Arzneim. Forsch. Drug Res., 47 (II): 847-854 (1997); the disclosures of each of which are incorporated by reference herein in their entirety. The methods of linking the heterocyclic nitric oxide donor group to compounds described in these references can be applied by one skilled in the art to produce any of the organic nitric oxide enhancing compounds described herein. Linking a nitrate group, a thionitrate group, a nitrite group and/or a thionitrite group to a compound can be achieved by the nitrosated and/or nitrosylated of a compound through one or more sites such as oxygen, sulfur and/or nitrogen using conventional methods known to one skilled in the art.

Known methods for nitrosating and/or nitrosylating compounds are described in U.S. Patent Nos. 5,380,758, 5,859,053, 5,703,073 and 6,297,260; and in WO 94/03421, WO 94/04484, WO 94/12463, WO 95/09831, WO 95/19952, WO 95/30641, WO 97/27749, WO 98/09948, WO 98/19672, WO 98/21193, WO 00/51988, WO 00/61604, WO 00/72838, WO 01/00563, WO 01/04082, WO 01/10814, WO 01/12584, WO 01/45703, WO 00/61541, WO 00/61537, WO 02/11707, WO 02/30866 and in Oae et al, Org. Prep. Proc. Int., 15(3):165-198 (1983), the disclosures of each of which are incorporated by reference herein in their entirety. The methods of nitrosating and/or nitrosylating the compounds described in these references can be applied by one skilled in the art to produce any of the nitrosated and/or nitrosylated compounds described herein.

Known methods of linking the nitrooxide group to compounds are described in U.S. Patent Nos. 6,448,267, 6,455,542, 6,759,430, and in WO 2004/050084, WO 03/088961, the disclosures of each of which are incorporated by reference herein in their entirety.

The organic nitric oxide enhancing salts of the angiotensin II antagonists are prepared by the following methods. When the angiotensin II antagonist to be salified is available as free base soluble in an organic solvent, which preferably does not contain hydroxyl groups, for example acetonitrile, ethyl acetate, tetrahydrofuran, and the like, the salt is prepared by dissolving the compound in the solvent at a concentration preferably equal to or higher than 10% w/v, adding the amount of organic nitric oxide enhancing compound corresponding to
the moles of the ionizable groups in the angiotensin II antagonist. The organic nitric oxide enhancing compound is preferably diluted in the same solvent. The salt is generally recovered by filtration and washed with the solvent.

When the angiotensin II antagonist is not very soluble, or is available in the form of a not very soluble salt in the above mentioned solvents, a hydroxylated solvent, such as, for examples, methyl alcohol, ethyl alcohol, water, and the like, can be used.

When the starting angiotensin II antagonist is an inorganic salt, the corresponding base can also be prepared by treatment with a saturated solution of sodium or potassium bicarbonate or carbonate, or with a diluted solution of sodium or potassium hydroxide. The base is then extracted with a suitable organic solvent (for example halogenated solvents, esters, ethers), which is then dried. The organic solution is evaporated and the organic nitric oxide enhancing salt is prepared as described herein.

Compounds contemplated for use in the invention, e.g., organic nitric oxide enhancing salts of angiotensin II antagonists, are, optionally, used in combination with nitric oxide enhancing compounds that release nitric oxide, increase endogeneous levels of nitric oxide or otherwise directly or indirectly deliver or transfer a biologically active form of nitrogen monoxide to a site of its intended activity, such as on a cell membrane in vivo.

Nitrogen monoxide can exist in three forms: NO- (nitroxy1), NO• (nitric oxide) and NO+ (nitrosonium). NO• is a highly reactive short-lived species that is potentially toxic to cells. This is critical because the pharmacological efficacy of NO depends upon the form in which it is delivered. In contrast to the nitric oxide radical (NO•), nitrosonium (NO+) does not react with O2 or O2- species, and functionalities capable of transferring and/or releasing NO+ and NO- are also resistant to decomposition in the presence of many redox metals. Consequently, administration of charged NO equivalents (positive and/or negative) does not result in the generation of toxic by-products or the elimination of the active NO group.

The term "nitric oxide" encompasses uncharged nitric oxide (NO•) and charged nitrogen monoxide species, preferably charged nitrogen monoxide species, such as nitrosonium ion (NO+) and nitroxy1 ion (NO-). The reactive form of nitric oxide can be provided by gaseous nitric oxide. The nitrogen monoxide releasing, delivering or transferring compounds have the structure F-NO, wherein F is a nitrogen monoxide releasing, delivering or transferring group, and include any and all such compounds which provide nitrogen monoxide to its intended site of action in a form active for its intended purpose.

The term "NO adducts" encompasses any nitrogen monoxide releasing, delivering or transferring compounds, including, for example, S-nitrosothiols, nitrites, nitrates, S-
nitrothiols, sydnonimines, 2-hydroxy-2-nitrosohydrozines, (NONOates), (E)-alkyl-2-((E)-hydroxyimino)-5-nitro-3-hexeneamido (FK-409), (E)-alkyl-2-((E)-hydroxyimino)-5-nitro-3-hexeneamides, N-((2Z, 3E)-4-ethyl-2-(hydroxyimino)-6-methyl-5-nitro-3-heptenyl)-3-pyridinecarboxamide (FR 146801), N-nitrosoamines, N-hydroxyl nitrosamines, nitrosamines, diazetine dioxides, oxatriazole 5-imines, oximes, hydroxylamines, N-hydroxyguanidines, hydroxyureas, benzofuroxanes, furoxans as well as substrates for the endogenous enzymes which synthesizes nitric oxide.

Suitable NONOates include, but are not limited to, (Z)-1-(N-methyl-N-(6-(N-methylammoniohexyl)amino))diazen-1-ium-1,2-diolate ("MAHMA/NO"), (Z)-1-(N-(3-ammoniopropyl)-N-(n-propyl)amino)diazen-1-ium-1,2-diolate ("PAPA/NO"), (Z)-1-(N-(3-aminopropyl)-N-(4-(3-aminopropanylammonio)butyl)-amino) diazen-1-ium-1,2-diolate (spermine NONOate or "SPER/NO") and sodium(Z)-1-(N,N-diethylamino)diazene-1,2-diolate (diethylamine NONOate or "DEA/NO") and derivatives thereof. NONOates are also described in U.S. Patent Nos. 6,232,336, 5,910,316 and 5,650,447, the disclosures of which are incorporated herein by reference in their entirety. The "NO adducts" can be mono-nitrosylated, poly-nitrosylated, mono-nitrosated and/or poly-nitrosated at a variety of naturally susceptible or artificially provided binding sites for biologically active forms of nitrogen monoxide.

Suitable furoxanes include, but are not limited to, CAS 1609, C93-4759, C92-4678, S35b, CHF 2206, CHF 2363, and the like.

Suitable sydnonimines include, but are not limited to, molsidomine (N-ethoxycarbonyl-3-morpholinosydnonimine), SIN-1 (3-morpholinosydnonimine) CAS 936 (3-(cis,2,6-dimethylpiperidino)-N-(4-methoxybenzoyl)-sydnonimine, pirsidomine), C87-3754 (3-(cis,2,6-dimethylpiperidino)sydnonimine, linsidomine, C4144 (3-(3,3-dimethyl-1,4-thiazane-4-yl)sydnonimine hydrochloride), C89-4095 (3-(3,3-dimethyl-1,1-dioxo-1,4-thiazane-4-yl)sydnonimine hydrochloride, and the like.

Suitable oximes, include, but are not limited to, NOR-1, NOR-3, NOR-4, and the like.

One group of NO adducts is the S-nitrosothiols, which are compounds that include at least one S-NO group. These compounds include S-nitroso-polypeptides (the term "polypeptide" includes proteins and polynzymo acids that do not possess an ascertained biological function, and derivatives thereof); S-nitrosylated amino acids (including natural and synthetic amino acids and their stereoisomers and racemic mixtures and derivatives thereof); S-nitrosylated sugars; S-nitrosylated, modified and unmodified, oligonucleotides (preferably of at least 5, and more preferably 5-200 nucleotides); straight or branched,
saturated or unsaturated, aliphatic or aromatic, substituted or unsubstituted S-nitrosylated hydrocarbons; and S-nitroso heterocyclic compounds. S-nitrosothiols and methods for preparing them are described in U.S. Patent Nos. 5,380,758 and 5,703,073; WO 97/27749; WO 98/19672; and Oae et al, Org. Prep. Proc. Int., 15(3):165-198 (1983), the disclosures of each of which are incorporated by reference herein in their entirety.

Another embodiment of the invention is S-nitroso amino acids where the nitroso group is linked to a sulfur group of a sulfur-containing amino acid or derivative thereof. Such compounds include, for example, S-nitroso-N-acetylcysteine, S-nitroso-captothi, S-nitroso-N-acetylpenicillamine, S-nitroso-homocysteine, S-nitroso-cysteine, S-nitroso-glutathione, S-nitroso-cysteinyl-glycine, and the like.

Suitable S-nitrosylated proteins include thiol-containing proteins (where the NO group is attached to one or more sulfur groups on an amino acid or amino acid derivative thereof) from various functional classes including enzymes, such as tissue-type plasminogen activator (TPA) and cathepsin B; transport proteins, such as lipoproteins; heme proteins, such as hemoglobin and serum albumin; and biologically protective proteins, such as immunoglobulins, antibodies and cytokines. Such nitrosylated proteins are described in WO 93/09806, the disclosure of which is incorporated by reference herein in its entirety. Examples include polynitrosylated albumin where one or more thiol or other nucleophilic centers in the protein are modified.

Other examples of suitable S-nitrosothiols include:

(i) $\text{HS(C(R_e)(R_f))_mSNO;}$

(ii) $\text{ONS(C(R_e)(R_f))_mR_c;}$ or

(iii) $\text{H_2N-CH(CO_2H)-(CH_2)_m-C(O)NH-CH(CH_2SNO)-C(O)NH-CH_2-CO_2H;}$

wherein $m$ is an integer from 2 to 20;

$R_e$ and $R_f$ are each independently a hydrogen, an alkyl, a cycloalkoxy, a halogen, a hydroxy, an hydroxyalkyl, an alkoxyalkyl, an arylheterocyclic ring, an alkylaryl, an alkylcycloalkyl, an alkylheterocyclic ring, a cycloalkylalkyl, a cycloalkylthio, an arylalkylthio, an arylalkylthioalkyl, an alkylthioalkyl, a cycloalkenyl, an heterocyclicalkyl, an alkoxy, a haloalkoxy, an amino, an alkylamino, a dialkylamino, an arylamino, a diarylamin, an alkylarylamin, an alkoxyhaloalkyl, a sulfonic acid, a sulfonic ester, an alkylsulfonic acid, an arylsulfonic acid, an arylalkoxy, an alklythio, an arylthio, a cyano, an aminoalkyl, an aminoaryl, an aryl, an arylalkyl, an alkylaryl, a carboxamido, an alkylcarboxamido, an arylcarboxamido, an amidyl, a carboxyl, a carbamoyl, an alkylcarboxylic acid, an arylcarboxylic acid, an alkylcarbonyl, an arylcarbonyl, an ester, a carboxylic ester, an
alkylcarboxylic ester, an arylcarboxylic ester, a sulfonamido, an alkylsulfonamido, an arylsulfonamido, an alkylsulfonyl, an alkylsulfonyleoxy, an arylsulfonyl, arylsulphonyloxy, a sulfonic ester, an alkyl ester, an aryl ester, a urea, a phosphoryl, a nitro, -U₃-V₅, V₆, -(C(Rₒ)(Rₚ))k₁-U₃-V₅, -(C(Rₒ)(Rₚ))k₁-U₃-V₆, -(C(Rₒ)(Rₚ))k₁-U₃-C(O)-V₆, or Rₑ and Rₖ taken together with the carbons to which they are attached form a carbonyl, a methanthial, a heterocyclic ring, a cycloalkyl group, an aryl group, an oxime, a hydrazone, a bridged cycloalkyl group,

Rₒ and Rₚ are each independently a hydrogen, an alkyl, a cycloalkoxy, a halogen, a hydroxy, an hydroxyalkyl, an alkoxycarbonyl, an arylheterocyclic ring, an alkylaryl, an alkylcycloalkyl, an alkylheterocyclic ring, a cycloalkylalkyl, a cycloalkylthio, an arylalkylthio, an arylalkylthioalkyl, an alkylthioalkyl, a cycloalkylthio, an heterocyclicalkyl, an alkoxy, a haloalkoxy, an amino, an alkylamino, a dialkylamino, an arylamino, a diarylamino, an alkylarylamino, an aralkylamino, a sulfonic acid, a sulfonic ester, an alkylsulfonic acid, an arylsulfonic acid, an aralkylthio, an alkylthio, an arylthio, a cyano, an aminoalkyl, an aminoaryl, an aryl, an aralkyl, an alkylaryl, a carbamido, an alkylcarboxamido, an arylcarboxamido, an amidyl, a carboxyl, a carbamoyl, an alkylcarboxylic acid, an arylcarboxylic acid, an alkylcarboxyl, an arylcarboxyl, an ester, a carboxylic ester, an alkylcarboxylic ester, an arylcarboxylic ester, a sulfonyl, an alkylsulfonyl, an arylsulfonyl, an alkylsulfonyl, an arylsulphonyloxy, a sulfonic ester, an alkyl ester, an aryl ester, a urea, a phosphoryl, a nitro, -U₃-V₅, V₆, or Rₑ and Rₖ taken together with the carbons to which they are attached form a carbonyl, a methanthial, a heterocyclic ring, a cycloalkyl group, an aryl group, an oxime, a hydrazone, a bridged cycloalkyl group,
$k_1$ is an integer form 1 to 3;

$U_3$ is an oxygen, sulfur- or -$N(R_a)R_i$;

$V_5$ is =NO or =NO$_2$ (i.e. an oxidized nitrogen);

$R_a$ is a lone pair of electrons, a hydrogen or an alkyl group;

$R_i$ is a hydrogen, an alkyl, an aryl, an alkylcarboxylic acid, an arylcarboxylic acid, an alkylcarboxylic ester, an arylcarboxylic ester, an alkylcarboxamido, an arylcarboxamido, an alkylaryl, an alkylsulfanyl, an alkylsulfonyl, an alkylsulfonyloxy, an arylsulfanyl, an arylsulfonyl, arylsulphonyloxy, a sulfonamido, a carboxamido, a carboxylic ester, an aminoalkyl, an aminocarbonyl, -$\text{CH}_2\text{-C}(U_3-V_5)(R_a)(R_i)$, a bond to an adjacent atom creating a double bond to that atom or -(N$_2$O$_2$)$^-$-$M_1^+$, wherein $M_1^+$ is an organic or inorganic cation.

In cases where $R_a$ and $R_i$ are independently a heterocyclic ring or taken together $R_a$ and $R_i$ are a heterocyclic ring, then $R_i$ can be a substituent on any disubstituted nitrogen contained within the radical wherein $R_i$ is as defined herein.

Nitrosothiols can be prepared by various methods of synthesis. In general, the thiol precursor is prepared first, then converted to the S-nitrosothiol derivative by nitrosation of the thiol group with NaNO$_2$ under acidic conditions (pH is about 2.5) which yields the S-nitroso derivative. Acids which can be used for this purpose include aqueous sulfuric, acetic and hydrochloric acids. The thiol precursor can also be nitrosylated by reaction with an organic nitrite such as tert-butyl nitrite, or a nitrosonium salt such as nitrosonium tetrafluoroborate in an inert solvent.

Another group of NO adducts for use in the invention, where the NO adduct is a compound that donates, transfers or releases nitric oxide, include compounds comprising at least one ON-O- or ON-N- group. The compounds that include at least one ON-O- or ON-N-group are preferably ON-O- or ON-N-polypeptides (the term "polypeptide" includes proteins and polyamino acids that do not possess an ascertained biological function, and derivatives thereof); ON-O- or ON-N-amino acids (including natural and synthetic amino acids and their stereoisomers and racemic mixtures); ON-O- or ON-N-sugars; ON-O- or -ON-N- modified or
unmodified oligonucleotides (comprising at least 5 nucleotides, preferably 5-200 nucleotides); ON-O- or ON-N- straight or branched, saturated or unsaturated, aliphatic or aromatic, substituted or unsubstituted hydrocarbons; and ON-O-, ON-N- or ON-C- heterocyclic compounds. Examples of compounds comprising at least one ON-O- or ON-N-group include butyl nitrite, isobutyl nitrite, tert-butyl nitrite, amyl nitrite, isoamyl nitrite, N-nitrosamines, N-nitrosamides, N-nitrosourea, N-nitrosoguanidines, N-nitrosocarbamates, N-acyl-N-nitroso compounds (such as, N-methyl-N-nitrosourea); N-hydroxy-N-nitrosamines, cupferron, alanosine, dopastin, 1,3-disubstituted nitrosiminobenzimidazoles, 1,3,4-thiadiazole-2-nitrosimines, benzothiazole-2(3H)-nitrosimines, thiazole-2-nitrosimines, oligonitroso sydnonimines, 3-alkyl-N-nitroso-sydnonimines, 2H-1,3,4-thiadiazine nitrosimines.

Another group of NO adducts for use in the invention include nitrates that donate, transfer or release nitric oxide, such as compounds comprising at least one O2N-O-, O2N-N- or O2N-S- group. Among these compounds are O2N-O-, O2N-N- or O2N-S- polypeptides (the term "polypeptide" includes proteins and also polyamino acids that do not possess an ascertained biological function, and derivatives thereof); O2N-O-, O2N-N- or O2N-S- amino acids (including natural and synthetic amino acids and their stereoisomers and racemic mixtures); O2N-O-, O2N-N- or O2N-S- sugars; O2N-O-, O2N-N- or O2N-S- modified and unmodified oligonucleotides (comprising at least 5 nucleotides, preferably 5-200 nucleotides); O2N-O-, O2N-N- or O2N-S- straight or branched, saturated or unsaturated, aliphatic or aromatic, substituted or unsubstituted hydrocarbons; and O2N-O-, O2N-N- or O2N-S- heterocyclic compounds. Examples of compounds comprising at least one O2N-O-, O2N-N- or O2N-S- group include isosorbide dinitrate, isosorbide mononitrate, clonitrate, erythrityl tetranitrate, mannitol hexanitrate, nitroglycerin, pentaerythritol tetranitrate, pentritol, propatyl nitrate and organic nitrates with a sulphhydryl-containing amino acid such as, for example SPM 3672, SPM 4757, SPM 5185, SPM 5186 and those disclosed in U.S. Patent Nos. 5,284,872, 5,428,061, 5,661,129, 5,807,847 and 5,883,122 and in WO 97/46521, WO 00/54756 and in WO 03/013432, the disclosures of each of which are incorporated by reference herein in their entirety.

Another group of NO adducts are N-oxo-N-nitrosoamines that donate, transfer or release nitric oxide and are represented by the formula: R"+R""N-N(O-M")-NO, where R" and R"" are each independently a polypeptide, an amino acid, a sugar, a modified or unmodified oligonucleotide, a straight or branched, saturated or unsaturated, aliphatic or aromatic, substituted or unsubstituted hydrocarbon, or a heterocyclic group, and where M"+ is an organic or inorganic cation, such, as for example, an alkyl substituted ammonium cation or a
Group I metal cation.

The invention is also directed to compounds that stimulate endogenous NO or elevate levels of endogenous endothelium-derived relaxing factor (EDRF) *in vivo* or are oxidized to produce nitric oxide and/or are substrates for nitric oxide synthase and/or cytochrome P450. Such compounds include, for example, L-arginine, L-homoarginine, and N-hydroxy-L-arginine, N-hydroxy-L-homoarginine, N-hydroxydebrisoquine, N-hydroxypentamidine including their nitrosated and/or nitrosylated analogs (e.g., nitrosated L-arginine, nitrosylated L-arginine, nitrosated N-hydroxy-L-arginine, nitrosylated N-hydroxy-L-arginine, nitrosated and nitrosylated L-homoarginine), N-hydroxyguanidine compounds, amidoxime, ketoximes, aldoxime compounds, that can be oxidized in vivo to produce nitric oxide. Compounds that may be substrates for a cytochrome P450, include, for example, imino(benzylamo)methylhydroxylamine, imino(((4-methylphenyl)methyl)amino)methylhydroxylamine, imino(((4-methoxyphenyl)methyl)amino)methylhydroxylamine, imino(((4-(trifluoromethyl)phenyl)methyl)amino)methylhydroxylamine, imino(((4-nitrophenyl) methyl)amino)methylhydroxylamine, (butylamo)iminomethylhydroxylamine, imino (propylamo)methylhydroxylamine, imino(pentylamo)methylhydroxylamine, imino (propylamo)methylhydroxylamine, imino ((methylamino)methylhydroxylamine, (cyclopropylamino)iminomethylhydroxylamine, imino-2,1,2,3,4-tetrahydroisoquinolyl methylhydroxylamine, imino(1-methyl(2,1,2,3,4-tetrahydroisoquinolyl)methylhydroxylamine, (1,3-dimethyl(2,1,2,3,4-tetrahydroisoquinolyl))iminomethylhydroxylamine, (((4-chlorophenyl)methyl) amino)iminomethylhydroxylamine, (((4-chlorophenyl)amino)iminomethylhydroxylamine, (4-chlorophenyl)(hydroxyimino)methylamine, and 1-(4-chlorophenyl)-1-(hydroxyimino) ethane, and the like, precursors of L-arginine and/or physiologically acceptable salts thereof, including, for example, citrulline, ornithine, glutamine, lysine, polypeptides comprising at least one of these amino acids, inhibitors of the enzyme arginase (e.g., N-hydroxy-L-arginine and 2(S)-amino-6-boronohexanoic acid), nitric oxide mediators and/or physiologically acceptable salts thereof, including, for example, pyruvate, pyruvate precursors, alpha-keto acids having four or more carbon atoms, precursors of alpha-keto acids having four or more carbon atoms (as disclosed in WO 03/017996, the disclosure of which is incorporated herein in its entirety), and the substrates for nitric oxide synthase, cytokines, adenosin, bradykinin, calreticulin, bisacodyl, and phenolphthalein. EDRF is a vascular relaxing factor secreted by the endothelium, and has been identified as nitric oxide (NO) or a closely related derivative thereof (Palmer et al, *Nature*, 327:524-526 (1987); Ignarro et al, *Proc. Natl. Acad. Sci. USA*, 84:9265-9269 (1987)).
The invention is also directed to nitric oxide enhancing compounds that can increase endogenous nitric oxide. Such compounds, include for example, nitrooxide containing compounds, include, but are not limited to, substituted 2,2,6,6-tetramethyl-1-piperidinolxy compounds, substituted 2,2,5,5-tetramethyl-3-pyrroline-1-oxyl compounds, substituted 2,2,5,5-tetramethyl-1-pyrollidinolxy compounds, substituted 1,1,3,3-tetramethylisindoline-2-yloxy compounds, substituted 2,2,4,4-tetramethyl-1-oxazolidinyl-3-oxyl compounds, substituted 3-imidazolin-1-ylxoy, 2,2,5,5-tetramethyl-3-imidazolin-1-ylxoyl compounds, OT-551, 4-hydroxy-2,2,6,6-tetramethyl-1-piperidinolxy (tempol), and the like. Suitable substituents, include, but are not limited to, aminomethyl, benzoyl, 2-bromoacetamido, 2-(2-bromoacetamido)ethoxy)ethylcarbamoyl, carbamoyl, carboxy, cyano, 5-(dimethylamino)-1-naphthalenesulfonylamido, ethoxyfluorophosphinoxy, ethyl, 5-fluoro-2, 4-dinitroanilino, hydroxy, 2-iodoacetamido, isothiocyanato, isothiocyanatomethyl, methyl, maleimido, maleimidoethyl, 2-(2-maleimidoethoxy)ethylcarbamoyl, maleimidomethyl, maleimido, oxo, phosphonoxy, and the like.

The invention is also based on the discovery that compounds and compositions of the invention may be used in conjunction with other therapeutic agents for co-therapies, partially or completely, in place of other therapeutic agents, such as, for example, aldosterone antagonists, α-adrenergic receptor agonists, α-adrenergic receptor antagonists, angiotensin II antagonists, angiotensin-converting enzyme (ACE) inhibitors, antidiabetic compounds, antihyperlipidemic compounds, antimicrobial compounds, antioxidants, antithrombotic and vasodilator compounds, β-adrenergic antagonists, calcium channel blockers, carbonic anhydrase inhibitors, digitalis, diuretics, endothelin antagonists, hydralazine compounds, H₂ receptor antagonists, neutral endopeptidase inhibitors, nonsteroidal antiinflammatory compounds (NSAIDs), phosphodiesterase inhibitors, potassium channel blockers, platelet reducing agents, prostaglandins, proton pump inhibitors, renin inhibitors, selective cyclooxygenase-2 (COX-2) inhibitors, steroids, and combinations of two or more thereof. The therapeutic agent may optionally be nitrosated and/or nitrosylated and/or contain at least one heterocyclic nitric oxide donor group and/or at least one nitrooxide group.

Suitable aldosterone antagonists include, but are not limited to, canrenone, potassium canrenoate, drospirenone, spironolactone, eplerenone (INSPIRA®), epoxymexrenone, fadrozole, pregn-4-ene-7,21-dicarboxylic acid, 9,11-epoxy-17-hydroxy-3-oxo, γ-lactone, methyl ester, (7α,11α,17β.); pregn-4-ene-7,21-dicarboxylic acid, 9,11-epoxy-17-hydroxy-3-oxo-dimethyl ester, (7α,11α,17β.); 3'H-cyclopropa(6,7)pregna-4,6-diene-21-carboxylic acid, 9,11-epoxy-6,7-dihydro-17-hydroxy-3-oxo-, γ-lactone, (6β,7β,11α,17β.); pregn-4-ene-7,21-
dicarboxylic acid, 9,11-epoxy-17-hydroxy-3-oxo-, 7-(1-methylethyl) ester, monopotassium salt, (7α,11α,17β)-; pregn-4-ene-7,21-dicarboxylic acid, 9,11-epoxy-17-hydroxy-3-oxo-, 7-methyl ester, monopotassium salt, (7α,11α,17β)-; 3'H-cyclopropa(6,7)pregna-1,4,6-triene-21-carboxylic acid, 9,11-epoxy-6,7-dihydro-17-hydroxy-3-oxo-, γ-lactone, (6β,7β,11α)-; 3'H-cyclopropa(6,7)pregna-4,6-diene-21-carboxylic acid, 9,11-epoxy-6,7-dihydro-17-hydroxy-3-oxo-, methyl ester, (6β,7β,11α,17β)-; 3'H-cyclopropa (6,7)pregna-4,6-diene-21-carboxylic acid, 9,11-epoxy-6,7-dihydro-17-hydroxy-3-oxo-, monopotassium salt, (6β,7β,11α,17β)-; 3'H-cyclopropa(6,7)pregna-1,4,6-triene-21-carboxylic acid, 9,11-epoxy-6,7-dihydro-17-hydroxy-3-oxo-, γ-lactone, (6β,7β,11α,17β)-; pregn-4-ene-7,21-dicarboxylic acid, 9,11-epoxy-17-hydroxy-3-oxo-, γ-lactone, ethyl ester, (7α,11α,17β)-; pregn-4-ene-7,21-dicarboxylic acid, 9,11-epoxy-17-hydroxy-3-oxo-, γ-lactone, 1-methylethyl ester, (7α,11α,17β)-; RU-28318, and the like. Suitable aldosterone antagonists are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, 13th Edition; and on STN Express, file phar and file registry.

In some embodiments the aldosterone antagonist is eplerenone or spironolactone (a potassium sparing diuretic that acts like an aldosterone antagonist). In more particular embodiments eplerenone is administered in an amount of about 25 milligrams to about 300 milligrams as a single dose or as multiple doses per day; the spironolactone is administered in an amount of about 25 milligrams to about 150 milligrams as a single dose or as multiple doses per day.

Suitable α-adrenergic receptor agonists, including, but are not limited to, agmatine, p-aminoclonidine, apraclonidine (IOPIpine®), 2-(arylamino) imidazolidine derivatives, azepoxole, azepin derivatives, such as for example, 2-amino-6-alkyl-4,5,7,8-tetrahydro-6H-thiazolo-(5,4,d) azepine, 2-amino-6-ethyl-4,5,7,8-tetrahydro-6H-thiazolo-(5,4,d) azepine, 2-amino-6-ethyl-4,5,7,8-tetrahydro-6H-oxazolo-(5,4,d) azepine, and the like; brimonidine, clonidine, clonidine derivatives, detomidine, dexmedetomidine, dipivefrin, dipivalylepinephrine, epinephrine, guanabenz, guanfacine, imidazolidine derivatives, such as, for example, 5-bromo-6-(2-imidazolidine-2-ylamino)quinoxaline, and the like; p-iodoclonidine, medetomidine, methoxamine (VASOXYL®), mepheterniene, metaraminol (ARAMINE®), methyldopa, mitodrine, napazoline (PRIVINE®, NAPHCON®), norepinephrine, oxymetazoline (AFRIN®, OCUCLEAR®), phenylepinephrine (NEOSYNEPHRINE®), rilmenidine, tetrahydrozoline (TYZINE®, VISINE®), tramazoline, xylazine, xylometazoline (OTRIVIN®), B-HT 920 (6-allyl-2-amino-5,6,7,8-tetrahydro-4H-
thiazolo(4,5-d)-azepine, B-HT 933 and UK 14,304, and the like. Suitable α-adrenergic receptor agonists are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, (1996); Merck Index on CD-ROM, 13th Edition; STN Express, file phar and file registry, the disclosures of each of which are incorporated by reference herein in their entirety.

In some embodiments the α-adrenergic receptor agonists are aminoclonidine, apraclonidine (IOPIDINE®), brimonidine, clonidine and clonidine derivatives.

Suitable α-adrenergic receptor antagonists receptor antagonists include, but are not limited to, phentolamine, tolazoline, idazoxan, deriglidole, RX 821002, BRL 44408, BRL 44409, BAM 1303, labelotel, ifenprodil, rauwolscine, corynathine, rauascine, tetrahydroalstonine, apoyohimbine, akuammigine, β-yohimbine, yohimbol, yohimbine, pseudoyohimbine, epi-3α-yohimbine, 10-hydroxy-yohimbine, 11-hydroxy-yohimbine, tamsulosin, benoxathian, atipamezole, BE 2254, WB 4101, HU-723, tedisamil, mirtazipine, setiptiline, reboxetine, delequamine, naftopil, saterinone, SL 89.0591, ARC 239, urapidil, 5-methylurapidil, monatepi, haloperidol, indoramin, SB 216469, moxisylyte, trazodone, dapiprazole, efaroxan, Recordati 15/2739, SNAP 1069, SNAP 5089, SNAP 5272, RS 17053, SL 89.0591, KMD 3213, spiperone, AH 11110A, chloroethyloclonidine, BMY 7378, niguldipine, and the like. Suitable alpha-adrenergic receptor antagonists are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, Thirteenth Edition; and on STN Express, file phar and file registry.

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Suitable angiotensin II antagonists are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-
Hill, 1995; and the Merck Index on CD-ROM, 13th Edition; and on STN Express, file phar and file registry.

In some embodiments the angiotensin II antagonists are candesartan, eprosartan, irbesartan, losartan, omlesartan, telmisartan or valsartan. In more particular embodiments the candesartan is administered as candesartan cilexetil in an amount of about 15 milligrams to about 100 milligrams as a single dose or as multiple doses per day; the eprosartan, is administered as eprosartan mesylate in an amount of about 400 milligrams to about 1600 milligrams as a single dose or as multiple doses per day; the irbesartan is administered in an amount of about 75 milligrams to about 1200 milligrams as a single dose or as multiple doses per day; the losartan is administered as losartan potassium in an amount of about 25 milligrams to about 100 milligrams as a single dose or as multiple doses per day; the omlesartan is administered as omlesartan medoxomil in an amount of about 5 milligrams to about 40 milligrams as a single dose or as multiple doses per day; the telmisartan is administered in an amount of about 20 milligrams to about 80 milligrams as a single dose or as multiple doses per day; the valsartan is administered in an amount of about 80 milligrams to about 320 milligrams as a single dose or as multiple doses per day.

Suitable angiotensin-converting enzyme inhibitors (ACE inhibitors) include, but are not limited to, alacepril, benazepril (LOTENSIN®, CIBACE®), benazeprilat, captopril, ceronapril, cilazapril, delapril, duinapril, enalapril, enalaprilat, fasidotril, fosinopril, fosinoprilat, gemopatrilat, glycopril, idrapril, imidapril, lisinopril, moexipril, moveltipril, naphthopidil, omapatriлат, pentopril, perindopril, perindoprilat, quinapril, quinaprilat, ramipril, ramiprilat, rentipril, saralasin acetate, spirapril, temocapril, tramadol, trandolapril, urapidil, zofenopril, acylmercapto and mercaptoalkanoyl pralines, carboxyalkyl dipeptides, carboxyalkyl dipeptide, phosphinylalkanoyl pralines, registry no.796406, AVE 7688, BP1.137, CHF 1514, E 4030, ER 3295, FPL-66564, MDL 100240, RL 6134, RL 6207, RL 6893, SA 760, S-5590, Z 13752A, and the like. Suitable angiotensin-converting enzyme inhibitors are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, Twelfth Edition, Version 12:1, 1996; and on STN Express, file phar and file registry.

In some embodiments the angiotensin-converting enzyme inhibitors are benazepril, captopril, enalapril, fosinopril, lisinopril, moexipril, quinapril, ramipril, tramadol, or trandolaprilat. In more particular embodiments the benazepril is administered as benazepril hydrochloride in an amount of about 5 milligrams to about 80 milligrams as a single dose or
as multiple doses per day; the captopril is administered in an amount of about 12.5 milligrams to about 450 milligrams as a single dose or as multiple doses per day; the enalapril is administered as enalapril maleate in an amount of about 2.5 milligrams to about 40 milligrams as a single dose or as multiple doses per day; the fosinopril is administered as fosinopril sodium in an amount of about 5 milligrams to about 60 milligrams as a single dose or as multiple doses per day; the lisinopril is administered in an amount of about 2.5 milligrams to about 75 milligrams as a single dose or as multiple doses per day; the moexipril is administered as moexipril hydrochloride in an amount of about 7.5 milligrams to about 45 milligrams as a single dose or as multiple doses per day; the quinapril is administered as quinapril hydrochloride in an amount of about 5 milligrams to about 40 milligrams as single or multiple doses per day; the ramipril hydrochloride in an amount of about 1.25 milligrams to about 40 milligrams as single or multiple doses per day; the trandolapril is administered as in an amount of about 0.5 milligrams to about 4 milligrams as single or multiple doses per day; the trandolaprilat is administered as in an amount of about 0.5 milligrams to about 4 milligrams as single or multiple doses per day.

Suitable antidiabetic compounds include but are not limited to, acarbose, acetoheaxamide, buformin, carbutamide, chlorpropamide, glibornuride, gliclazide, glimepiride, glipizide, gliquidone, glitazepid, glyburide, glybuthiazol(e), glybuzole, glyhexamide, glymidine, glypinamide, insulin, metformin, miglitol, nateglinide, phenbutamide, phenformin, pioglitazone, repaglinide, rosiglitazone, tolazamide, tolbutamide, tolcyclamide, troglitazone, voglibose, and the like. Suitable anti-diabetic compounds are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, Thirteenth Edition; and on STN Express, file phar and file registry.

Suitable anti-hyperlipidemic compounds include, but are not limited to, statins or HMG-CoA reductase inhibitors, such as, for example, atorvastatin (LIPITOR®), bervastatin, cerivastatin (BAYCOL®), dalvastatin, fluviodastatin (Sandoz XU-62-320), fluvasatin, glenvastatin, lovastatin (MEVACOR®), mevastatin, pravastatin (PRAVACHOL®), rosuvastatin (CRESTRO®), simvastatin (ZOCOR®), velostatin (also known as synvinolin), VYTORIN™ (ezetimibe/simvastatin), GR-95030, SQ 33,600, BMY 22089, BMY 22,566, CI 980, and the like; gemfibrozil, cholesteryamine, colestipol, niacin, nicotinic acid, bile acid sequestrants, such as, for example, cholestyramine, coleselvalam, colestipol, poly(methyl-(3-trimethylaminopropyl) imino-trimethylene dihalide) and the like; probucol; fibrac acid agents or fibrates, such as, for example, bezafibrate (Bezalip™), beclobrate, binifibrate, ciprofibrate,
clinofibrate, clofibrate, efofibrate, fenofibrate (Lipidil™, Lipidil Micro™), gemfibrozil (Lopid™), nicofibrate, pirifibrate, ronifibrate, simofibrate, theofibrate and the like; cholesterol ester transfer protein (CETP) inhibitors, such as for example, CGS 25159, CP-529414 (torcetrapib), JTT-705, substituted N-[3-(1,1,2,2-tetrafluoroethoxy)benzyl]-N-(3-phenoxyphenyl)-trifluoro-3-amino-2-propanols, N,N-disubstituted trifluoro-3-amino-2-propanols, PD 140195 (4-phenyl-5-tridecyl-4H-1,2,4-triazole-3-thiol), SC-794, SC-795, SCH 58149, and the like.

In some embodiments the anti-hyperlipidemic compounds are atorvastatin, fluvastatin, lovastatin, pravastatin, rosuvastatin or simvastatin. In more particular embodiments the atorvastatin is administered in an amount of about 10 milligrams to about 80 milligrams as a single dose or as multiple doses per day; the fluvastatin is administered in an amount of about 20 milligrams to about 80 milligrams as a single dose or as multiple doses per day; the lovastatin is administered in an amount of about 10 milligrams to about 80 milligrams as a single dose or as multiple doses per day; the pravastatin is administered in an amount of about 10 milligrams to about 80 milligrams as a single dose or as multiple doses per day; the rosuvastatin is administered in an amount of about 5 milligrams to about 40 milligrams as a single dose or as multiple doses per day; the simvastatin is administered in an amount of about 5 milligrams to about 80 milligrams as a single dose or as multiple doses per day.

Suitable antimicrobial compounds, include, but are not limited to, acediasulfone, acetate, acetyl sulfametossipirazine, acetyl sulfamethoxypyrazine, acranil, albendazole, alecidine, amatadine, ambazone, aminocillin, amikacin, p-aminosalicylic acid, p-aminosalicylic acid hydrazine, amoxicillin, ampicillin, anisomycin, apacillin, apicyclin, apramycin, arbekacin, argininsa, aspoxicillin, azadamfenicol, azidocillin, azithromycin, azlocillin, aztreonam, bacampicillin, bacitracin, benzoylpenicillin, benzyl penicillin acid, benzyl sulfamide, bicozamycin, bipenam, brodimprom, capreomycin, carbenicillin, carbomycin, cafazedone, carindacillin, carumonam, cefcapene pivoxil, cefaclor, cefadroxil, cefafroxil, cefamandole, cefatamet, cefatrizine, cefazedone, cefazolin, cefbuperazone, cefclidin, cefdinir, cefditoren, cefixime, cefmenoxime, cefmetazole, cefminox, cefodizime, cefonicid, cefoperazone, ceforanide, cefotaxime, cefotetan, cefotiam, cefoxitin, cefozopran, cefpimizole, cefpiramide, cefpirome, cepodoxime proxetil, cefprozil, cefroxadine, cefsulodin, cefazidine, cefteram, ceftezole, ceftributen, ceftiofur, ceftrizoxime, ceftriaxone, cefuroxime, cefuzonam, cepachetrelute sodium, cephradin, cephalaxin, cephaloglycin, cephaloridine, cephalosporin C, cephalothin, cepahaprin sodium, cephradine, chibrorifamycin, chloramphenicol, chlorotetracycline, cinoxacin, ciprofloxacin, clariromycin, clavulanic acid,
clinafloxacín, clindamycín, clofazimín, clofocatal, clometocilín, clomocyclín, cloxacilín, cloxyquin, colistín, cyclacilín, cycloserín, danoflaxcín, dapsone, deoxycyclín, deoxydihydrostreptomycín, dibekacín, dicloxacilín, diflaxcín, dihydrostreptomycín, dimetridazole, dimenazène, dirırtomicín, duramycín, efomithine, enrofloxacín, enviomycin, epicilín, erythromycín, etacillín, ethambutol, ethionamid, famcyclovir, fenbecilín, fleroxácin, flomoxef, floxacilín, flumequín, n-formamidoylthienamycin, furonazide, fortimycín, furazolium clorid, gentamycín, glycoxázide, gramicidín, grepafloxacín, guacuteclín, halofuginone, hetacilín, homidium, hydroxyl-stilbamidina, ibostamycín, imidocarb, imipenem, ipronidazole, isoniázide, josámycín, inosine, kanamycin, lauroguañidina, lenampilícín, lincomycín, lomefloxacín, loracarbef, lymecyclín, mafenide, mebendazolé, mecloclín, meropenem, metampicilín, metacilín, metaclyclín, methicilín sodíum, metronidazolé, 4'-(methylsulfamoyl) sulfanilamide, mezlocilín, mezicilín, micronomicín, midécamycín A₁, minocyclín, mioacilín, miokamacín, morfazamidina, moxalactam, mupirocin, myxín, nadifloxacín, nalidixic acid, negamycín, neomycín, netilmicín, nifurfóline, nifurpinirló, nifurprázine, nimorazolé, nitroxoline, norfloxacín, novobicín, ofloxacín, oleandomycín, opinazidé, oxacilín, oxophenarsine, oxolinic acid, oxytetracycline, panipenam, paromycín, pazufloxacín, pefloxacín, penicilín G potassium salt, penicilín N, penicilín O, penicilín V, penethamate hydroiodide, pentamidina, phenamidina, phenethicilín potassium salt, phenyl aminosalicylate, pipacyclín, pipemidic acid, pipéralicín, pirlimycín, piromidic acid, pivampicilín, pivefalexín, polimyixin B, proflorimycín, propamidina, propícilín, protoniamidina, puraltadone, puromycín, pyrazinamidé, pyrithromycín, quinacilín, quinacrine, quinapyramine, quintine, ribostamycín, rifabutíne, rifamidé, rifampín, rifamycín, rifanpin, rifapentine, rifaxymine, ritipeném, rokitamycín, rolitetracycline, rosamycín, rufloxacín, salazosulfadimidina, salinazid, sancycline, sarafloxacín, sedacamycin, secnidazole, sismycín, sparfloxacín, spectinomycin, spiramycín, spirámicin I, spirámicin II, spirámicin III, stilbamidina, streptomycín, streptonicidizid, sulbactam, sulbenicilín, succisulfone, sulfanilamide, sulfabenzamidina, sulfacetamidina, sulfachloropyridazina, sulfachrysoidina, sulfactylin, sulfadiazina, sulfadicamidina, sulfadimetoxine, sulfadoxine, sulfadrazone, sulfacetidol, sulfafenazol, sulfaguanidina, sulfaguanolé, sulfalene, sulfamerazine, sulfameter, sulfamethazine, sulfamethizolé, sulfamethomidina, sulfamethoxazolé, sulfamethoxypyridazina, sulfamethylicolazol, sulfamethethylazolé, sulfametrolé, sulfamidochrysoidina, sulfamoxolé, sulfanilamidina, 4-sulfanilamido salicylic acid, 4-4'-sulfanilylbenzylamine, p-sulfanilylbenzylamine, 2-p-sulfinylanilinoethanol, sulfanilylurea, sulfoniazide, sulfaperine,
sulfaphenazole, sulfaproxyline, sulfapyrazine, sulfapyridine, sulfathiazole, sulfathidole, sulfathiourea, sulfisomidine, sulfasomizole, sulfasymazine, sulfisoxazole, 4,4'-sulfinylidianieline, N4-sulfanilylsulfanilamide, N-sulfanilyl-3,4-xylamide, sulfamicillin, talampicillin, tambutol, taurolidine, teicoplanin, temocillin, tetracycline, tetroxoprim, thiabendazole, thiazolsulfone, tibezonium iodide, ticarcillin, tigemonam, tinidazole, tobramycin, tosufloxacin, trimethoprim, troleandomycin, trospasmycin, trovafloxacin, tubercidine, miokamycin, oleandomycin, troleandomycin, vancomycin, verazide, viomycin, virginiamycin, zalcitabine, PA-1806 and PA-2794, and the like. Suitable antimicrobial compounds are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, (1996); Merck Index on CD-ROM, 13th Edition; STN Express, file phar and file registry, the disclosures of each of which are incorporated by reference herein in their entirety.

In some embodiments the antimicrobial compound amikacin, azithromycin, azetreonam, bacitracin, carbencillin, cefazolin, cefoxitin, cephaloridine, chibrorifamycin, chloramphenicol, colistin, duramycin, n-formamidothienamycin, gentamycin, gramicidin, kanamycin, neomycin, penicillin G, polymyxin B, sisomicin, tetracyclines, tigecycline, tobramycin, vancomycin, PA-1806 and PA-2794.

In other embodiments the antimicrobial compound is an antiviral compound, including but not limited to, acyclovir, amatadine, cidofovir, cytarabine, didanosine, dideoxyadenosine, edoxudine, famciclovir, floxuridine, gancyclovir, idoxuridine, indanavir, kethoxal, lamivudine, MADU, penciclovir, podophyllotoxin, ribavirine, rimantadine, saquinavir, sorivudine, stavudine, trifuridine, valacyclovir, vidarabine, xenazoic acid, zalcitabine, zidovudine, and the like.

Suitable antioxidants include, but are not limited to, small-molecule antioxidants and antioxidant enzymes. Suitable small-molecule antioxidants include, but are not limited to, hydralazine compounds, glutathione, vitamin C, vitamin E, cysteine, N-acetyl-cysteine, β-carotene, ubiquinone, ubiquinol-10, tocopherols, coenzyme Q, superoxide dismutase mimetics, such as, for example, 2,2,6,6-tetramethyl-1-piperidinylxoxo (TEMPO), DOXYL, PROXYL nitroxide compounds; 4-hydroxy-2,2,6,6-tetramethyl-1-piperidinylxoxo (Tempol), M-40401, M-40403, M-40407, M-40419,M-40484, M-40587, M-40588, and the like. Suitable antioxidant enzymes include, but are not limited to, superoxide dismutase, catalase, glutathione peroxidase, NADPH oxidase inhibitors, such as, for example, apocynin, aminoguanidine, ONO 1714, S17834 (benzo(b)pyran-4-one derivative), and the like; xanthine oxidase inhibitors, such as, for example, allopurinol, oxypurinol, amflutizole,
diethylidithiocarbamate, 2-styrylchromones, chrysin, luteolin, kaempferol, quercetin, myricetin, isorhamnetin, benzophenones such as 2,2',4,4'-tetrahydroxybenzophenone, 3,4,5,2',3',4'-hexahydroxybenzophenone and 4,4'-dihydroxybenzophenone; benzothiazinone analogues such as 2-amino-4H-1,3-benzothiazine-4-one, 2-guanidino-4H-1,3-benzothiazin-4-one and rhodamine; N-hydroxyguanidine derivative such as, PR5 (1-(3, 4-dimethoxy-2-chlorobenzylideneamino)-3-hydroxyguanidine); 6-formylpterin, and the like. The antioxidant enzymes can be delivered by gene therapy as a viral vector and/or a non-viral vector. Suitable antioxidants are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, Thirteenth Edition; and on STN Express, file phar and file registry.

In some embodiments the antioxidants are apocynin, hydralazine compounds and superoxide dimutase mimetics.

Suitable antithrombotic and vasodilator compounds include, but are not limited to, abciximab, acetermin, acetylsalicylic acid, argatroban, benemid, benfurodil, benziodarone, betaistine, bisaramil, brovincamine, bufeniode, citicoline, clobenfurul, clopidogrel, cyclandelate, dalteparin, dipyriramol, droprenilamine, enoxaparin, fendiline, ifenprodil, iloprost, indobufen, isobogrel, isoaxsuprine, heparin, lamifiban, midrodine, nadroparin, nicotinyl alcohol, nylidrin, ozagrel, perhexiline, phenylpropanolamine, prenylamine, papaveroline, reviparin sodium salt, ridogrel, sulocitidil, tinoferidine, tizaparin, trifusal, vintoperol, xanthinal niacinate, and the like. Suitable antithrombotic and vasodilator compounds are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, Thirteenth Edition; and on STN Express, file phar and file registry.

Suitable β-adrenergic antagonists include, but are not limited to, acebutolol, alpenrolol, amosulalol, arotinolol, atenolol, befunolol, betaxolol, betvantolol, bisoprolol, bopindolol, bucinodolol, bumomolol, bufetolol, bufuralol, bunitrolol, bupranolol, butofilolol, carazolol, capsinolol, catecolol, carvedilol (COREG®), celiprolol, cetamolol, cindolol, cloranolol, dilevalol, diprafenone, epanolol, esentilide, esmolol, esprolol, hedroxalol, indenolol, labetalol, landiolol, lanisolol, levobunolol, mepindolol, methylpranol, metindol, metipranolol, metrizoranolol, metoprolol, moproolol, nadolol, nadoxolol, nebulonol, nifenalol, nipradilol, oxprenolol, penbutolol, pindolol, practolol, pronethalol, propranolol, sotalol, sotalolnadolol, sulfinalol, taliprolol, talinolol, tertatolol, tilisolol, timolol, toliprolol, tomalolol, trimepranol, xamoterol, xibenolol, 2-(3-1,1-dimethylethyl)-amino-2-hydroxypropoxy)-3-pyridenecarbonitriHCl, 1-butylamino-3-(2,5-dichlorophenox)-2-

In some embodiments the β-adrenergic antagonists are atenolol, bisoprolol, carvedilol, metoprolol, nebivolol, propranolol or timolol. In more particular embodiments the atenolol is administered in an amount of about 50 milligrams to about 200 milligrams as a single dose or as multiple doses per day; the bisoprolol is administered as bisoprolol fumarate in an amount of about 2.5 milligrams to about 30 milligrams as a single dose or as multiple doses per day; the carvedilol is administered in an amount of about 3.125 milligrams to about 200 milligrams as a single dose or as multiple doses per day; the metoprolol is administered as metoprolol tartarate or metoprolol succinate in an amount of about 25 milligrams to about 300 milligrams as a single dose or as multiple doses per day; the nebivolol is administered as nebivolol hydrochloride in an amount of about 2.5 milligrams to about 20 milligrams as a single dose or as multiple doses per day; the propranolol is administered as propranolol hydrochloride in an amount of about 40 milligrams to about 240 milligrams as a single dose or as multiple doses per day; the timolol is administered as timolol maleate in an amount of about 10 milligrams to about 30 milligrams as a single dose or as multiple doses per day.

Suitable calcium channel blockers include, but are not limited to, amlodipine (NORVASC®), anipamil, aranidipine, amrinone, azelnidipine, barnidipine, benycyclane, benidipine, bepridil, cilnidipine, cinnarizine, clentiazem, diltiazem, dotarizine, efondipine, elgodipine, fantofarone, felodipine, fendiline, flunarizine, fluspirilene, furnidipine, gallopamil, ipenoxazine, isradipine, lacidipine, lemilidipine, lercanidipine, lomerizine, manidipine, mibebradil, monatepl, nicardipine, nifedipine, niguldipine, niludipine, nilvadipine, nimodipine, nisoldipine, nitrendipine, nivaldipine, oxodipine, verhexilene, phenytoin, phenytoxylidamine, prandipine, ranolazine, ryosidine, semotiadil, tamolarizine, temiverine hydrochloride, terodipine, tiapamil, vatanidipine hydrochloride, verapamil, ziconotide, AE-0047, CAI, JTV-519, CHF-1521, L-651582, NS-7, NW-1015, RO-2933, SB-237376, SL-34.0829-08, S-312d, SD-3212, TA-993, YM-430, and the like. Suitable calcium channel blockers are described more fully in the literature, such as in Goodman

In some embodiments the calcium channel blockers are amlodipine, diltiazem, isradipine, nicardipine, nifedipine, nimodipine, nisoldipine, nitrendipine, verapamil.

Suitable carbonic anhydrase inhibitors, include, but are not limited to, acetazolamide, brinzolamide, dorzolamide, ethoxzolamide, 6-hydroxy-2-benzothiazolesulfonamide, methazolamide, thiophene sulfonamide, an aromatic sulfonamide, an ester of 6-hydroxy-2-benzothiazolesulfonamide, an ester of 5-hydroxy-2-benzothiazolesulfonamide, and the like. Suitable carbonic anhydrase inhibitors are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, 13th Edition; and on STN Express, file phar and file registry.

In some embodiments the carbonic anhydrase inhibitors are brinzolamide and dorzolamide.

Suitable digitals include but are not limited to, digoxin and digoxitin. In some embodiments the digoxin is administered to achieve a steady state blood serum concentration of at least about 0.7 nanograms per ml to about 2.0 nanograms per ml.

Suitable diuretics include but are not limited to, thiazides (such as, for example, althiazide, bendroflumethiazide, benzclortriazide, benzhydrochlorothiazide, benzthiazide, buthiazide, chlorothiazide, cyclopenethiazide, cyclothiazide, epithiazide, ethiazide, hydrobenzthiazide, hydrochlorothiazide, hydroflumethiazide, methylclothiazide, methylcyclothiazide, penflutazide, polythiazide, teclothiazide, trichlormethiazide, triflumethiazide, and the like); alilusem, ambuside, amiloride, aminometradine, azosemide, bemetizide, bumetanide, butazolamide, butizide, canrenone, carperitide, chloraminophenamide, chlorazanil, chlormerodrin, chlorthalidone, cicletanide, clofenamide, clopamide, clorexolone, conivaptan, daglitril, dichlorphenamide, disulfamide, ethacrynic acid, ethoxzolamide, etozolon, fenoldopam, fenquizone, furosemide, indapamide, mebutizide, mefruside, meralluride, mercaptopmerin sodium, mercumallylic acid, mersanyl, methazolamide, meticaine, metolazone, moazavaptan, muzolimine, N-(5-1,3,4-thiadiazol-2-yl)acetamide, nesiritide, pamabrom, paraflutizide, piretanide, protheobromine, quinetahzone, scoparius, spironolactone, theobromine, ticrynafen, torsemide, torvaptan, triamterene, tripamide, ularitide, xipamide or potassium, AT 189000, AY 31906, BG 9928, BG 9791, C 2921, DTI 0017, JDL 961, KW 3902, MCC 134, SLV 306, SR 121463, WAY 140288, ZP
120, and the like. Suitable diuretics are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, 13th Edition; and on STN Express, file phar and file registry.

Depending on the diuretic employed, potassium may also be administered to the patient in order to optimize the fluid balance while avoiding hypokalemic alkalosis. The administration of potassium can be in the form of potassium chloride or by the daily ingestion of foods with high potassium content such as, for example, bananas or orange juice. The method of administration of these compounds is described in further detail in U.S. Patent No. 4,868,179, the disclosure of which is incorporated by reference herein in its entirety.

In some embodiments the diuretics are amiloride, furosemide, chlorthalidone, hydrochlorothiazide or triamterene. In more particular embodiments the amiloride is administered as amiloride hydrochloride in an amount of about 5 milligrams to about 15 milligrams as a single dose or as multiple doses per day; the furosemide is administered in an amount of about 10 milligrams to about 600 milligrams as a single dose or as multiple doses per day; the chlorthalidone is administered in an amount of about 15 milligrams to about 150 milligrams as a single dose or as multiple doses per day; the hydrochlorothiazide is administered in an amount of about 12.5 milligrams to about 300 milligrams as a single dose or as multiple doses per day; the triamterene is administered in an amount of about 35 milligrams to about 225 milligrams as a single dose or as multiple doses per day.

Suitable endothelin antagonists include, but are not limited to, atrasentan, bosentan, darusentan, endothelin, enrasentan, sitaxsentan, sulfonamide endothelin antagonists, tezosentan, BMS 193884, BQ-123, SQ 28608, and the like. Suitable endothelin antagonists are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, Thirteenth Edition; and on STN Express, file phar and file registry.

Suitable hydralazine compounds include, but are not limited to, compounds having the formula:

\[
\begin{array}{c}
\text{R}_1 \\
\text{R}_2 \\
\text{R}_3 \\
\text{R}_4 \\
\end{array}
\]

wherein \(a\), \(b\) and \(c\) are independently a single or double bond; \(R_1\) and \(R_2\) are each independently a hydrogen, an alkyl, an ester or a heterocyclic ring, wherein alkyl, ester and
heterocyclic rind are as defined herein; R₃ and R₄ are each independently a lone pair of electrons or a hydrogen, with the proviso that at least one of R₁, R₂, R₃ and R₄ is not a hydrogen. Exemplary hydralazine compounds include budralazine, cadralazine, dihydralazine, endralazine, hydralazine, pilodralazine, todralazine, and the like. Suitable hydralazine compounds are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, Thirteenth Edition; and on STN Express, file phar and file registry.

In some embodiments the hydralazine compound is hydralazine or a pharmaceutically acceptable salt thereof such as hydralazine hydrochloride. In more particular embodiments the hydralazine is administered as hydralazine hydrochloride in an amount of about 10 milligrams to about 300 milligrams as a single dose or as multiple doses per day.

Suitable H₂ receptor antagonists include, but are not limited to, burimamide, cimetidine, ebrotidin, famotidine, nizatidine, roxatidine, rantidine, tiotidine, and the like. Suitable H₂ receptor antagonists are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995, Pgs. 901-915; the Merck Index on CD-ROM, 13th Edition; and in WO 00/28988 assigned to NitroMed Inc., the disclosures of which are incorporated herein by reference in their entirety.

Suitable neutral endopeptidase inhibitors include, but are not limited to, atrial natriuretic peptides, diazapsins, azepinones, ecadotril, fasidotril, fasidotrilat, omapatrilat, sampatrilat, BMS 189,921, Z 13752 A, and the like. Neutral endopeptidase inhibitors are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, Thirteenth Edition; and on STN Express, file phar and file registry.

Suitable NSAIDs include, but are not limited to, acetaminophen, acemetacin, acelofenac, alminoprofen, amfenac, bendazac, benoxaprofen, bromfenac, bucloloyxid acid, butibufen, carprofen, cinmetacin, clopirac, diclofenac, etodolac, felbinac, fencloczic acid, fenbufen, fenoprofen, fentiazac, flunoxaprofen, flurbiprofen, ibufenac, ibuprofen, indomethacin, isofezolac, isoxepac, indoprofen, ketoprofen, lonazolac, loxoprofen, metazinonic acid, mofezolac, miroprofen, naproxen, oxaprozin, pirozolac, piroprofen, pranoprofen, protizinonic acid, salicylamide, sulindac, suprofen, suxibuzone, tiaprofenic acid, tolnetin, xenbucin, ximoprofen, zaltoprofen, zomepirac, aspirin, acemetacin, bumadizon,
carprofenac, clidanac, diflunisal, enfenamic acid, fendosal, flufenamic acid, flunixin, genistic acid, ketorolac, meclofenamic acid, mefenamic acid, mesalamine, prodrugs thereof, and the like. Suitable NSAIDs are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995, Pgs. 617-657; the Merck Index on CD-ROM, 13th Edition; and in U.S. Patent Nos. 6,057,347 and 6,297,260 assigned to NitroMed Inc., the disclosures of which are incorporated herein by reference in their entirety.

In some embodiments the NSAIDs are acetaminophen, diclofenac, flurbiprofen, ibuprofen, indomethacin, ketoprofen, naproxen or aspirin. In more particular embodiments the acetaminophen is administered in an amount of about 325 milligrams to about 4 grams as a single dose or as multiple doses per day; the diclofenac is administered in an amount of about 50 milligrams to about 250 milligrams as a single dose or as multiple doses per day; the flurbiprofen is administered in an amount of about 100 milligrams to about 300 milligrams as a single dose or as multiple doses per day; the ibuprofen is administered in an amount of about 400 milligrams to about 3.2 grams as a single dose or as multiple doses per day; the indomethacin is administered in an amount of about 25 milligrams to about 200 milligrams as a single dose or as multiple doses per day; the ketoprofen is administered in an amount of about 50 milligrams to about 300 milligrams as a single dose or as multiple doses per day; the naproxen is administered in an amount of about 250 milligrams to about 1.5 grams as a single dose or as multiple doses per day; the aspirin is administered in an amount of about 10 milligrams to about 2 grams as a single dose or as multiple doses per day.

Suitable phosphodiesterase inhibitors, include but are not limited to, filaminast, piclamilast, rolipram, Org 20241, MCI-154, roflumilast, toborinone, posicar, lixazinone, zaprinast, sildenafil, pyrazolopyrimidinones, motapizone, pimobendan, zardaverine, siguazodon, CI 930, EMD 53998, imazodon, saterinone, loprinone hydrochloride, 3-pyridinecarbonitrile derivatives, acefylline, albifylline, bamifylline, denbufyllene, diphylline, doxofylline, etofylline, torbafylline, theophylline, nanterinone, pentoxofylline, proxyphylline, cilostazol, cilostamide, MS 857, piroximone, milrinone, amrinone, tolafentrine, diprydamole, papaveroline, E4021, thienopyrimidine derivatives, triflusil, ICOS-351, tetrahydropipеразино(1,2-b)beta-carboline-1,4-dione derivatives, carboline derivatives, 2-pyrazolin-5-one derivatives, fused pyridazine derivatives, quinazoline derivatives, anthranilic acid derivatives, imidazoquinazoline derivatives, tadafil, vardenafil, and in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Ed.), McGraw-Hill, Inc. (1995), The Physician's Desk Reference (49th Ed.), Medical Economics (1995), Drug Facts and
Comparisons (1993 Ed), Facts and Comparisons (1993), and the Merck Index on CD-ROM, 13th Edition; and the like. Phosphodiesterase inhibitors and their nitrosated and/or nitrosylated derivatives are also disclosed in U. S. Patent Nos. 5,932,538, 5,994,294, 5,874,437, 5,958,926 reissued as U.S. Patent No. RE 03772346, 172,060, 6,197,778, 6,177,428, 6,172,068, 6,221,881, 6,232,321, 6,197,782, 6,133,272, 6,211,179, 6,316,457 and 6,331,542, the disclosures of each of which are incorporated herein by reference in their entirety.

Suitable potassium channel blockers include but are not limited to, nicorandil, pinacidil, cromakalim (BRL 34915), aprikalim, bimakalim, emakalim, lemakalim, minoxidil, diazoxide, 9-chloro-7-(2-chlorophenyl)-5H-pyrimido(5,4-d)(2)-benzazepine, Ribi, CPG-11952, CGS-9896, ZD 6169, diazoxide, Bay X 9227, P1075, Bay X 9228, SDZ PCO 400, WAY-120,491, WAY-120,129, Ro 31-6930, SR 44869, BRL 38226, S 0121, SR 46142A, CPG 42500, SR 44994, artilide fumarate, lorazepam, temazepam, rilmazafone, nimetazepam, midazolam, lormetazepam, loprazolam, ibutilide fumarate, haloxazolam, flunitrazepam, estazolam, doxepazine, clonazepam, cinnazolam, brotizolam, and the like. Suitable potassium channel blockers are described more fully in the literature, such as in Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the Merck Index on CD-ROM, Thirteenth Edition; and on STN Express, file phar and file registry.

Suitable platelet reducing agents include but are not limited to, fibrinolytic agents such as for example, ancrod, anistreplase, bisobrin lactate, brinolase, Hageman factor (i.e. factor XII) fragments, plasminogen activators such as, for example, streptokinase, tissue plasminogen activators (TPA), urokinase, pro-Urokinase, recombinant TPA, plasmin, plasminogen, and the like; anti-coagulant agents including but are not limited to, inhibitors of factor Xa, factor TFPI, factor VIIa, factor IXc, factor Va, factor VIIIa, inhibitors of other coagulation factors, and the like; vitamin K antagonists, such as, for example, coumarin, coumarin derivatives (e.g., warfarin sodium); glycosaminoglycans such as, for example, heparins both in unfractionated form and in low molecular weight form; ardeparin sodium, bivalirudin, bromindione, coumarin, dalteparin sodium, danaparoid sodium; dazoxiben hydrochloride, desirudin, dicumarol, efegatran sulfate, enoxaparin sodium, ifetroban, ifetroban sodium, lyapolate sodium, nafamostat mesylate, phenprocoumon, sulfatide, tinzaparin sodium, retaplace; trifenagrel, warfarin, dextrins and the like; abciximab, acadesine, anipamil, argatroban, aspirin, clopidogrel, diadenosine 5',5''-P1,P4-tetraphosphate (Ap4A) analogs, difibrotide, dilazep dihydrochloride, dipyridamole, dopamine, 3-methoxytyramine, glucagon, glycoprotein IIb/IIIa antagonists, such as, for example, Ro-43-
8857, L-700,462, iloprost, isocarbacyclin methyl ester, itazigrel, ketanserin, BM-13.177,
lanifiban, lifarizine, molsidomine, nifedipine, oxagrelate, prostaglandins, platelet activating
factor antagonists such as, for example, lexipafant, prostacyclins, pyrazines, pyridinol
carbamate, ReoPro (i.e., abciximab), sulfipyrazone, synthetic compounds BN-50727, BN-
52021, CV-4151, E-5510, FK-409, GU-7, KB-2796, KBT-3022, KC-404, KF-4939, OP-
41483, TRK-100, TA-3090, TFC-612, ZK-36374, 2,4,5,7-tetrahydrooctane, 2,4,5,7-
tetrahydrooctane 2,2-dioxide, 2,4,5-trithiahexane, theophyllin pentoxifyllin, thromboxane and
thromboxane synthetase inhibitors such as, for example, picotamide, sulotroban, ticlopidine,
tirofiban, trapidil, ticlodipine, trifenagrel, trilinolein, 3-substituted 5,6-bis(4-methoxyphenyl)-
1,2,4-triazines; antibodies to glycoprotein IIb/IIIa; anti-serotonin drugs, such as, for example,
clopidogrel; sulfipyrazone and the like; aspirin; dipyridamole; clofibrate; pyridinol
carbamate; glucagon, caffeine; theophyllin pentoxifyllin; ticlopidine, and the like.

Suitable prostaglandins, include but are not limited to, naturally occurring
prostaglandins such as, for example, arbaprostil, alprostadil, beraprost, carboprost,
cloprostenol, dimoxaprost, enprostil, enisoprost, fluprostrenol, fenprostalene, gemprost,
latanoprost, limaprost, meteneprost, mexiprostil, misoprostol, misoprostol acid, nocloprost,
ornoprostil, prostanlene, PGE_{1}, PGE_{2}, PGF_{1}, PGF_{2α}, rioprostil, rosaprostol,
remiprostol, salprostone, trimoprostil, tiprostanide, travoprost, unoprostone, viprostol,
viprostol. Suitable prostaglandins are described more fully in the literature, such as in
Goodman and Gilman, The Pharmacological Basis of Therapeutics (9th Edition), McGraw-
Hill, 1995; and the Merck Index on CD-ROM, 13th Edition; and on STN Express, file phar
and file registry.

In some embodiments the prostaglandins are cloprostenol, fluprostrenol and
tavoprost.

Suitable proton pump inhibitors include, but are not limited to, disulprazole,
esomeprazole, lansoprazole, leminoprazole, omeprazole, pantoprazole, rabeprazole,
timoprazole, tenatoprazole, 2-(2-benzimidazoly)-pyridine, tricyclic imidazole,
thenopydine benzimidazole, fluoroalkoxy substituted benzimidazole, dialkoxy
benzimidazole, N-substituted 2-(pyridylalkenesulfinyl) benzimidazole, cycloheptenepridine,
5-pyrrolyl-2-pyridylmethydsulfinyl benzimidazole, alkylsulfinyl benzimidazole, fluoro-
pyridylmethylsulfinyl benzimidazole, imidazo(4,5-b)pyrididine, RO 18-5362, IY 81149, 4-
amino-3-carbonyl quinoline, 4-amino-3-aclyphthryride, 4-aminoquinoline, 4-amino-3-
acryquinoline, 3-butyryl-4-(2-methylphenylamino)-8-(2-hydroxyethoxy)quinoline,
quinazoline, tetrahydroisoquinolin-2-yl pyrimidine, YH 1885, 3-substituted 1,2,4-
thiadiazolo(4,5-a) benzimidazole, 3-substituted imidazo(1,2-d)-thiadiazole, 2-
sulfinynicotinamide, pyridylsulfinylbenz imidazole, pyridylsulfinyl thieno imidazole,
theinoimidazole-toluidine, 4,5-dihydrooxazole, thienoimidazole-toluidine, Hoe-731,
imidazo(1,2-a)pyridine, pyrrolo(2,3-b)pyridine, and the like. Suitable proton pump inhibitors
are described more fully in the literature, such as in Goodman and Gilman, The
Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; the Merck Index
on CD-ROM, 13th Edition; and in WO 00/50037 assigned to NitroMed Inc., the disclosures of
which are incorporated herein by reference in their entirety.

Suitable renin inhibitors include, but are not limited to, aldosterone, aliskiren (SPP-
100), ditekiren, enalkrein (A-64662), medullipin, terlikiren, tonin, zankiren, RO 42-5892
(remikiren), A 62198, A 64662, A 65317, A 69729, A 72517 (zankiren), A 74273, CP
80794, CGP 29287, CGP-38560A, EMD 47942, ES 305, ES 1005, ES 8891, FK 906, FK
744, H 113, H-142, KRI 1314, pepstatin A, RO 44-9375 (ciprokinren), RO 42-5892, RO 66-
1132, RO 66-1168, SP 500, SP 800, SR-43845, SQ 34017, U 71038, YM-21095, YM-
26365, urea derivatives of peptides, amino acids connected by nonpeptide bonds, di- and
tri-peptide derivatives (e.g., Act-A, Act-B, Act-C, ACT-D, and the like), amino acids and
derivatives thereof, diol sulfonamides and sulfinyls, modified peptides, peptidyl beta-
aminoacetyl aminodiol carbamates, monoclonal antibodies to renin. Suitable renin inhibitors
are described more fully in U.S. Patent Nos. 5,116,835, 5,114,937, 5,106,835, 5,104,869,
5,095,119, 5,098,924), 5,095,006, 5,089,471, 5,075,451, 5,066,643, 5,063,208, 4,845,079,
5,055,466, 4,980,283, 4,885,292), 4,780,401, 5,071,837, 5,064,965, 5,063,207, 5,036,054,
5,036,053, 5,034,512, and 4,894,437, the disclosures of each of which are incorporated
herein by reference in their entirety; and in the literature, such as in Goodman and Gilman,
The Pharmacological Basis of Therapeutics (9th Edition), McGraw-Hill, 1995; and the
Merck Index on CD-ROM, Thirteenth Edition; and on STN Express, file phar and file
registry.

Suitable COX-2 inhibitors include, but are not limited to, nimesulide, celecoxib
(CELEBREX®), etoricoxib (ARCOXIA®), flusulide, lumiracoxib (PREXIG®, COX-189),
parecoxib (DYNSTAT®), rofecoxib (VIOLIX®), tircaxix (JTE-522), valdecoxib
(BEXTRA®), ABT 963, BMS 347070, CS 502, DuP 697, GW-406381, NS-386, SC-57666,
SC-58125, SC-58635, and the like, and mixtures of two or more thereof. Suitable COX-2
inhibitors are in U.S. Patent Nos. 5,344,991, 5,380,738, 5,393,790, 5,409,944, 5,434,178,
5,436,265, 5,466,823, 5,474,995, 5,510,368, 5,536,752, 5,550,142, 5,552,422, 5,604,253,
5,604,260, 5,639,780, 5,932,598 and 6,633,272, and in WO 94/03387, WO 94/15723, WO

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In some embodiments the COX-2 inhibitors are celecoxib, etoricoxib, lumiracoxib, paracoxib, rofecoxib or valdecoxib. In more particular embodiments the celecoxib is administered in an amount of about 100 milligrams to about 800 milligrams as a single dose or as multiple doses per day; the etoricoxib is administered in an amount of about 50 milligrams to about 200 milligrams as a single dose or as multiple doses per day; the lumiracoxib is administered in an amount of about 40 milligrams to about 1200 milligrams as a single dose or as multiple doses per day; the paracoxib is administered in an amount of about 20 milligrams to about 100 milligrams as a single dose or as multiple doses per day; the rofecoxib is administered in an amount of about 12.5 milligrams to about 50 milligrams as a single dose or as multiple doses per day; the valdecoxib is administered in an amount of about 10 milligrams to about 40 milligrams as a single dose or as multiple doses per day.

Suitable steroids include, but are not limited to, 21-acetoxypregneneolone, alcolometasone, algestone, amcinonide, beclomethasone, betamethasone, budesonide, chlorprednisone, clobetasol, clobentasone, clocortolone, cloprednol, corticosterone, cortisine, corticazol (cortivatol), deflazacort, desonide, desoximetasone, dexamethasone, diflorasone, diflucortolone, difluprednate, enoxolone, fluzacort, flucloronide, flumethasone, flunisolide, flucinolone acetonide, flucinidine, flucortin butyl, fluocortolone, fluorometholone, fluperoxolone acetate, fluprednidene acetate, fluprednisolone, flurandrenolide, fluticasone propionate, fluticasone propionate, formocort, halcinonide, halobetasol propionate, halometasone, haloprednol acetate, hydrocortamate, hydrocortisone and its derivatives (such as phosphate, 21-sodium succinate and the like), hydrocortisone terbutate, isoflupredone, loteprednol etabonate, mazipredone, medrysone, meprednisone, methylprednisolone, mometasone furoate, paremethasone, prednicarbate, prednisolone and its derivatives (such as 21-stearoylglycolate, sodium phosphate and the like), prednisone, prednival, prednylidene and its derivatives (such as 21-diethylaminoacetate and the like), rimexolone, tixocortol, trimcinolone and its derivatives (such as acetonide, benetonide and the like), and the like. Suitable NSAIDs are described more fully in the literature, such as in Goodman and Gilman,

In some embodiments the steroids are dexamethasone, fluorometholone, hydrocortisone, and prednisolone.

The invention provides compositions comprising (i) at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and (ii) at least one compound is selected from the group consisting of an aldosterone antagonist, an angiotensin II antagonist, an angiotensin-converting enzyme (ACE) inhibitor, a β-adrenergic antagonist, a calcium channel blocker, a diuretic, a hydralazine compound and a renin inhibitor in one or more pharmaceutically acceptable carriers. In other embodiments of the invention the aldosterone antagonist is eplerenone or spironolactone; the angiotensin II antagonist is candesartan, candesartan cilexetil, eprosartan mesylate, irbesartan, losartan, medoxomil, telmisartan, trandolapril, trandolaprilat or valsartan; the angiotensin-converting enzyme inhibitor is benazepril hydrochloride, captopril, enalapril maleate, fosinopril sodium, lisinopril, moexipril hydrochloride, quinapril hydrochloride, ramipril; the β-adrenergic antagonist is bisoprolol fumarate, carvedilol, metoprolol tartrate, propranolol hydrochloride or timolol maleate; the calcium channel blockers is amlodipine, diltiazem, isradipine, nicardipine, nifedipine, nimodipine, nisoldipine, nitrendipine, verapamil; the diuretic is amiloride hydrochloride, chlorthalidone, hydrochlorothiazide or triamterene; the hydralazine compound is hydralazine hydrochloride; and the renin inhibitor is aliskiren, ciprokiren, ditekiren, enalkrein, medullipin, remikiren, terlikiren, tonin or zankiren.

The invention provides compositions comprising (i) an organic nitric oxide enhancing salt of an angiotensin II antagonist (ii) a nitric oxide enhancing compound, such as, isosorbide dinitrate and/or isosorbide mononitrate (preferably isosorbide dinitrate), and (i) a hydralazine compound (such as hydralazine hydrochloride). In one embodiment, the hydralazine hydrochloride can be administered in an amount of about 30 milligrams per day to about 400 milligrams per day; the isosorbide dinitrate can be administered in an amount of about 10 milligrams per day to about 200 milligrams per day; or the isosorbide mononitrate can be administered in an amount of about 5 milligrams per day to about 120 milligrams per day. In another embodiment, the hydralazine hydrochloride can be administered in an amount of about 50 milligrams per day to about 300 milligrams per day; the isosorbide dinitrate can be administered in an amount of about 20 milligrams per day to about 160 milligrams per day; or
the isosorbide mononitrate can be administered in an amount of about 15 milligrams per day to about 100 milligrams per day. In yet another embodiment, the hydralazine hydrochloride can be administered in an amount of about 37.5 milligrams to about 75 milligrams one to four times per day; the isosorbide dinitrate can be administered in an amount of about 20 milligrams to about 40 milligrams one to four times per day; or the isosorbide mononitrate can be administered in an amount of about 10 milligrams to about 20 milligrams one to four times per day. In another embodiment of the methods of the invention, the patient can be administered a composition comprising about 225 mg hydralazine hydrochloride and about 120 mg isosorbide dinitrate once per day (i.e., q.d.). In another embodiment of the methods of the invention, the patient can be administered a composition comprising about 112.5 mg hydralazine hydrochloride and about 60 mg isosorbide dinitrate twice per day (i.e., b.i.d.). In another embodiment of the methods of the invention, the patient can be administered a composition comprising about 56.25 mg hydralazine hydrochloride and about 30 mg isosorbide dinitrate twice per day (i.e., b.i.d.). In another embodiment of the methods of the invention, the patient can be administered a composition comprising about 75 mg hydralazine hydrochloride and about 40 mg isosorbide dinitrate three times per day (i.e., t.i.d.). In another embodiment of the methods of the invention, the patient can be administered a composition comprising about 37.5 mg hydralazine hydrochloride and about 20 mg isosorbide dinitrate three times per day (i.e., t.i.d.). The particular amounts of hydralazine and isosorbide dinitrate or isosorbide mononitrate can be administered as a single dose once a day; or in multiple doses several times throughout the day; or as a sustained-release oral formulation, or as an injectable formulation.

The invention provides methods for treating cardiovascular disorders by administering to the patient in need thereof an effective amount of the compounds and/or compositions described herein. For example, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist. In another embodiment, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and at least one nitric oxide enhancing compound. In yet another embodiment, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, at least one therapeutic agent, including but not limited to, such as, for example, aldosterone antagonists, \( \alpha \)-adrenergic receptor agonists, \( \alpha \)-adrenergic receptor antagonists, angiotensin II antagonists, angiotensin-converting enzyme (ACE) inhibitors, antidiabetic compounds, anti-hyperlipidemic compounds, antimicrobial compounds, antioxidants, antithrombotic and
vasodilator compounds, β-adrenergic antagonists, calcium channel blockers, carbonic anhydrase inhibitors, digitalis, diuretics, endothelin antagonists, hydralazine compounds, H₂ receptor antagonists, neutral endopeptidase inhibitors, nonsteroidal antiinflammatory compounds (NSAIDs), phosphodiesterase inhibitors, potassium channel blockers, platelet reducing agents, prostaglandins, proton pump inhibitors, renin inhibitors, selective cyclooxygenase-2 (COX-2) inhibitors, steroids, and combinations of two or more thereof. In another embodiment, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, at least one therapeutic agent, and, at least one nitric oxide enhancing compound. In one embodiment the cardiovascular disorder is is hypertension, heart failure, arterial stiffness, postmyocardial infarction, stroke and/or diastolic dysfunction. The organic nitric oxide enhancing salts of angiotensin II antagonists, nitric oxide enhancing compounds, and/or therapeutic agents can be administered separately or as components of the same composition in one or more pharmaceutically acceptable carriers.

The invention provides methods for treating renovascular diseases by administering to the patient in need thereof an effective amount of the compounds and/or compositions described herein. For example, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonista. In another embodiment, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonista, and at least one nitric oxide enhancing compound. In yet another embodiment, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonista, and, at least one therapeutic agent, including but not limited to, such as, for example, aldosterone antagonists, α-adrenergic receptor agonists, α-adrenergic receptor antagonists, angiotensin II antagonists, angiotensin-converting enzyme (ACE) inhibitors, antidiabetic compounds, antihyperlipidemic compounds, antimicrobial compounds, antioxidants, antithrombotic and vasodilator compounds, β-adrenergic antagonists, calcium channel blockers, carbonic anhydrase inhibitors, digitalis, diuretics, endothelin antagonists, hydralazine compounds, H₂ receptor antagonists, neutral endopeptidase inhibitors, nonsteroidal antiinflammatory compounds (NSAIDs), phosphodiesterase inhibitors, potassium channel blockers, platelet reducing agents, prostaglandins, proton pump inhibitors, renin inhibitors, selective cyclooxygenase-2 (COX-2) inhibitors, steroids, and combinations of two or more thereof. In another embodiment, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonista, and, at least one
therapeutic agent, and, at least one nitric oxide enhancing compound. In one embodiment the renovascular disease is renal failure, renal insufficiency, renal deterioration associated with severe hypertension or renovascular hypertension. The organic nitric oxide enhancing salts of angiotensin II antagonists, nitric oxide enhancing compounds, and/or therapeutic agents can be administered separately or as components of the same composition in one or more pharmaceutically acceptable carriers.

The invention provides methods for treating diabetes; treating diseases resulting from oxidative stress; treating endothelial dysfunctions; treating diseases caused by endothelial dysfunctions; treating cirrhosis; treating pre-eclampsia; treating osteoporosis; treating nephropathy; treating peripheral vascular diseases; treating portal hypertension; treating metabolic syndrome; and treating hyperlipidemia by administering to the patient in need thereof an effective amount of the compounds and/or compositions described herein. For example, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist. In another embodiment, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and at least one nitric oxide enhancing compound. In yet another embodiment, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, at least one therapeutic agent, including but not limited to, such as, for example, aldosterone antagonists, α-adrenergic receptor agonists, α-adrenergic receptor antagonists, angiotensin II antagonists, angiotensin-converting enzyme (ACE) inhibitors, antidiabetic compounds, anti-hyperlipidemic compounds, antimicrobial compounds, antioxidants, antithrombotic and vasodilator compounds, β-adrenergic antagonists, calcium channel blockers, carbonic anhydrase inhibitors, digitalis, diuretics, endothelin antagonists, hydralazine compounds, H₂ receptor antagonists, neutral endopeptidase inhibitors, nonsteroidal antiinflammatory compounds (NSAIDs), phosphodiesterase inhibitors, potassium channel blockers, platelet reducing agents, prostaglandins, proton pump inhibitors, renin inhibitors, selective cyclooxygenase-2 (COX-2) inhibitors, steroids, and combinations of two or more thereof. In another embodiment, the patient can be administered an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, at least one therapeutic agent, and, at least one nitric oxide enhancing compound. The organic nitric oxide enhancing salts of angiotensin II antagonists, nitric oxide enhancing compounds, and/or therapeutic agents can be administered separately or as components of the same composition in one or more
pharmaceutically acceptable carriers.

The invention provides methods for treating ophthalmic disorders in a patient in need thereof comprising administering to the patient an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, optionally, at least one therapeutic agent, such as, for example, α-adrenergic receptor agonists, angiotensin-converting enzyme (ACE) inhibitors, antimicrobial compounds, β-adrenergic antagonists, carbonic anhydrase inhibitors, nonsteroidal antiinflammatory compounds, prostaglandins, selective cyclooxygenase-2 (COX-2) inhibitors, steroids and combinations of two or more thereof. The methods can optionally further comprise the administration of at least one nitric oxide enhancing compound. In this embodiment of the invention, the methods can involve (i) administering the organic nitric oxide enhancing salt of an angiotensin II antagonist, (ii) administering the organic nitric oxide enhancing salt of an angiotensin II antagonist, (iii) administering the organic nitric oxide enhancing salt of an angiotensin II antagonist and therapeutic agents, or (iv) administering organic nitric oxide enhancing salt of an angiotensin II antagonist, nitric oxide enhancing compounds, and therapeutic agents. In one embodiment the at least one therapeutic agent is selected from the group consisting of an α-adrenergic receptor agonist, an angiotensin-converting enzyme (ACE) inhibitor, an antimicrobial compound, a β-adrenergic antagonist, a carbonic anhydrase inhibitor, a nonsteroidal antiinflammatory compound, a prostaglandin, a selective cyclooxygenase-2 (COX-2) inhibitor, and a steroid. In one embodiment the ophthalmic disorder is ophthalmic infection, glaucoma, elevated ocular pressure, macular degeneration and diabetic retinopathy. The organic nitric oxide enhancing salts of angiotensin II antagonists, nitric oxide enhancing compounds, and/or therapeutic agents can be administered separately or as components of the same composition in one or more pharmaceutically acceptable carriers.

When administered separately, the organic nitric oxide enhancing salt of the angiotensin II antagonist, nitric oxide enhancing compound and/or therapeutic agent can be administered about the same time as part of the overall treatment regimen, i.e., as a combination therapy. "About the same time" includes administering the organic nitric oxide enhancing salt of the angiotensin II antagonist, simultaneously, sequentially, at the same time, at different times on the same day, or on different days, as long as they are administered as part of an overall treatment regimen, i.e., combination therapy or a therapeutic cocktail.

When administered in vivo, the compounds and compositions of the invention can be administered in combination with pharmaceutically acceptable carriers and in dosages
described herein. When the compounds and compositions of the invention are administered as a combination of at least organic nitric oxide enhancing salt of an angiotensin II antagonist and/or at least one nitric oxide enhancing compound and/or therapeutic agent, they can also be used in combination with one or more additional compounds which are known to be effective against the specific disease state targeted for treatment. The nitric oxide enhancing compounds, therapeutic agents and/or other additional compounds can be administered simultaneously with, subsequently to, or prior to administration of the organic nitric oxide enhancing salt of the angiotensin II antagonist.

The compounds and compositions of the invention can be administered by any available and effective delivery system including, but not limited to, orally, bucally, parenterally, by inhalation, by topical application, by injection, transdermally, or rectally (e.g., by the use of suppositories) in dosage unit formulations containing conventional nontoxic pharmaceutically acceptable carriers, adjuvants, and vehicles, as desired. Parenteral includes subcutaneous injections, intravenous, intramuscular, intrasternal injection, or infusion techniques. In one embodiment of the invention the organic nitric oxide enhancing salt of an angiotensin II antagonist is administered orally, parentally or by inhalation.

Transdermal compound administration, which is known to one skilled in the art, involves the delivery of pharmaceutical compounds via percutaneous passage of the compound into the systemic circulation of the patient. Topical administration can also involve the use of transdermal administration such as transdermal patches or iontophoresis devices. Other components can be incorporated into the transdermal patches as well. For example, compositions and/or transdermal patches can be formulated with one or more preservatives or bacteriostatic agents including, but not limited to, methyl hydroxybenzoate, propyl hydroxybenzoate, chlorocresol, benzalkonium chloride, and the like. Dosage forms for topical administration of the compounds and compositions can include creams, sprays, lotions, gels, ointments, eye drops, nose drops, ear drops, and the like. In such dosage forms, the compositions of the invention can be mixed to form white, smooth, homogeneous, opaque cream or lotion with, for example, benzyl alcohol 1% or 2% (wt/wt) as a preservative, emulsifying wax, glycerin, isopropyl palmitate, lactic acid, purified water and sorbitol solution. In addition, the compositions can contain polyethylene glycol 400. They can be mixed to form ointments with, for example, benzyl alcohol 2% (wt/wt) as preservative, white petrolatum, emulsifying wax, and tenox II (butylated hydroxyanisole, propyl gallate, citric acid, propylene glycol). Woven pads or rolls of bandaging material, e.g., gauze, can be impregnated with the compositions in solution, lotion, cream, ointment or other such form
can also be used for topical application. The compositions can also be applied topically using a transdermal system, such as one of an acrylic-based polymer adhesive with a resinous crosslinking agent impregnated with the composition and laminated to an impermeable backing.

The compositions can also be applied topically using a transdermal system, such as one of an acrylic-based polymer adhesive with a resinous crosslinking agent impregnated with the composition and laminated to an impermeable backing. In a particular embodiment, the compositions of the invention are administered as a transdermal patch, more particularly as a sustained-release transdermal patch. The transdermal patches of the invention can include any conventional form such as, for example, adhesive matrix, polymeric matrix, reservoir patch, matrix or monolithic-type laminated structure, and are generally comprised of one or more backing layers, adhesives, penetration enhancers, an optional rate controlling membrane and a release liner which is removed to expose the adhesives prior to application. Polymeric matrix patches also comprise a polymeric-matrix forming material. Suitable transdermal patches are described in more detail in, for example, U. S. Patent Nos. 5,262,165, 5,948,433, 6,010,715 and 6,071,531, the disclosure of each of which are incorporated herein in their entirety.

Solid dosage forms for oral administration can include capsules, sustained-release capsules, tablets, sustained release tablets, chewable tablets, sublingual tablets, effervescent tablets, pills, powders, granules and gels. In such solid dosage forms, the active compounds can be admixed with at least one inert diluent such as sucrose, lactose or starch. Such dosage forms can also comprise, as in normal practice, additional substances other than inert diluents, e.g., lubricating agents such as magnesium stearate. In the case of capsules, tablets, effervescent tablets, and pills, the dosage forms can also comprise buffering agents. Soft gelatin capsules can be prepared to contain a mixture of the active compounds or compositions of the invention and vegetable oil. Hard gelatin capsules can contain granules of the active compound in combination with a solid, pulverulent carrier such as lactose, saccharose, sorbitol, mannitol, potato starch, corn starch, amylopectin, cellulose derivatives of gelatin. Tablets and pills can be prepared with enteric coatings.

Liquid dosage forms for oral administration can include pharmaceutically acceptable emulsions, solutions, suspensions, syrups, and elixirs containing inert diluents commonly used in the art, such as water. Such compositions can also comprise adjuvants, such as wetting agents, emulsifying and suspending agents, and sweetening, flavoring, and perfuming agents.
Suppositories for vaginal or rectal administration of the compounds and compositions of the invention, such as for treating pediatric fever and the like, can be prepared by mixing the compounds or compositions with a suitable nonirritating excipient such as cocoa butter and polyethylene glycols which are solid at room temperature but liquid at rectal temperature, such that they will melt in the rectum and release the drug.

Injectable preparations, for example, sterile injectable aqueous or oleaginous suspensions can be formulated according to the known art using suitable dispersing agents, wetting agents and/or suspending agents. The sterile injectable preparation can also be a sterile injectable solution or suspension in a nontoxic parenterally acceptable diluent or solvent, for example, as a solution in 1,3-butanediol. Among the acceptable vehicles and solvents that can be used are water, Ringer's solution, and isotonic sodium chloride solution. Sterile fixed oils are also conventionally used as a solvent or suspending medium.

The compositions of this invention can further include conventional excipients, i.e., pharmaceutically acceptable organic or inorganic carrier substances suitable for parenteral application which do not deleteriously react with the active compounds. Suitable pharmaceutically acceptable carriers include, for example, water, salt solutions, alcohol, vegetable oils, polyethylene glycols, gelatin, lactose, amylose, magnesium stearate, talc, surfactants, silicic acid, viscous paraffin, perfume oil, fatty acid monoglycerides and diglycerides, petroethral fatty acid esters, hydroxymethyl-cellulose, polyvinylpyrrolidone, and the like. The pharmaceutical preparations can be sterilized and if desired, mixed with auxiliary agents, e.g., lubricants, preservatives, stabilizers, wetting agents, emulsifiers, salts for influencing osmotic pressure, buffers, colorings, flavoring and/or aromatic substances and the like which do not deleteriously react with the active compounds. For parenteral application, particularly suitable vehicles consist of solutions, preferably oily or aqueous solutions, as well as suspensions, emulsions, or implants. Aqueous suspensions may contain substances which increase the viscosity of the suspension and include, for example, sodium carboxymethyl cellulose, sorbitol and/or dextran. Optionally, the suspension may also contain stabilizers.

The composition, if desired, can also contain minor amounts of wetting agents, emulsifying agents and/or pH buffering agents. The composition can be a liquid solution, suspension, emulsion, tablet, pill, capsule, sustained release formulation, or powder. The composition can be formulated as a suppository, with traditional binders and carriers such as triglycerides. Oral formulations can include standard carriers such as pharmaceutical grades of mannitol, lactose, starch, magnesium stearate, sodium saccharine, cellulose, magnesium
carbonate, and the like.

The compounds of the invention, can be incorporated into various types of pharmaceutical compositions, such as, for example, ophthalmic formulations for delivery to the eye (e.g., topically, intracamerally, or via an implant). The compounds are preferably incorporated into topical ophthalmic formulations, such as for example, solutions, suspensions, gels, ointments, implants, and the like. The compounds of the invention may be combined with ophthalmologically acceptable preservatives, viscosity enhancers, penetration enhancers, buffers, sodium chloride, water to form an aqueous, sterile ophthalmic suspensions or solutions, and the like.

Suitable preservatives include, but are not limited to, benzalkonium chloride, thimerosal, chlorobutanol, methyl paraben, propyl paraben, phenylethyl alcohol, edetate disodium, sorbic acid, ONAMER®, and the like. The preservatives are typically employed at a concentration between about 0.001% and about 1.0% by weight. Appropriate co-solvents include, but are not limited to, Polysorbate 20, 60 and 80; Pluronic F-68, F-84 and P-103; Tyloxapol®; Cremophor® EL; sodium dodecyl sulfate; glycerol; PEG 400; propylene glycol; cyclodextrins, and the like. The co-solvents are typically employed at a concentration between about 0.01% and about 2% by weight. Viscosity enhancers are required as a viscosity greater than that of simple aqueous solutions may be desirable to increase ocular absorption of the active compound, to decrease variability in dispensing the formulations, to decrease physical separation of components of a suspension or emulsion of formulation and/or otherwise to improve the ophthalmic formulation. Suitable viscosity enhancers, include, but are not limited to, polyvinyl alcohol, methyl cellulose, hydroxy propyl carboxymethyl cellulose, hydroxyethylcellulose, hydroxyethylcellulose, hydroxypropylmethylcellulose, methylcellulose, polyvinylpyrrolidone, and the like. Gelling agents can also be used, including, but not limited to, gellan and xanthan gum, and the like. Viscosity enhancers are typically employed at a concentration between about 0.01% and about 2% by weight.

Ophthalmic solution formulations may be prepared by dissolving a compound in a physiologically acceptable isotonic aqueous buffer. Alternatively, the ophthalmic solution may include an ophthalmologically acceptable surfactant to assist in dissolving the compound. Additionally for sterile ophthalmic ointment formulations, the compounds of the invention may be combined with a preservative in an appropriate vehicle, such as, mineral oil, liquid lanolin, or white petrolatum. Sterile ophthalmic gel formulations may be prepared by suspending the active ingredient in a hydrophilic base prepared from the combination of, for example, carbopol-974, and the like.
Various delivery systems are known and can be used to administer the compounds or compositions of the invention, including, for example, encapsulation in liposomes, microbubbles, emulsions, microparticles, microcapsules and the like. The required dosage can be administered as a single unit or in a sustained release form.

The bioavailability of the compositions can be enhanced by micronization of the formulations using conventional techniques such as grinding, milling, spray drying and the like in the presence of suitable excipients or agents such as phospholipids or surfactants.

Sustained release dosage forms of the invention may comprise microparticles and/or nanoparticles having a therapeutic agent dispersed therein or may comprise the therapeutic agent in pure, crystalline, solid form. The therapeutic dosage forms of this aspect of the invention may be of any configuration suitable for sustained release.

Nanoparticle sustained release therapeutic dosage forms are preferably biodegradable and, optionally, bind to the vascular smooth muscle cells and enter those cells, primarily by endocytosis. The biodegradation of the nanoparticles occurs over time (e.g., 30 to 120 days; or 10 to 21 days) in prelyosomal vesicles and lysosomes. Larger microparticle therapeutic dosage forms of the invention release the therapeutic agents for subsequent target cell uptake with only a few of the smaller microparticles entering the cell by phagocytosis. A practitioner in the art will appreciate that the precise mechanism by which a target cell assimilates and metabolizes a dosage form of the invention depends on the morphology, physiology and metabolic processes of those cells. The size of the particle sustained release therapeutic dosage forms is also important with respect to the mode of cellular assimilation. For example, the smaller nanoparticles can flow with the interstitial fluid between cells and penetrate the infused tissue. The larger microparticles tend to be more easily trapped interstitially in the infused primary tissue, and thus are useful to deliver anti-proliferative therapeutic agents.

Particular sustained release dosage forms of the invention comprise biodegradable microparticles or nanoparticles. More particularly, biodegradable microparticles or nanoparticles are formed of a polymer containing matrix that biodegrades by random, nonenzymatic, hydrolytic scissioning to release therapeutic agent, thereby forming pores within the particulate structure.

In a particular embodiment, the compositions of the invention are orally administered as a sustained release tablet or a sustained release capsule. For example, the sustained release formulations can comprise an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and, optionally at least one nitric oxide enhancing
compound, or the sustained release formulations can comprise an effective amount of at least one organic nitric oxide enhancing salt of an angiotensin II antagonist, and at least one nitric oxide enhancing compound, and, optionally at least one therapeutic agent.

While individual needs may vary, determination of optimal ranges for effective amounts of the compounds and/or compositions is within the skill of the art. Generally, the dosage required to provide an effective amount of the compounds and compositions, which can be adjusted by one of ordinary skill in the art, will vary depending on the age, health, physical condition, sex, diet, weight, extent of the dysfunction of the recipient, frequency of treatment and the nature and scope of the dysfunction or disease, medical condition of the patient, the route of administration, pharmacological considerations such as the activity, efficacy, pharmacokinetic and toxicology profiles of the particular compound used, whether a drug delivery system is used, and whether the compound is administered as part of a drug combination.

The amount of a given organic nitric oxide enhancing salt of an angiotensin II antagonist that will be effective in the treatment of a particular disorder or condition will depend on the nature of the disorder or condition, and can be determined by standard clinical techniques, including reference to Goodman and Gilman, supra; The Physician's Desk Reference, Medical Economics Company, Inc., Oradell, N.J., 1995; and Drug Facts and Comparisons, Inc., St. Louis, MO, 1993. The precise dose to be used in the formulation will also depend on the route of administration, and the seriousness of the disease or disorder, and should be decided by the physician and the patient's circumstances.

The invention also provides pharmaceutical kits comprising one or more containers filled with one or more of the ingredients of the pharmaceutical compounds and/or compositions of the invention, including, at least, one or more of the novel organic nitric oxide enhancing salts of angiotensin II antagonists and one or more of the nitric oxide enhancing compounds described herein. Associated with such kits can be additional therapeutic agents or compositions (e.g., aldosterone antagonists, α-adrenergic receptor agonists, α-adrenergic receptor antagonists, angiotensin II antagonists, angiotensin-converting enzyme (ACE) inhibitors, antidiabetic compounds, anti-hyperlipidemic compounds, antimicrobial compounds, antioxidants, antithrombotic and vasodilator compounds, β-adrenergic antagonists, calcium channel blockers, carbonic anhydrase inhibitors, digitalis, diuretics, endothelin antagonists, hydralazine compounds, H₂ receptor antagonists, neutral endopeptidase inhibitors, nonsteroidal anti-inflammatory compounds (NSAIDs), phosphodiesterase inhibitors, potassium channel blockers, platelet reducing agents,
prostaglandins, proton pump inhibitors, renin inhibitors, selective cyclooxygenase-2 (COX-2) inhibitors, steroids, and the like, and combinations of two or more thereof), devices for administering the compositions, and notices in the form prescribed by a governmental agency regulating the manufacture, use or sale of pharmaceuticals or biological products which reflects approval by the agency of manufacture, use or sale for humans.

**EXAMPLES**

**Example 1:** L-Valine, N-(1-oxopentyl)-N-[[2’-(1H-tetrazol-5-yl)[1,1’-biphenyl]-4-yl][methyl]-; salt with 3-pyridinecarboxamide, N-[3-(nitrooxy)propyl]- (1:2)

1a. 3-Pyridinecarboxamide, N-[3-(nitrooxy)propyl]-

![Chemical structure](image)

To a mixture of sodium bicarbonate (5.0 g, 59.5 mmol), 1-propanol, 3-amino-, nitrate (ester), nitrate (1:1) (salt) (1.83 g, 10.0 mmol, prepared according to WO 2005/030135, Example 8a) in water (15 mL) and chloroform (20 mL) was added nicotinoyl chloride hydrochloride (2.5 g, 14.04 mmol) over a period of 10 minutes at 0 °C. After addition, the reaction was warmed to room temperature and stirred at room temperature for 16 hours. The reaction mixture was extracted with water. The organic layer was evaporated and the resulting mixture was purified by column chromatography (silica gel, eluting with 95:5 dichloromethane:methanol) to give the title compound (1.2 g, 53% yield) as a white solid: mp 56 -58 °C; ¹H NMR (400 Mz, d₆-DMSO) δ 8.96 (m, 1H), 8.72 (m, 1H), 8.67 (dd, J = 1.6, 4.8 Hz, 1H), 8.13 (m, 1H), 7.47 (dd, J = 4.8, 8 Hz, 1H), 4.56 (t, J = 6.3 Hz, 2H), 3.34 (q, J = 6.3 Hz, 2H), 1.91 (m, 2H); Mass spectrum (API-TIS) m/z 226 (MH⁺), 248 (MNa⁺).

1b. L-Valine, N-(1-oxopentyl)-N-[[2’-(1H-tetrazol-5-yl)[1,1’-biphenyl]-4-yl][methyl]-, compd. with N-[3-(nitrooxy)propyl]-3-pyridinecarboxamide (1:2)

![Chemical structure](image)
A solution of valsartan (0.46 g, 1.06 mmol, Onbio, Inc.) in ethyl acetate (10 mL) was added to a solution of the product of Example 1a (0.60 g, 2.67 mmol) in ethyl acetate (10 mL). The reaction mixture was stirred at room temperature for 10 minutes and the solvent was evaporated. The resulting oil was washed with ethyl ether and then 5% methanol in ethyl ether, dried to yield the title compound (0.2 g, 21% yield) as a white foam: mp 60-63 °C; 1H NMR (400 MHz, d6-DMSO) δ 8.96 (m, 1H), 8.72-8.67 (m, 3H), 8.13 (m, 2H), 7.65-7.46 (m, 6H), 7.18-6.93 (m, 4H), 4.70-4.55 (m, 5H), 4.44-4.04 (m, 2H), 3.34 (q, J = 6.5 Hz, 4H), 2.40-2.00 (m, 3H), 1.89 (m, 4H), 1.52-1.08 (m, 4H), 0.90-0.66 (m, 9H); Mass spectrum (API-TIS) m/z 436 (MH+), 458 (MNa+).

Example 2: L-Valine, N-(1-oxopentyl)-N-[[2'-((1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl]methyl]-; salt with 1-propanol, 3-amino-, nitrate (ester) (1:2)

2a. 1-Propanol, 3-amino-, nitrate (ester)

To a solution of 1-propanol, 3-amino-, nitrate (ester), nitrate (1:1) (salt) (1.50 g, 8.19 mmol, prepared according to WO 2005/030135, Example 8a) in water (15 mL) and dichloromethane (20 mL) at 0 °C was added sodium hydroxide solid (0.328 g, 8.19 mmol). The reaction was stirred at room temperature for 10 minutes and dichloromethane layer was separated. The aqueous layer was extracted with dichloromethane. The combined organic layer was evaporated to give the title compound as an oil which was used directly to the next step without further purification.

2b. L-Valine, N-(1-oxopentyl)-N-[[2'-((1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl]methyl]-, compd. with 1-propanol, 3-amino-, nitrate (ester) (1:2)

To a stirred solution of valsartan (0.80 g, 1.84 mmol, Onbio, Inc.) in 10% methanol in ethyl ether (15 mL) was added a solution of the product of Example 2a (0.62 g, 4.63 mmol) in 10% methanol in ethyl ether (15 mL). The reaction mixture was stirred at room temperature for 10 minutes and resulting oil was separated. The oil was washed with 10% methanol in ethyl ether and was then dissolved in dichloromethane (1 mL). The dichloromethane solution
was diluted with ethyl ether and hexane. The resulting semisolid was washed with ethyl acetate, dried on a vacuum pump to give title compound (0.52 g, 42% yield) as a white solid: mp 108 -111 °C; 1H NMR (400 Mz, d6-DMSO) δ 8.30 (br s, 4H), 7.59 (t, J = 8.2 Hz, 1H), 7.37-7.30 (m, 3H), 7.04-6.88 (m, 4H), 4.68-3.76 (m, 7H), 2.79 (t, J = 7.3 Hz, 4H), 2.55-2.09 (m, 3H), 1.93 (m, 4H), 1.58-1.11 (m, 4H), 0.90-0.71 (m, 9H); Mass spectrum (API-TIS) m/z 436 (MH+)

Example 3: L-Valine, N-(1-oxopentyl)-N-[[2'(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl]methyl]-; salt with 1,3-propanediol, 2-amino-2-[(nitrooxy)methyl]-, dinitrate (ester)

![Chemical Structure Image]

To a stirred solution of 1,3-propanediol, 2-amino-2-[(nitrooxy)methyl]-, dinitrate (ester) (0.44 g, 1.72 mmol, prepared according to US 2004/0024057; WO 2004/004648, Example 8) in 5% methanol in ethyl ether (10 mL) was added a solution of valsartan (0.35 g, 0.80 mmol, Onbio, Inc.) in 5% methanol in ethyl ether (10 mL). The reaction mixture was stirred at room temperature for 30 minutes and solvent was then removed. The resulting oil was washed with 5 % methanol in ethyl ether to give the title compound as a white foam (0.23 g, 41% yield): mp 78 -82 °C; 1H NMR (400 Mz, d6-DMSO) δ 7.65-7.50 (m, 4H), 7.17-6.93 (m, 4H), 4.62 (s, 6H), 4.59-4.03 (m, 3H), 2.21-1.99 (m, 3H), 1.53-1.06 (m, 4H), 0.90-0.66 (m, 9H); Mass spectrum (API-TIS) m/z 436 (MH+), 458 (MNa+)

Example 4: L-Valine, N-(1-oxopentyl)-N-[[2'(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl]methyl]-; salt with 2-propanediol, 3-amino-, dinitrate (ester), (2R)-4a. 1,2-Propanediol, 3-amino-, dinitrate (ester), (2R)-

![Chemical Structure Image]

To a solution of 1,2-propanediol, 3-amino-, dinitrate (ester), (2R)-, nitrate (1:1) (salt) (0.50 g, 2.05 mmol, prepared according to US 2005/0059655; WO 2005/023183, Example 12a) in water (5 mL) and dichloromethane (15 mL) at 0 °C was added sodium hydroxide solid
The reaction was stirred at room temperature for 10 minutes and dichloromethane layer was separated. The aqueous layer was extracted with dichloromethane. The combined organic layers were evaporated and dried to give the title compound as a light yellow oil which was used directly to the next step without further purification. $^1$H NMR (400 MHz, d$_6$-DMSO) $\delta$ 5.28 (m, 1H), 4.90 (dd, $J = 2.6$, 12.6 Hz, 1H), 4.71 (m, 1H), 2.83 (m, 2H), 2.25 (br s, 2H).

4b. L-Valine, N-(1-oxopentyl)-N-[[2'-[(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl]methyl]-; salt with 2-propanediol, 3-amino-, dinitrate (ester), (2R)-

To a stirred solution of the product of Example 4a (0.33 g, 1.82 mmol) in 5% methanol in ethyl ether (10 mL) was added a solution of valsartan (0.37 g, 0.85 mmol, Onbio, Inc.) in 5% methanol in ethyl ether (10 mL). The reaction mixture was stirred at room temperature for 30 minutes and solvent was removed. The resulting solid was washed with 5% methanol in ethyl ether and dried to give the title compound (0.36 g, 69% yield) as a light yellow solid: mp 102-105 °C; $^1$H NMR (400 MHz, d$_6$-DMSO) $\delta$ 7.55-7.34 (m, 4H), 7.09-6.92 (m, 4H), 5.50 (m, 1H), 4.92 (dd, $J = 3$, 12.7 Hz, 1H), 4.74 (m, 1H), 4.58-3.94 (m, 3H), 3.13-2.98 (m, 2H), 2.41-1.98 (m, 3H), 1.53-1.04 (m, 4H), 0.89-0.68 (m, 9H); Mass spectrum (APITIS) m/z 436 (MH$^+$).

The disclosure of each patent, patent application and publication cited or described in the present specification is hereby incorporated by reference herein in its entirety.

Although the invention has been set forth in detail, one skilled in the art will appreciate that numerous changes and modifications can be made to the invention, and that such changes and modifications can be made without departing from the spirit and scope of the invention.
What is claimed is:

1. A compound of Formula (I), (II), (III), (IV), (V), (VI), (VII), (VIII):

wherein the compound of Formula (I) is:

![Chemical Structure](image)

wherein:

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<th>X₃ is:</th>
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<tr>
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| (7) ![Structure Image](image) | }
(8) -N(D₁)-C(O)-N(D₁)-CH₂-CH₂-CH₃;
(9) -R₁₃;
(10) -C(O)-CH₂-NH(D₄);
(11) -S(O)₂-N(H⁺ Z)-C(O)-C₆H₅;
(12) -S(O)₂-N(H⁺ Z)-C(O)-ND₄⁻ CH₂-CH₂-CH₃; or
(13) -S(O)₂-N(D₄)-OD₁;
Z₃ is a carbon, -CH or a nitrogen atom;
R₁₀ is a fluorine or a hydrogen atom;
Y₃ is:
$Z_4$ is C-$R_{29}$ or a nitrogen;
$R_{37}$ is $-\text{(O)U}_{3}D_1$ or $-\text{COOH}\cdot Z$;
$R_{11}$ is:

1. $-\text{CH}_{2}\cdot \text{OD}_1$;
2. $-\text{R}_37$;

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SUBSTITUTE SHEET (RULE 26)
(3) -C(O)-O-CH(CH₃)-O-C(O)-OR₁₅; or

(4) -CH₂-N(D₁)-C(O)-OR₁₃;

R₁₂ is a chlorine, -SCH₃ or a haloalkyl;
R₁₃ is a lower alkyl or K;
R₁₄ is a lower alkyl or a cycloalkyl;
R₁₅ is:

(1) hydrogen;
(2) a lower alkyl;
(3)

(4)

(5) -R₃₇;

R₁₆ is a hydrogen, a lower alkyl, an alkoxy, -OD₁, a cyano, -R₃₇, NH(D₁) or an alkylcarbonyl;
R₁₇ is an aryl or a cycloalkyl;
R₁₈ at each occurrence is independently selected from a lower alkyl, an alkoxyalkyl, an alkylcarboxylic acid, an alkyl-R₃₇, an hydroxyalkyl, an arylalkoxy, an arylalkyl or an aryl;
R₁₉ is a hydrogen or -R₃₇;
R₂₀ is a hydrogen, a lower alkyl or -R₃₇;
R₂₁ is:

(1)

(2) D₁O

R₂₂ is a hydrogen, -R₃₇ or

SUBSTITUTE SHEET (RULE 26)
$R_{37}$ is a lower alkyl or an alkoxyalkyl;
$R_{32}$ is a hydrogen, an alkyl or an aryl;
$R_{33}$ is $-(\text{CH}_2)_2$-OD$_1$ or

$R_{34}$ is a hydrogen, a lower alkyl, a lower haloalkyl, an aryl or an arylalkyl;
$R_{27}$ is a lower alkyl, an aryl an arylalkyl or $-(\text{CH}_2)_k$-$R_{37}$;
$R_{28}$ is $-\text{OD}_1$, $-\text{S(O)}_2$-$\text{N(D)}_1$H, -$\text{N(D)}_1$H, -$R_{37}$ or CH$_2$-OD$_1$;
$R_{29}$ is a hydrogen, a lower alkyl or -$R_{37}$;
$R_{30}$ is a lower alkyl or a haloalkyl;

$R_{31}$ is:

(1)

\[
\begin{array}{c}
\text{or} \\
\end{array}
\]

(2)

$R_{45}$ is a hydrogen or a lower alkyl;
$R_{46}$ is an alkoxy, an amino group or $-\text{N}(R_{13})$(R$_{13}$);
$R_{42}$ is a lower alkyl or $-(\text{C(R}_o\text{R}_p)_{k-V}}$;

$R_{43}$ and $R_{44}$ taken together are:

(1)

(2)

$o_1$ is an integer from 0 to 3;
k is an integer from 1 to 3;
\[ D_1 \text{ is a hydrogen, } -\text{H} \div \text{Z or K;} \]
\[ D_2 \text{ is a hydrogen or K;} \]
\[ Z \text{ is an organic base or } -\text{N}(R_{38})(R_{39})(R_{40}); \]
\[ R_{38}, R_{39} \text{ and } R_{40} \text{ are each independently selected from } K \text{ or } R_e, \text{ or } R_{38} \text{ and } R_{39} \text{ taken together with the nitrogen to which they are attached are a heterocyclic ring, with the proviso that when the heterocyclic ring is an aromatic ring it can be substituted at any position by } L \text{ and } R_{39} \text{ is not present; } \]
\[ L = -(W_3)_a-E_6-(C(R_e)(R_i))_{p1}-E_4-(C(R_e)(R_i))_{x4}-(W_3)_a-(C(R_e)(R_i))_{y4}-(W_3)_a-E_4-(W_3)_a-(C(R_e)(R_i))_{z4} \]
\[ K = -(W_3)_a-E_6-(C(R_e)(R_i))_{p1}-E_4-(C(R_e)(R_i))_{x4}-(W_3)_a-(C(R_e)(R_i))_{y4}-(W_3)_a-E_4-(W_3)_a-(C(R_e)(R_i))_{z4} \]
\[ a, b, c, d, g, i \text{ and } j \text{ are each independently an integer from 0 to 3;} \]
\[ p, x, y \text{ and } z \text{ are each independently an integer from 0 to 10;} \]
\[ V_4 \text{ is } V_3, R_e, -U_3-V_5 \text{ or } V_6; \]
\[ V_3 \text{ is:} \]

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R₂₄ is -C₆H₄R₂₇, -CN, -S(O)₂-C₆H₄R₂₇, -C(O)-N(R₆)(R₇), -NO₂ or -C(O)-OR₂₅;
R₂₅ is an aryl group, a lower alkyl group, a haloalkyl group, a hydroxyalkyl group or an arylalkyl group;
R₂₆ is -C(O)- or -S(O)₂-;
R₂⁷ is a hydrogen, -CN, -S(O)₂-R₂₅, -C(O)-N(R₆)(R₇), -NO₂ or -C(O)-OR₂₅;
T' is oxygen, sulfur or NR₆;
R₆ is a hydrogen, a lower alkyl group, an aryl group;

V₆ is:

Z₅ is -CH₃ or oxygen;
$Z_6$ is $-\text{CH}$ or nitrogen;

$W_3$ at each occurrence is independently $-\text{C(O)}$, $-\text{C(S)}$, $-\text{T}_3$, $-(\text{C}(\text{R}_a)(\text{R}_b))_{\text{h}}$, $-\text{N}(\text{R}_a)$, an alkyl group, an aryl group, a heterocyclic ring, an arylheterocyclic ring, $-(\text{CH}_2\text{CH}_2\text{O})_{\text{q}}$ or a heterocyclic nitric oxide donor;

$E$ at each occurrence is independently $-\text{T}_3$, an alkyl group, an aryl group, $-(\text{C}(\text{R}_a)(\text{R}_b))_{\text{h}}$, a heterocyclic ring, an arylheterocyclic ring, $-(\text{CH}_2\text{CH}_2\text{O})_{\text{q}}$ or $Y_3$;

$Y_3$ is:

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$T$ is a $-\text{S(O)}_o$; a carbonyl or a covalent bond;

$o$ is an integer from 0 to 2;

$R_j$ and $R_k$ are independently selected from an alkyl group, an aryl group, or $R_j$ and $R_k$ taken together with the nitrogen atom to which they are attached are a heterocyclic ring;

$T_3$ at each occurrence is independently a covalent bond, a carbonyl, an oxygen, $-\text{SO(O)}_o$ or $-\text{N}(\text{R}_a)$;

$h$ is an integer form 1 to 10;
q₁ is an integer from 1 to 5;

Rₑ and Rₜ are each independently a hydrogen, an alkyl, a cycloalkoxy, a halogen, a hydroxy, an hydroxyalkyl, an alkoxyalkyl, an arylheterocyclic ring, an alkylaryl, an alkylcycloalkyl, an alkylheterocyclic ring, a cycloalkylalkyl, a cycloalkylthio, an arylalkylthioalkyl, an alkylthioalkyl, a cycloalkenyl, an heterocyclicalkyl, an alkoxy, a haloalkoxy, an amino, an alkylamino, a dialkylamino, an arylamino, a diarylamino, an alkylarylamino, an alkoxyhaloalkyl, a sulfonic acid, a sulfonic ester, an alkylsulfonic acid, an arylsulfonic acid, an arylalkoxy, an alkythio, an arylthio, an aminoalkyl, an aminoaryl, an aryl, an arylalkyl, an alkylaryl, a carboxamido, an alkylcarboxamido, an arylcarboxamido, an amidyl, a carboxyl, a carbamoyl, an alkylcarboxylic acid, an arylcarboxylic acid, an alkylcarbonyl, an arylcarbonyl, an ester, a carboxylic ester, an alkylcarboxylic ester, an arylcarboxylic ester, a sulfonamido, an alkylsulfonamido, an arylsulfonamido, an alkylsulfon, an alkylsulfonyloxy, an arylsulfon, arylsulphonyloxy, a sulfonic ester, an alkyl ester, an aryl ester, a urea, a phosphoryl, a nitro, -U₃-V₅, V₆, -(C(R₀)(R₉))ₖ₁-U₃-V₅, -(C(R₀)(R₉))ₖ₁-U₃-V₅, -(C(R₀)(R₉))ₖ₁-U₃-V₅, -(C(R₀)(R₉))ₖ₁-U₃-C(O)-V₆, or Rₑ and Rₜ taken together with the carbons to which they are attached form a carbonyl, a methanthione, a heterocyclic ring, a cycloalkyl group, an aryl group, an oxime, an imine, a hydrazone, a bridged cycloalkyl group.

R₀ and R₉ are each independently a hydrogen, an alkyl, a cycloalkoxy, a halogen, a hydroxy, an hydroxyalkyl, an alkoxyalkyl, an arylheterocyclic ring, an alkylaryl, an alkylcycloalkyl, an alkylheterocyclic ring, a cycloalkylalkyl, a cycloalkylthio, an arylalkylthioalkyl, an alkylthioalkyl, a cycloalkenyl, an heterocyclicalkyl, an alkoxy, a haloalkoxy, an amino, an alkylamino, a dialkylamino, an arylamino, a diarylamino, an alkylarylamino, an alkoxyhaloalkyl, a sulfonic acid, a sulfonic ester, an alkylsulfonic acid, an arylsulfonic acid, an arylalkoxy, an alkythio, an arylthio, a cyano an aminoalkyl, an aminoaryl, an aryl, an arylalkyl, an alkylaryl, a carboxamido, an alkylcarboxamido, an
arylcarboxamido, an amidyl, a carboxyl, a carbamoyl, an alkylcarboxylic acid, an alkylcarboxylic acid, an alkylcarbonyl, an arylcarbonyl, an ester, a carboxylic ester, an alkylcarboxylic ester, an arylcarboxylic ester, a sulfonamido, an alkylsulfonamido, an arylsulfonamido, an alkylsulfonyl, an alkylsulfonfolyxoy, an arylsulfonyl, arylsulphonyloxy, a sulfonic ester, an alkyl ester, an aryl ester, a urea, a phosphoryl, a nitro, -$\text{U}_3$-$\text{V}_5$, $\text{V}_6$, or $\text{R}_1$ and $\text{R}_2$ taken together with the carbons to which they are attached form a carbonyl, a methanthial, a heterocyclic ring, a cycloalkyl group, an aryl group, an oxime, an imine, a hydrazone a bridged cycloalkyl group,

![Diagram](image)

$\text{U}_3$ is an oxygen, sulfur or -N($\text{R}_4$)$\text{R}_4$;

$\text{V}_5$ is -NO or -NO$_2$ (i.e. an oxidized nitrogen);

$k_1$ is an integer from 1 to 3;

$\text{R}_3$ is a lone pair of electrons, a hydrogen or an alkyl group;

$\text{R}_1$ is a hydrogen, an alkyl, an aryl, an alkylcarboxylic acid, an arylcarboxylic acid, an alkylcarboxylic ester, an arylcarboxylic ester, an alkylcarboxamido, an arylcarboxamido, an alkylaryl, an alkylsulfanyl, an alkylsulfonfolyxoy, an arylsulfanyl, an arylsulfonfolyxoy, an alkylsulfonamido, a sulfonamido, a carboxamido, a carboxylic ester, an aminoalkyl, an aminoaryl, -CH$_2$-C-(U$_3$-V$_5$)(R$_4$)(R$_4$), a bond to an adjacent atom creating a double bond to that atom or -(N$_2$O$_2$)-$\text{M}_1^+$, wherein $\text{M}_1^+$ is an organic or inorganic cation; and

with the proviso that the compound of Formula (I) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., *) to at least one carboxylic acid group and/or acidic nitrogen group;

wherein the compound of Formula (II) is:
wherein

$R_{37}$ is as defined herein; and

with the proviso that the compound of Formula (II) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group;

wherein the compound of Formula (III) is:

(III)

wherein:

$X_3$ and $Y_3$ are as defined herein; and

with the proviso that the compound of Formula (III) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group and/or acidic nitrogen group;

wherein the compound of Formula (IV) is:
wherein:
$X_3$ and $Y_3$ are as defined herein; and

with the proviso that the compound of Formula (IV) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group and/or acidic nitrogen group;

wherein the compound of Formula (V) is:

wherein:
$X_3$ and $Y_3$ are as defined herein; and

with the proviso that the compounds of Formula (V) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., •) to at least one carboxylic acid group and/or acidic nitrogen group;

wherein the compound of Formula (VI) is:

wherein:
$X_3$ and $Y_3$ are as defined herein; and
with the proviso that the compounds of Formula (VI) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., \( \cdot \)) to at least one carboxylic acid group and/or acidic nitrogen group;

wherein the compound of Formula (VII) is:

\[
\begin{align*}
\text{(VII)} \\
\end{align*}
\]

\[
\begin{align*}
Y_3 & \quad R_{47} \quad X_3 \quad R_{47} \\
\end{align*}
\]

\[
\begin{align*}
\text{wherein:} \\
R_{47} & \text{ is a lower alkyl or } -(C(R_p)R_h)_k-V_4; \\
X_3, Y_3, R_p, R_h, V_4, K & \text{ and } k \text{ are as defined herein; and} \\
\end{align*}
\]

with the proviso that the compounds of Formula (VII) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., \( \cdot \)) to at least one carboxylic acid group and/or acidic nitrogen group;

wherein the compound of Formula (VIII) is:

\[
\begin{align*}
\text{(VIII)} \\
\end{align*}
\]

\[
\begin{align*}
Y_3 & \quad \text{N} \\
\end{align*}
\]

\[
\begin{align*}
\text{wherein:} \\
X_3 & \text{ and } Y_3 \text{ are as defined herein; and} \\
\end{align*}
\]

with the proviso that the compounds of Formula (VIII) must contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., \( \cdot \)) to at least one carboxylic acid group and/or acidic nitrogen group.

2. A composition comprising the compound of claim 1 and a pharmaceutically acceptable carrier.

3. The compound of claim 1, wherein the compound of Formula (I) is an organic nitric oxide enhancing salt of abitesartan, an organic nitric oxide enhancing salt of candesartan, an organic nitric oxide enhancing salt of CV-11974, an organic nitric oxide enhancing salt of elisartan analogue, an organic nitric oxide enhancing salt of embutartan, an

4. The compound of claim 1, wherein the compound of Formula (I) is an organic nitric oxide enhancing salt of abitesartan of Formula (IX), an organic nitric oxide enhancing salt of CV-11974 of Formula (X), an organic nitric oxide enhancing salt of elisartan of Formula (XI), an organic nitric oxide enhancing salt of embusartan of Formula (XII), an organic nitric oxide enhancing salt of enoltasosartan of Formula (XIII), an organic nitric oxide enhancing salt of fonsartan of Formula (XIV), an organic nitric oxide enhancing salt of forasartan of Formula (XV), an organic nitric oxide enhancing salt of glycylosoartan of Formula (XVI), an organic nitric oxide enhancing salt of irbesartan of Formula (XVII), an organic nitric oxide enhancing salt of losartan of Formula (XVIII), an organic nitric oxide enhancing salt of olmesartan of Formula (XIX), an organic nitric oxide enhancing salt of milfasartan of Formula (XX), an organic nitric oxide enhancing salt of pomisartan of Formula (XXI), an organic nitric oxide enhancing salt of ripisartan of Formula (XXII), an organic nitric oxide enhancing salt of tasosartan of Formula (XXIII), an organic nitric oxide enhancing salt of telmisartan of Formula (XXIV), an organic nitric oxide enhancing salt of valsartan of Formula (XXV), a nitric oxide enhancing analogue related to EMD 66684 of Formula (XXVI); a nitric oxide enhancing EXP 3134 of Formula (XXVII); a nitric oxide enhancing MK-996 of Formula (XXVIII); the organic nitric oxide enhancing salt of angiotensin II antagonists of Formula (II) is an organic nitric oxide enhancing salt of eprosartan of Formula (XXIX); and the organic nitric oxide enhancing salt of angiotensin II antagonist of Formula (III) is an organic nitric oxide enhancing salt of saprisartan of Formula (XXX), or an organic nitric oxide enhancing salt of zolasartan of Formula (XXXI),

wherein the compound of Formula (IX) is:

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SUBSTITUTE SHEET (RULE 26)
and the compound of Formula (X) is:

and the compound of Formula (XI) is:

and the compound of Formula (XII) is:
and the compound of Formula (XIII) is:

(XIII)

and the compound of Formula (XIV) is:

(XIV)

and the compound of Formula (XV) is:

(XV)

and the compound of Formula (XVI) is:
and the compound of Formula (XVII) is:

and the compound of Formula (XVIII) is:

and the compound of Formula (XIX) is:

and the compound of Formula (XX) is:
and the compound of Formula (XXI) is:

and the compound of Formula (XXII) is:

and the compound of Formula (XXIII) is:
and the compound of Formula (XXIV) is:

and the compound of Formula (XXV) is:

and the compound of Formula (XXVI) is:
and the compound of Formula (XXVII) is:

![Formula XXVII](image)

(XXVII)

and the compound of Formula (XXVIII) is:

![Formula XXVIII](image)

(XXVIII)

and the compound of Formula (XXIX) is:

![Formula XXIX](image)

(XXIX)

and the compound of Formula (XXX) is:
and the compound of Formula (XXXI) is:

wherein
$R_{35}$ is $-\text{CO}(\text{O})\text{U}_{3}D_{1}$ or $-\text{COOH} \cdot Z_{1}$;
$R_{36}$ is a hydrogen or $-\text{H} \cdot Z_{1}$;
nBu is the lower alkyl group $\text{CH}_{3} - \text{CH}_{2} - \text{CH}_{2} - \text{CH}_{2}$;
nPr is the lower alkyl group $\text{CH}_{3} - \text{CH}_{2} - \text{CH}_{2}$;
$\text{iPr}$ is the lower alkyl group $(\text{CH}_{3})_{2} - \text{CH}$;
$\text{OEt}$ is the alkoxy group $-\text{OCH}_{2} - \text{CH}_{3}$;
$Z_{1}$ is:

(1) ![Diagram](image1)
(2) ![Diagram](image2)
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**SUBSTITUTE SHEET (RULE 26)**
U₃ and D₁ are as defined herein; and

with the proviso that the compounds of Formula (IX) to Formula (XXXI) must
contain at least one organic nitric oxide enhancing compound linked via a salt bridge (i.e., • or □) to at least one carboxylic acid group and/or acidic nitrogen group in the compounds of
Formula (IX) to Formula (XXXI).

5. A method for treating a cardiovascular disease in a patient in need thereof
comprising administering to the patient an effective amount of the composition of claim 2.

6. The method of claim 5, wherein the cardiovascular disease is heart failure,
restenosis, hypertension, diastolic dysfunction, a coronary artery disease, myocardial
infarction, cerebral infarction, arterial stiffness, atherosclerosis, atherogenesis,
cerebrovascular disease, angina, aneurysm, ischemic heart disease, cerebral ischemia,
myocardial ischemia, thrombosis, platelet aggregation, platelet adhesion, smooth muscle cell
proliferation, a vascular or non-vascular complications associated with the use of a medical
device, a wound associated with the use of a medical device, vascular or non-vascular wall
damage, peripheral vascular disease, neointimal hyperplasia following percutaneous
transluminal coronary angiograph, vascular grafting, coronary artery bypass surgery,
thromboembolic events, post-angioplasty restenosis, coronary plaque inflammation,
hypercholesterolemia, embolism, stroke, shock, arrhythmia, atrial fibrillation or atrial flutter,
thrombotic occlusion and reclusion cerebrovascular incidents, left ventricular dysfunction and
hypertrophy.

7. The method of claim 6, wherein the cardiovascular disease is hypertension,
heart failure, arterial stiffness, postmyocardial infarction, stroke and/or diastolic dysfunction.

8. A method for treating a renovascular disease in a patient in need thereof
comprising administering to the patient an effective amount of the composition of claim 2.

9. The method of claim 8, wherein the renovascular disease is renal failure, renal
insufficiency, renal deterioration associated with severe hypertension or renovascular
hypertension.

10. A method for treating diabetes; treating diseases resulting from oxidative
stress; treating endothelial dysfunctions; treating a disease caused by endothelial

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SUBSTITUTE SHEET (RULE 26)
dysfunctions; treating cirrhosis; treating pre-eclampsia; treating osteoporosis; treating nephropathy; treating a peripheral vascular disease; treating portal hypertension; treating an ophthalmic disorder; treating metabolic syndrome; or treating hyperlipidemia in a patient in need thereof comprising administering to the patient an effective amount of the composition of claim 2.

11. The composition of claim 2, further comprising (i) at least one therapeutic agent; (ii) at least one nitric oxide enhancing compound; or (iii) at least one therapeutic agent and at least one nitric oxide enhancing compound.

12. The composition of claim 11, wherein the therapeutic agent is an aldosterone antagonist, an α-adrenergic receptor agonist, an α-adrenergic receptor antagonist, an angiotensin II antagonist, an angiotensin-converting enzyme inhibitor, an antidiabetic compound, an anti-hyperlipidemic compound, an antimicrobial compound, an antioxidant, an antithrombotic and vasodilator compound, a β-adrenergic antagonist, a calcium channel blocker, a carbonic anhydrase inhibitor, a digitalis, a diuretic, an endothelin antagonist, a hydralazine compound, a H₂ receptor antagonist, an neutral endopeptidase inhibitor, a nonsteroidal antiinflammatory compound, a phosphodiesterase inhibitor, a potassium channel blocker, a platelet reducing agent, a prostaglandin, a proton pump inhibitor, a renin inhibitor, a selective cyclooxygenase-2 inhibitor, a steroid, or a combination of two or more thereof.

13. The composition of claim 12, wherein the therapeutic agent is at least one compound selected from the group consisting of an aldosterone antagonist, an angiotensin II antagonist, an angiotensin-converting enzyme (ACE) inhibitor, a β-adrenergic antagonist, a calcium channel blocker, a diuretic, a hydralazine compound and a renin inhibitor.

14. The composition of claim 13, wherein the aldosterone antagonist is eplerenone or spironolactone; the angiotensin II antagonist is candesartan, candesartan cilexetil, eprosartan mesylate, irbesartan, losartan, medoxomil, telmisartan, trandolapril, trandolaprilat or valsartan; the angiotensin-converting enzyme inhibitor is benazepril hydrochloride, captopril, enalapril maleate, fosinopril sodium, lisinopril, moexipril hydrochloride, quinapril hydrochloride, ramipril; the β-adrenergic antagonist is bisoprolol fumarate, carvedilol, metoprolol tartrate, propranolol hydrochloride or timolol maleate; the calcium channel blockers is amlopidine, diltiazem, isradipine, nicardipine, nifedipine, nimodipine, nisoldipine, nitrendipine, verapamil; the diuretic is amiloride hydrochloride, chlorthalidone, hydrochlorothiazide or triamterene; the hydralazine compound is hydralazine hydrochloride; and the renin inhibitor is aliskiren, ciprokiren, ditekiren, enalkrein, medullipin, remikiren, terikiren, tonin or zankiren.
15. The composition of claim 11, wherein the nitric oxide enhancing compound is selected from the group consisting of a S-nitrosothiol, a nitrite, a nitrate, a S-nitrosothiol, a sydnonimine, a NONOate, a N-nitrosoamine, a N-hydroxyl nitrosamine, a nitrosimine, a diazetine dioxide, an oxatriazole 5-imine, an oxime, a hydroxylamine, a N-hydroxyguanidine, a hydroxyurea, a furoxan or a nitroxide.

16. The method of claims 5, 8 or 10, further comprising administering (i) at least one therapeutic agent; (ii) at least one nitric oxide enhancing compound (iii) at least one therapeutic agent and at least one nitric oxide enhancing compound.

17. The method of claim 16, wherein the therapeutic agent is is an aldosterone antagonist, an α-adrenergic receptor agonist, an α-adrenergic receptor antagonist, an angiotensin II antagonist, an angiotensin-converting enzyme inhibitor, an antidiabetic compound, an anti-hyperlipidemic compound, an antimicrobial compound, an antioxidant, an antithrombotic and vasodilator compound, a β-adrenergic antagonist, a calcium channel blocker, a carbonic anhydrase inhibitor, a digitali, a diuretic, an endothelin antagonist, a hydralazine compound, a H₂ receptor antagonist, a neutral endopeptidase inhibitor, a nonsteroidal antiinflammatory compound, a phosphodiesterase inhibitor, a potassium channel blocker, a platelet reducing agent, a prostaglandin, a proton pump inhibitor, a renin inhibitor, a selective cyclooxygenase-2 inhibitor, a steroid, or a combination of two or more thereof.

18. The method of claim 16, wherein the nitric oxide enhancing compound is selected from the group consisting of a S-nitrosothiol, a nitrite, a nitrate, a S-nitrosothiol, a sydnonimine, a NONOate, a N-nitrosoamine, a N-hydroxyl nitrosamine, a nitrosimine, a diazetine dioxide, an oxatriazole 5-imine, an oxime, a hydroxylamine, a N-hydroxyguanidine, a hydroxyurea, a furoxan or a nitroxide.

19. A kit comprising at least one compound of claim 1.

20. The kit of claim 19, further comprising further comprising (i) at least one therapeutic agent; (ii) at least one nitric oxide enhancing compound; or (iii) at least one therapeutic agent and at least one nitric oxide enhancing compound.

21. The kit of claim 20, wherein the (i) at least one therapeutic agent; (ii) at least one nitric oxide enhancing compound; or (iii) at least one therapeutic agent and at least one nitric oxide enhancing compound are in the form of separate components in the kit.

22. A compound selected from the group consisting of: L-valine, N-(1-oxopentyl)-N-[[2'-(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl]methyl]; salt with 3-pyridinecarboxamide, N-[3-(nitroxypropyl)- (1:2); L-valine, N-(1-oxopentyl)-N-[[2'-(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl]methyl]; salt with 1-
propanol, 3-amino-, nitrate (ester) (1:2); 
L-valine, N-(1-oxopentyl)-N-[[2'-(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl]methyl]; salt with 1,3-propanediol, 2-amino-2-[(nitrooxy)methyl]-, dinitrate (ester);
L-valine, N-(1-oxopentyl)-N-[[2'-(1H-tetrazol-5-yl)[1,1'-biphenyl]-4-yl]methyl]; salt with 2-propanediol, 3-amino-, dinitrate (ester), (2R)-.