Title
Multiple drug delivery from a balloon and a prosthesis

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Applicant(s)
Abbott Laboratories

Inventor(s)
Cromack, Keith R.; Von Oepen, Randolf; Burke, Sandra E.; Toner, John L.

Agent / Attorney
Spruson & Ferguson, Level 35 St Martins Tower 31 Market Street, Sydney, NSW, 2000

Related Art
US 5591227 (Dinh et al) 7th January 1997
Title: MULTIPLE DRUG DELIVERY FROM A BALLOON AND A PROSTHESIS

Abstract: Disclosed is an interventional device for delivery of therapeutic agents from an angioplasty balloon and from a prosthesis such as an intraluminal stent. The invention also relates to the method of loading the beneficial agents onto the balloon and the device, as well as the method of delivery of the agents from separate surfaces. The invention also relates to an interventional device having a prosthesis surface that is loaded with a first beneficial agent, and a balloon surface loaded with a second beneficial agent. The invention also relates to a method of loading multiple beneficial agents onto the prosthesis surfaces and the balloon surfaces, and to a method of manufacturing an interventional device for the delivery of a first beneficial agent and a second beneficial agent from separate surfaces.
MULTIPLE DRUG DELIVERY FROM A BALLOON AND A PROSTHESIS

BACKGROUND OF THE INVENTION

Related Application

The present invention relates to an interventional device for delivery of therapeutic agents from an angioplasty balloon and from a prosthesis such as an intraluminal stent. The invention also relates to the method of loading the beneficial agents onto the balloon and the medical device, as well as the method of delivery of the agents from separate surfaces. The invention also relates to an interventional device having a prosthesis surface that is loaded with a first beneficial agent, and a balloon surface loaded with a second beneficial agent. The invention also relates to a method of loading multiple beneficial agents onto the prosthesis surfaces and the balloon surfaces, and to a method of manufacturing an interventional device for the delivery of a first beneficial agent and a second beneficial agent from separate surfaces.

Description of Related Art

Balloon angioplasty associated with the implantation of a vascular stent is a procedure designed to expand occluded blood vessels, resulting in adequate perfusion of distal tissues. The stent, which is crimped onto the balloon, is introduced via a peripheral artery, and advanced to the lesion site over a guidewire. Inflation of the balloon results in compression of plaque and simultaneous implantation of the stent, which acts as a scaffold to keep the vessel expanded to its normal diameter. The balloon is then deflated, allowing removal of the catheter assembly, leaving the stent in place to maintain patency of the vessel.

This percutaneous intervention, described as PCI when associated with coronary balloon angioplasty, has been effective in normalizing the vessel lumen, and providing relief of pain often associated with myocardial ischemia. The procedure is not restricted to the coronary vasculature, but may also be applied to other vessels, including renal, carotid, iliac and superficial femoral arteries. However, although the success of the intervention is generally high, the long-term patency of the vessel is often reduced by restenosis of the vessel at the site of the original lesion. This restenotic process is the consequence of a variety of
factors acting in concert to re-occlude the vessel, reducing blood flow and nutrient supply to tissues. These include progression of the underlying disease, as well as the generation of cytokines and other growth factors which promote cell proliferation. These factors emanate from a variety of inflammatory cell types including monocytes and macrophages. In addition to inflammation and cell proliferation, migration of cells from the medial or adventitial layers of the vessel wall may contribute to the growth of a new layer, described as neointima, which re-occludes the vessel. In recent years, the use of bare metal stents, while effective in the short-term, has been associated with a significant rate of restenosis. Therefore, many investigators have sought to provide technologies to reduce the restenosis rate, while maintaining the beneficial effects offered by these metal scaffolds. The coating of stents with bioinert polymers has been somewhat effective, but the most important advance in this field has been the loading of these polymers with drugs known to block cell proliferation. One commonly applied technique for the local delivery of a drug is through the use of a polymeric carrier coated onto the surface of a stent, as disclosed in Berg et al., U.S. Pat. No. 5,464,650, the disclosure of which is incorporated herein by reference. Such conventional methods and products generally have been considered satisfactory for their intended purpose. The gradual elution of drug from the polymer is known to impact the restenotic process, providing beneficial concentrations of the beneficial agent at a time when the inflammatory and proliferative processes are thought to be most prevalent. The introduction of these drug-eluting stents (DES) has reduced the restenosis rate from 20 – 30% to less than 10% in several clinical trials. However, many are attempting to reduce the rate even further, providing nearly all patients who receive a DES with long-term vessel patency and minimal chance of return to the cath lab for repeat procedures. The delivery of multiple drugs, using both the stent and the balloon itself as delivery platforms, may help to achieve this goal.

As evident from the related art, conventional methods of loading interventional devices with beneficial agents, such as drugs, often requires coating the entire prosthesis with a polymer capable of releasing beneficial drugs, as disclosed in Campbell, U.S. 5,649,977 and Dinh et al., U.S. Patent No. 5,591,227, the disclosures of which are incorporated by reference.
Object of the Invention

It is the object of the present invention to substantially overcome or ameliorate one or more of the disadvantages of the prior art.

Summary of the Invention

The present invention proposes the use of one or more beneficial agents, applied to the surface of the balloon material by any method, and the application of one or more beneficial agents applied to either the bare-metal surface of a second device, or incorporated with the polymer which coats the second device. The delivery of the beneficial agent from the balloon is expected to occur during either pre-dilatation of the vessel at the lesion site, or from the balloon during the delivery of the device during a stenting procedure. Additionally, the delivery of the beneficial agent can be from the balloon during a final stent sizing balloon expansion. The delivery of the beneficial agent from the prosthesis is expected to occur over a longer period, as the drug is released from the polymer or from the surface of the device. The associated prosthesis may be placed directly when the balloon is inflated at the lesion site, immediately after as commonly practiced in pre-dilatation procedures, or within a suitable time period in a second interventional procedure.

The present invention provides a system for delivering a beneficial agent, comprising:

a) a balloon having an outer surface that is at least partially coated with a therapeutically effective amount of a first beneficial agent; and

b) a prosthesis that can be delivered or implanted having a surface that is at least partially coated with a therapeutically effective amount of a second beneficial agent

wherein the system is useful for the treatment and prevention of vascular diseases.

Preferably, the balloon is an angioplasty balloon.

Preferably, the prosthesis is a stent.

Preferably, said prosthesis is crimped onto said balloon.

Preferably, said prosthesis is at least partially retained on said balloon by said coating on said balloon.

Preferably, the stent is a self-expanding stent.

Preferably, the coating on the balloon further comprises a carrier for said first beneficial agent.

Preferably, the carrier is selected from the group consisting of complex sugars (mannitol) starches (cellulose), collagens, and polymeric materials.
Preferably, the coating on the prosthesis further comprises a carrier for said second beneficial agent.

Preferably, said carrier for said second beneficial agent is selected from a group consisting of phosphorylcholine, polycaprolactone, poly-D,L-lactic acid, poly-L-lactic acid, poly(lactide-co-glycolide), poly(hydroxybutyrate), poly(hydroxybutyrate-co-valerate), polydioxanone, polyorthoester, polyanhydride, poly(glycolic acid), poly(glycolic acid-co-trimethylene carbonate), polyphosphoester, polyphosphoester urethane, poly(amino acids), cyanoacrylates, poly(trimethylene carbonate), poly(iminocarbonate), polyalkylene oxalates, polyphosphazenes, polyiminocarbonates, and aliphatic polycarbonates, fibrin, fibrinogen, cellulose, starch, collagen, one or more of a series of polymers synthesized by the pyrolytic, vapor-decomposition polymerization of p-xylylene dimers, or a plasma enhanced coating thereof, polyurethane, polycarbonate urethanes, polyethylene, polyethylene terephthalate, ethylene vinyl acetate, ethylene vinyl alcohol, silicone polysiloxanes, substituted polysiloxanes, polyethylene oxide, polybutylene terephthalate-co-PEG, PCL-co-PEG, PLA-co-PEG, polyacrylates, polyvinyl pyrrolidone, polyacrylamide, thermoplastic elastomers, polyolefin elastomers, EPDM rubbers, polyamide elastomers, biostable plastic, acrylic polymers, nylon, polyesters, epoxies and derivatives or combinations thereof.

Preferably, the system further comprises a balloon which is substantially free of a beneficial agent.

Preferably, the first beneficial agent is the same as the second beneficial agent.

Preferably, a surface of said prosthesis is loaded with therapeutically effective amounts of more than one beneficial agent.

Preferably, the first beneficial agent and the second beneficial agent are individually selected from the group consisting of antithrombotics, anticoagulants, antiplatelet agents, anti-lipid agents, thrombolytics, antiproliferatives, antiinflammatories, agents that inhibit hyperplasia, smooth muscle cell inhibitors, antibiotics, growth factor inhibitors, cell adhesion inhibitors, cell adhesion promoters, antimitotics, antifibrins, antioxidants, antineoplastics, agents that promote endothelial cell recovery, antiallergic substances, viral vectors, nucleic acids, monoclonal antibodies, antisense compounds, oligonucleotides, cell permeation enhancers, radiopaque agent markers, HMG CoA reductase inhibitors, indomethacin, phenyl salicylate, vinblastine, ABT-627 (atrasentan), ABT-578, testosterone, progesterone, paclitaxel, taxanes, cyclosporin A, vincristine, carvedilol, vindesine, dipyridamole, methotrexate, folic acid, thrombospondin mimetics, estradiol, dexamethasone, metrizamide, iopamidol, iohexol, iopromide, iobitridol,
iomeprol, iopentol, ioversol, ioxilan, iodixanol, iotrolan, rapamycin, rapamycin
derivatives, pimecrolimus, everolimus, fenofibrate, carvedilol, taxoteres, tacrolimus,
peroxisomal proliferator activated receptor γ (PPAR-γ) agonists, paricalcitol, butylated
hydroxytoluene, butylated hydroxyanisole, vitamin E, danazol, probucol, tocopherols,
tocotrienols, colchicine, betamethasone, cortisone, budesonide, prednisolone,
methylprednisolone, hydrocortisone, flurbiprofen, ibuprofen, ketoprofen, fenoprofen,
naproxen, diclofenac, diflunisal, acetylsalicylic acid, sulindac, etodolac, ketorolac,
meclomenamic acid, piroxicam, phenylbutazone, altretamine, bendamucine, carboplatin,
carmustine, cisplatin, cyclophosphamide, fotemustine, ifosfamide, lomustine, nimustine,
prednimustine, treosulfin, docetaxel, mercaptopurine, pentostatin, trimetrexate,
gemcitabine, azathioprine, fluorouracil, doxorubicin hydrochloride, mitomycin, RPR-
101511 A, rosiglitazone, pioglitazone, lovastatin, atorvastatin, simvastatin, pravastatin,
cerivastatin, fluvastatin, ABT-518, permirolast, potassium nitroprusside,
phosphodiesterase inhibitors, prostaglandin inhibitors, suramin, serotonin blockers,
steroids, thioprotease inhibitors, triazolopyrimidine, nitric oxide, dalteparin, enoxaparin,
nadroparin, reviparin, ar doparin, certaparin, heparinoids, hirudin, argatroban, forskolin,
vapriprost, protacyclin, dextran, D-phe-pro-5-arg-chlormethylketonedipyridamole,
glycoprotein IIb/IIa, recombinant hirudin, thrombin inhibitors, urokinase, recombinant
urokinase, pro-urokinase, tissue plasminogen activator, tenecteplase, angiopeptin,
captopril, cilazapril, lisinopril, nifedipine, amlodipine, cilnidipine, lercanidipine,
benidipine, trifluperazine, diltiazem, verapamil, fibroblast growth factor antagonists, fish
oil (omega 3-fatty acid), histamine antagonists, etoposide, topotecan, tamoxifen and
combinations thereof.

Preferably, said HMG CoA reductase inhibitor is selected from the group
consisting of atorvastatin, simvastatin, pravastatin, lovastatin, cerivastatin, fluvastatin.

Preferably, the PPAR-γ agonist is a thiazolidinedione.

Preferably, the beneficial agent is a nucleic acid that encodes a pharmaceutically
useful peptide or an anti-sense oligonucleotide used to control a gene of interest in a cell
of the patient.

Preferably, the first beneficial agent has a varied local areal density as a measure
of beneficial agent per unit local surface area of the balloon surface.

Preferably, the coating on the balloon is discontinuous or forms a pattern.

Preferably, said coating on the balloon extends beyond said prosthesis cramped
on said balloon.
Preferably, said coated balloon is delivered after said stent is implanted to accomplish final sizing of said stent.

Preferably, said first beneficial agent is transferred from said coating on said balloon into said coating on said prosthesis.

Preferably, said coating on said prosthesis comprises a barrier to transfer of said first beneficial agent from said coating on said balloon into said coating on said prosthesis or of said second beneficial agent from said coating on said prosthesis into said coating on said balloon.

Preferably, said barrier is polymeric.

Preferably, the polymeric component is selected from a group consisting of phosphorylcholine, polycaprolactone, poly-D,L-lactic acid, poly-L-lactic acid, poly(lactide-co-glycolide), poly(hydroxybutyrate), poly(hydroxybutyrate-co-valerate), polydioxanone, polyorthoester, polyanhydride, poly(glycolic acid), poly(glycolic acid-co-trimethylene carbonate), polyphosphoesterurethane, poly(aminocids), cyanoacrylates, poly(trimethylene carbonate), poly(iminocarbonates), polyalkylene oxalates, polyphosphazenes, poly-iminocarbonates, and aliphatic polycarbonates, fibrin, fibrinogen, cellulose, starch, collagen, one or more of a series of polymers synthesized by the pyrolytic, vapor-decomposition polymerization of p-xylylene dimers, or a plasma enhanced coating thereof, polyurethane, polycarbonate urethanes, polyethylene, polyethylene terephthalate, ethylene vinyl acetate, ethylene vinyl alcohol, silicone polysiloxanes, substituted polysiloxanes, polyethylene oxide, polybutylene terephthalate-co-PEG, PCL-co-PEG, PLA-co-PEG, polyacrylates, polyvinyl pyrrolidone, polyacrylamide, thermoplastic elastomers, polyolefin elastomers, EPDM rubbers, polyamide elastomers, biostable plastic, acrylic polymers, nylon, polyesters, epoxies and derivatives or combinations thereof.

The present invention also provides a method of treating and preventing a vascular disease using the above-described system.

Preferably, said first beneficial agent is delivered from said coating on said prosthesis into a target site.

Preferably, at least one of said coating on said balloon and said coating on said prosthesis contains multiple beneficial agents.

The present invention further provides a method of treating and preventing a vascular disease, comprising the steps of:

1. delivering a balloon having an outer surface that is at least partially coated with a therapeutically effective amount of a first beneficial agent; and
b) delivering a prosthesis having a surface that is at least partially coated with a therapeutically effective amount of a second beneficial agent.

Preferably, the first beneficial agent is not the same as the second beneficial agent.

Preferably, step a and step b are performed simultaneously.

Preferably, said balloon is adhered to said prosthesis by said coating on said balloon.

Preferably, step a and step b are performed in sequence.

Preferably, the method further comprises a step of delivering an uncoated balloon.

Preferably, said prosthesis is a self-expanding stent.

Preferably, said coated balloon is used for final sizing of said coated stent.

The present invention further provides a method of providing a device for treatment and prevention of vascular disease, comprising the steps of:

- providing a beneficial agent; and
- coating said beneficial agent onto a balloon, wherein the balloon includes a balloon surface having a working length, the working length of the balloon surface is at least partially coated with the beneficial agent, the beneficial agent having a varied local areal density as a measure of beneficial agent per unit local surface area on the working length of the balloon surface.

Preferably, said coating step is accomplished by dipping, spraying, layering, painting or jetting.

Preferably, said beneficial agent is coated from a solution.

Preferably, said solution comprises at least one solvent.

Preferably, said at least one solvent is selected from the group consisting of water, alcohols, dichloromethane, dimethylsulfoxide, ethanol, isopropyl alcohol, isobutyl alcohol, tetrahydrofuran, alkanes, dimethylformamide and mixtures thereof.

Preferably, the method further comprises a step of providing a prosthesis for use in association with said coated balloon.

Preferably, the method further comprises a step of enhancing retention of said prosthesis by mounting said prosthesis on said coated balloon prior to complete drying or curing of said coating on said balloon.

The present invention further provides a method for treating vascular disease in a bifurcated vessel, comprising:
deploying a balloon within a first branch of a bifurcated vessel, wherein the balloon includes a balloon surface having a working length, the working length of the balloon surface is at least partially coated with at least one beneficial agent, the at least one beneficial agent having a varied local areal density as a measure of beneficial agent per unit local surface area on the working length of the balloon surface; and delivering a therapeutically effective amount of the at least one beneficial agent to the first branch of the bifurcated vessel.

Preferably, the method further comprises deploying a first stent device within the first branch of the bifurcated vessel.
It is to be understood that both the foregoing general description and the following
detailed description are exemplary and are intended to provide further explanation of the
invention claimed.

The accompanying Figures, which are incorporated in and constitute part of this
specification, are included to illustrate and provide a further understanding of the method and
system of the invention. Together with the description, the Figures serve to explain the
principles of the invention.

BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1 is a schematic representation of an angioplasty procedure and stent
placement equipment showing a balloon on a catheter and the syringe systems used to inflate
the balloon.

Figure 2a is a schematic representation of a stent crimped onto a catheter balloon.
Figure 2b shows a blowup of the balloon and stents section of the catheter with the shading
on the balloon representing a coating of a second beneficial agent and the shading of the stent
struts representing a coating of a first beneficial agent.

Figure 3 is a schematic representation of an embodiment of the system of the present
invention showing a cross section through a stent crimped onto a catheter balloon. The dark
center is the catheter body, the white is the balloon, the squares are the individual struts of the
stent, the shading on the balloon representing a coating of a second beneficial agent on the
balloon and the shading of the stent struts representing a coating of a first beneficial agent on
the stent.

Figure 4 is a schematic representation of the embodiment of the system of the present
invention for the delivery of the beneficial agents to a vessel wall. The drawing shows the
process of delivering a stent from a balloon to expand the lumen of a narrowed vessel. 4a.
Shows the placement of the balloon-stent combination at the site of delivery. 4b. shows the
expansion of the balloon, which results in the expansion of the stent against the vessel wall.
4c show the result after the balloon is deflated and removed leaving the stent behind.

Figure 5a-c is a schematic representation of a prosthesis or balloon loaded with
beneficial agent having a first portion and a second portion having different local areal
densities of beneficial agent in accordance with the present invention, and graph depicting
 corresponding areal density.
DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

Reference will now be made in detail to the present preferred embodiments of the method and system for loading a first beneficial agent onto a prosthesis, and a second beneficial agent onto a balloon. Wherever possible, the same reference characters will be used throughout the drawings to refer to the same or like parts.

In accordance with the present invention, a system is provided for delivery of beneficial agents within a lumen. Particularly, the present invention provides a system including a prosthesis having a first beneficial agent and a balloon having second beneficial agent where the beneficial agents are delivered for treatment and prevention of vascular or other intraluminal diseases.

As used herein "interventional device" refers broadly to any device suitable for intraluminal delivery or implantation. For purposes of illustration and not limitation, examples of such interventional devices include stents, grafts, stent-grafts, and the like. As is known in the art, such devices may comprise one or more prostheses, each having a first cross-sectional dimension or profile for the purpose of delivery and a second cross-sectional dimension or profile after deployment. Each prosthesis may be deployed by known mechanical techniques such as balloon expansion deployment techniques, or by electrical or thermal actuation, or self-expansion deployment techniques, as well known in the art.


For purposes of explanation and illustration, and not limitation, an exemplary embodiment of the interventional device in accordance with the invention is shown schematically in Figure 2. In accordance with one aspect of the invention, as shown schematically in Figure 2, the interventional device generally includes a prosthesis loaded with beneficial agent to provide a local delivery of a first beneficial agent across a treatment zone and a balloon with a second beneficial agent delivered a cross a second overlapping treatment zone. Particularly, as embodied herein the prosthesis may be a stent, a graft or a stent-graft, as previously noted, for intravascular or coronary delivery and implantation. However, the prosthesis may be any type of implantable member capable of being loaded
with beneficial agent. The balloon may be any type of catheter based expandable entity that can act to expand the prosthesis, the local tissue, or push the second beneficial agent against the lumen wall.

The prosthesis can be in an expanded or unexpanded state during the loading of beneficial agent. The underlying structure of the prosthesis can be virtually any structural design and the prosthesis can be composed of any suitable material such as, but not limited to, stainless steel, “MP35N,” “MP20N,” elastinite (Nitinol), tantalum, nickel-titanium alloy, platinum-iridium alloy, gold, magnesium, polymer, ceramic, tissue, or combinations thereof. “MP35N” and “MP20N” are understood to be trade names for alloys of cobalt, nickel, chromium, and molybdenum available from Standard Press Steel Co., Jenkintown, PA. “MP35N” consists of 35% cobalt, 35% nickel, 20% chromium, and 10% molybdenum. “MP20N” consists of 50% cobalt, 20% nickel, 20% chromium and 10% molybdenum. The prosthesis can be made from bioabsorbable or biostable polymers. In some embodiments, the surface of the prosthesis can include one or more reservoirs or cavities formed therein, as described further below.

The prosthesis can be fabricated utilizing any number of methods known in the art. For example, the prosthesis can be fabricated from a hollow or formed tube that is machined using lasers, electric discharge milling, chemical etching or other known techniques. Alternatively, the prosthesis can be fabricated from a sheet that is rolled into a tubular member, or formed of a wire or filament construction as known in the art.

The balloon can be in an expanded or unexpanded state during the loading of beneficial agent. Additionally, the balloon can be in a rolled or unrolled state during the loading of beneficial agent. The underlying structure of the balloon can be virtually any structural design and the balloon can be composed of any suitable material such as, but not limited to, polyester, pTFE (Teflon), nylon, Dacron, or combinations thereof. “Teflon” and “Dacron” are understood to be trade names for polymers available from DuPont Co., Wilmington, DE. In some embodiments, the surface of the balloon can include one or more reservoirs or cavities formed therein or ports for solution delivery.

The balloon can be fabricated utilizing any number of methods known in the art. For example, the balloon can be fabricated from a hollow or formed tube that is cover with thin membranes of polymer that is solution or physically (by laser or ultrasonically) welded to the tube. The inner volume of the balloon is then in direct contact with the tube such that air or...
aqueous solutions can be injected into the space under pressure to expand the balloon into any predefined shape that is of use. The surface of the balloon can be rolled to reduce the outer diameter of the final catheter balloon assemble.

The balloons can be loaded with a beneficial agent from a dilute solution of the agent made in an appropriate solvent (for example Ethanol) (if desired this solution could also contain multiple beneficial agents) and allowed to dry before the stent is crimped onto it. Alternatively, the coating could not be allowed to dry or cure past a "tacky" state before the stent is crimped onto it. This would enable the adhesion of the beneficial agent coating on the balloon to the inside of the prosthesis. This process increases the retention of the prosthesis onto the balloon (acting as a prosthesis retention enhancer) thus reducing the chance that the stent will move on the angioplasty balloon during the torturous trip to the coronary arteries. To prevent the film on the balloon from drying too quickly (i.e. becoming hard before the stent was placed over the balloon) the solution can contain a second liquid that has a higher boiling point (preferable water) and thus a slower drying time than the main solvent. Additionally, the use of a two solvent system (i.e. Ethanol-water) would allow the solvent to be adjusted such that the balloons beneficial agent (for example dexamethasone) is soluble enough to be laid down but the beneficial agent (for example ABT-578, rapamycin, and rapamycin analogies) on the prosthesis is not soluble enough to leach out of the prosthesis into the balloon coating or out of the balloon coating into the prosthesis coating during the drying time. Additionally, polymer barriers, timing layers, top or capcoats, especially on the luminal side of the prosthesis, or the use of bare metal interfaces can be used to prevent drug transfer from the balloon surface into the delivery polymer of the prosthesis. Alternately, some of the beneficial agent from the balloon could be allowed to transfer to the stent creating a gradient of the two beneficial agents released from the stent into the tissue. The binder can be composed of complex sugars (mannitol), starches (e.g., cellulose), collagens. In general the binder would be noncrystalline, have low water solubility, have good film forming characteristics, good solubility with solvents that may be used to dissolve the drug, biocompatible, inert (nonreactive with respect to the drug and also body tissues, fluids, etc), polymer, (e.g., hydrogel), can be hydrophobic if not hydrogel, especially if it is not permanently attached to balloon (if permanently attached, then can use hydrogel, can be used to absorb drug and then when balloon inflated, will squeeze out the drug into ablumenal tissue), low blood solubility if not permanently attached to balloon
The prosthesis, balloon combination can be fabricated utilizing any number of methods known in the art. For example, the prosthesis can be slipped over the end of the balloon and aligned at the center of the balloon. The prosthesis can be reduced in diameter such that as it is slipped over the end of the balloon there is a tight fit between the prosthesis and the balloon surface. Additionally, the prosthesis can be crimped onto the balloon to ensure that the prosthesis does not move during delivery of the prosthesis. The envisioned steps for this process would be: Dip or spray coat the balloon with the balloons beneficial agent, place the previously beneficial agent coated prosthesis onto a dry or tacky balloon and place Balloon/Stent into crimper and crimping.

As noted above, the prosthesis and the balloon are at least partially loaded with beneficial agent (10a, 10b, 10c). "Beneficial agent" as used herein, refers to any compound, mixture of compounds, or composition of matter consisting of a compound, which produces a beneficial or useful result. The beneficial agent can be a polymer, a marker, such as a radiopaque dye or particles, or can be a drug, including pharmaceutical and beneficial agents, or an agent including inorganic or organic drugs without limitation. The agent or drug can be in various forms such as uncharged molecules, components of molecular complexes, pharmacologically-acceptable salts such as hydrochloride, hydrobromide, sulfate, laurate, palmitate, phosphate, nitrate, borate, acetate, maleate, tartrate, olate, and salicylate.

An agent or drug that is water insoluble can be used in a form that is a water-soluble derivative thereof to effectively serve as a solute, and on its release from the device, is converted by enzymes, hydrolyzed by body pH, or metabolic processes to a biologically active form. Additionally, the agents or drug formulations can have various known forms such as solutions, dispersions, pastes, particles, granules, emulsions, suspensions and powders. The drug or agent may or may not be mixed with polymer or a solvent as desired.

For purposes of illustration and not limitation, the drug or agent can include antithrombotics, anticoagulants, antiplatelet agents, thrombolytics, lipid-lowering agents, antiproliferatives, anti-inflammatory agents, agents that inhibit hyperplasia, inhibitors of smooth muscle cell proliferation, antibiotics, growth factor inhibitors, cell adhesion promoters, or cell adhesion inhibitors. Other drugs or agents include but are not limited to antineoplastics, antimitotics, antifibrins, antioxidants, agents that promote endothelial cell recovery, antiallergic substances, radiopaque agents, viral vectors, antisense compounds, oligonucleotides, cell permeation enhancers, angiogenesis agents, and combinations thereof.
Examples of such antithrombotics, anticoagulants, antiplatelet agents, and thrombolytics include unfractionated heparin, low molecular weight heparins, such as dalteparin, enoxaparin, nadroparin, reviparin, ardoeparin and certaparin, heparinoids, hirudin, argatroban, forskolin, vaptireprost, prostacyclin and prostacyclin analogues, dextran, D-phe-pro-arg-chloromethylketone (synthetic antithrombin), dipyriramole, glycoprotein IIb/IIIa (platelet membrane receptor antagonist antibody), recombinant hirudin, and thrombin inhibitors such as Angiomax™, from Biogen, Inc., Cambridge, Mass; and thrombolytic agents, such as urokinase, e.g., Abbokinase™ from Abbott Laboratories Inc., North Chicago, IL, recombinant urokinase and pro-urokinase from Abbott Laboratories Inc., tissue plasminogen activator (Alteplase™ from Genentech, South San Francisco, CA and tenecteplase (TNKtPA).

Examples of such cytostatic or antiproliferative agents include rapamycin and its analogs such as ABT-578, i.e.,


Examples of such anti-inflammatories include colchicine and glucocorticoids such as betamethasone, cortisone, dexamethasone, budesonide, prednisolone, methylprednisolone and hydrocortisone. Non-steroidal anti-inflammatory agents include flurbiprofen, ibuprofen, ketoprofen, fenoprofen, naproxen, diclofenac, diflunisal, acetaminophen, indomethacin, sulindac, etodolac, diclofenac, ketorolac, meclofenamic acid, piroxicam and phenylbutazone.
EDITORIAL NOTE

APPLICATION NO.: 2005222719

The following Description page 15 has not been numbered.
Examples of such antineoplastics include alkylating agents such as altretamine, bendamucine, carboplatin, carmustine, cisplatin, cyclophosphamide, fotemustine, ifosfamide, lomustine, nimustine, prednimustine, and treosulfin, antimitotics such as vincristine, vinblastine, paclitaxel, e.g., TAXOL® by Bristol-Myers Squibb Co., Stamford, Conn., docetaxel, e.g., Taxotere® from Aventis S.A., Frankfort, Germany, antimetabolites such as methotrexate, mercaptopurine, pentostatin, trimetrexate, gemcitabine, azathioprine, and fluorouracil, and antibiotics such as doxorubicin hydrochloride, e.g., Adriamycin® from Pharmacia & Upjohn, Peapack, NJ, and mitomycin, e.g., Mutamycin® from Bristol-Myers Squibb Co., Stamford, Conn, agents that promote endothelial cell recovery such as Estradiol.

Additional drugs which may be utilized in this application include inhibitors of tyrosine kinase such as RPR-101511A, PPAR-alpha agonists such as Tricor™ (fenofibrate) from Abbott Laboratories Inc., North Chicago, IL, PPAR-gamma agonists selected from a group consisting of rosiglitazone (Glaxo Smith Kline) and Pioglitazone (Takeda), HMG CoA reductase inhibitors selected from a group consisting of lovastatin, atorvastatin, simvastatin, pravastatin, cerivastatin and fluvastatin, endothelin receptor antagonists such as ABT-627 having general formula C_{29}H_{38}N_{2}O_{6}.ClH, and the following structural formula.

![Chemical Structure](image1)

from Abbott Laboratories Inc., North Chicago, IL; matrix metalloproteinase inhibitors such as ABT-518 having general formula C_{21}H_{22}F_{3}NO_{6}S and having the following structural formula.

![Chemical Structure](image2)
from Abbott Laboratories Inc., North Chicago, IL, antiallergic agents such as permethylast potassium nitroprusside, phosphodiesterase inhibitors, prostaglandin inhibitors, suramin, serotonin blockers, steroids, thioprotease inhibitors, triazolopyrimidine, and nitric oxide.

While the foregoing beneficial agents are known for their preventive and treatment properties, the substances or agents are provided by way of example and are not meant to be limiting. Further, other beneficial agents that are currently available or may be developed are equally applicable for use with the present invention.

If desired or necessary, the beneficial agent can include a binder to carry, load, or allow sustained release of an agent, such as but not limited to a suitable polymer or similar carrier. The term "polymer" is intended to include a product of a polymerization reaction inclusive of homopolymers, copolymers, terpolymers, etc., whether natural or synthetic, including random, alternating, block, graft, branched, cross-linked, blends, compositions of blends and variations thereof. The polymer may be in true solution, saturated, or suspended as particles or supersaturated in the beneficial agent. The polymer can be biocompatible, or biodegradable.

For purpose of illustration and not limitation, the polymeric material include phosphorylcholine linked macromolecules, such as a macromolecule containing pendant phosphorylcholine groups such as poly(MPC<sub>x</sub>:LMA<sub>y</sub>:HPMA<sub>z</sub>:TSMA<sub>z</sub>), where MPC is 2-methacryloyloxyethylphosphorylcholine, LMA is lauryl methacrylate, HPMA is hydroxypropyl methacrylate and TSMA is trimethoxysilylpropyl methacrylate, polycaprolactone, poly-D,L-lactic acid, poly-L-lactic acid, poly(lactide-co-glycolide), poly(hydroxybutyrate), poly(hydroxybutyrate-co-valerate), polydioxanone, polymers, polyanhydride, poly(glycolic acid), poly(glycolic acid-co-trimethylene carbonate), polyphosphoester, polyphosphoester urethane, poly(amino acids), cyanoacrylates, poly(trimethylene carbonate), poly(iminocarbonates), polyalkylene oxalates, polyphosphazenes, polyiminocarbonates, and aliphatic polycarbonates, fibrin, fibrinogen, cellulose, starch, collagen, Parylene®, Parylast®, polyurethane including polycarbonate urethanes, polyethylene, polyethylene terephthalate, ethylene vinyl acetate, ethylene vinyl alcohol, silicone including polysiloxanes and substituted polysiloxanes, polyethylene oxide, polybutylene terephthalate-co-PEG, PCL-co-PEG, PLA-co-PEG, polyacrylates, polyvinyl pyrrolidone, polyacrylamide, and combinations thereof. Non-limiting examples of other
suitable polymers include thermoplastic elastomers in general, polyolefin elastomers, EPDM rubbers and polyamide elastomers, and biostable plastic material such as acrylic polymers, and its derivatives, nylon, polyesters and epoxies. Preferably, the polymer contains pendant phosphoryl groups as disclosed in U.S. Patent Nos. 5,705,583 and 6,090,901 to Bowers et al. and U.S. Patent No. 6,083,257 to Taylor et al., which are all incorporated herein by reference.

The beneficial agent can include a solvent. The solvent can be any single solvent or a combination of solvents. For purpose of illustration and not limitation, examples of suitable solvents include water, aliphatic hydrocarbons, aromatic hydrocarbons, alcohols, ketones, dimethyl sulfoxide, tetrahydrofuran, dihydrofuran, dimethylacetamide, acetates, and combinations thereof. Preferably, the solvent is ethanol. More preferably, the solvent is isobutanol. Additionally, in another aspect of the invention, multiple beneficial agents are dissolved or dispersed in the same solvent. For purpose of illustration and not for limitation, dexamethasone, estradiol, and paclitaxel are dissolved in isobutanol. Alternatively, dexamethasone, estradiol, and paclitaxel are dissolved in ethanol. In yet another example, dexamethasone, estradiol, and ABT-578, i.e., the rapamycin analog, 3S,6R,7E,9R,10R,12R,14S,15E,17E,19E,21S,23-
S,26R,27R,34aS)9,10,12,13,14,21,22,23,24,25,26,27,32,33,34,34a-Hexadecahydro-9,27-
dihydroxy-3-[(1R)-2-[(1S,3R,4R)-3-methoxy-4-tetrazol-1-yl)cyclohexyl]-1-methylethyl]-10,21-dimethoxy-6,8,12,14,20,26-hexamethyl-23,27-epoxy-3H-pyrido[2,1-c][1,4]oxaaazacyclohentriacontine-1,5,11,12,28,29(4H,6H,31H)-pentone; 23,27-Epoxy-3H-pyrido[2,1-c] [1,4]oxaaazacyclohentriacontine-1,5,11,28,29(4H,6H,31H)-pentone, are dissolved together in one solvent. Preferably, the solvent is ethanol. More preferably, the solvent is isobutanol.

Additionally, the beneficial agent includes any of the aforementioned drugs, agents, polymers, and solvents either alone or in combination.

A number of methods can be used to load the beneficial agent onto the surface of the prosthesis or balloon to provide for a controlled local areal density of beneficial agent. For example, the prosthesis or balloon can be constructed to include pores or reservoirs which are impregnated or filled with beneficial agent or multiple beneficial agents. The pores can be sized or spaced apart to correspond to or limit the amount of beneficial agent contained therein in accordance with the desired local areal density pattern along the length of the interventional device, wherein larger pores or more dense spacing would be provided in such portions intended to have a greater local areal density. Alternatively, uniform pores sizes can
be provided but the amount of beneficial agent loaded therein is limited accordingly. Additionally, if desired, a membrane of biocompatible material can then be applied over the pores or reservoirs for sustained or controlled release of the beneficial agent from the pores or reservoirs.

According to some of the embodiments, the beneficial agent can be loaded directly onto the prosthesis or balloon or alternatively, the beneficial agent is loaded onto a base material layer that is applied to a surface of the prosthesis or balloon. For example and not limitation, a base coating, such as a binder or suitable polymer, is applied to a selected surface of the prosthesis or balloon such that a desired pattern is formed on the prosthesis or balloon surface. Beneficial agent is then applied directly to the pattern of the base material.

In one aspect of the invention, the desired pattern corresponds to the desired controlled local areal density. For example, a greater amount of base material layer is applied to portions of the prosthesis or balloon intended to have a greater local areal density of beneficial agent, and a lesser amount of base material is applied to portions of the prosthesis or balloon intended to have a lower local areal density of beneficial agent.

Alternatively, a suitable base coating capable of retaining beneficial agent therein can be applied uniformly over the surface of the prosthesis or balloon, and then selected portions of the base coating can be loaded with the beneficial agent in accordance with the invention. A greater amount of beneficial agent would be loaded over a unit surface area of the base coating intended to have a greater local areal density and a lower amount of beneficial agent would be loaded over a unit surface area intended to have a lower local areal density.

In yet another embodiment of the present invention, the beneficial agent can be applied directly to the surface of the prosthesis or balloon. Generally a binder or similar component can be required to ensure sufficient adhesion. For example, this coating technique can include admixing the beneficial agent with a suitable binder or polymer to form a coating mixture, which is then coated onto the surface of the prosthesis or balloon. The coating mixture is prepared in higher or lower concentrations of beneficial agent as desired, and then applied to selected portions of the prosthesis or balloon appropriately. In general the binder used with the beneficial agent for the prosthesis may be different than the binder used for the beneficial agent for the balloon.
In any of the embodiments disclosed herein, a porous or biodegradable membrane or layer made of biocompatible material can be coated over the beneficial agent for sustained release thereof, if desired.

Conventional coating techniques can be utilized to coat the beneficial agent onto the surface of the prosthesis or balloon such as spraying, dipping or sputtering and still provide the desired effect if performed appropriately. With such techniques, it may be desirable or necessary to use known masking or extraction techniques to control the location and amount in which beneficial agent is loaded. Although not required, prior to coating the prosthesis or balloon with beneficial agent, optical machine vision inspection of the prosthesis or balloon may be utilized to ensure that no mechanical defects exist. Defective prostheses or balloons may be rejected before wasting beneficial agent, some of which may be very costly.

In accordance with one aspect of the invention, a method of loading beneficial agent onto a prosthesis for delivery within a lumen is disclosed. The method comprises the steps of providing a prosthesis, beneficial agent to be delivered from the prosthesis, and a fluid-dispenser having a dispensing element capable of dispensing the beneficial agent in discrete droplets, wherein each droplet has a controlled trajectory. The method further includes creating relative movement between the dispensing element and the prosthesis to define a dispensing path and selectively dispensing the beneficial agent in a raster format to a predetermined portion of the prosthesis along the dispensing path. In particular, the beneficial agent is selectively dispensed from the dispensing element to a predetermined portion of the prosthesis in a raster format along a dispensing path. As used herein “raster format” refers to a continuous or non-continuous dispensing pattern of droplets of beneficial agent.

According to another aspect of the invention, the method of loading beneficial agent onto the prosthesis includes providing a prosthesis including a tubular member having a central axis defined along a length of the tubular member. This method further includes dispensing beneficial agent.

In accordance with another aspect of the invention, additional beneficial agents or multiple beneficial agents can be loaded onto the prosthesis as described above. Therefore, further in accordance with the invention, an interventional device comprising a prosthesis loaded with a beneficial agent and additional beneficial agents is provided.
Particularly, the method described in detail above for one beneficial agent can be modified to allow for loading multiple beneficial agents onto a prosthesis and/or a balloon, which might ordinarily lead to undesirable results when using conventional loading techniques. For example and not limitation, the first beneficial agent and the second beneficial agent may have different physical and/or chemical characteristics preventing the beneficial agents from being capable of dissolving in the same solvent, or at the same pH or temperature. In particular, the first beneficial agent can be dissolved in a solvent that is immiscible with the solvent in which the second beneficial agent is dissolved. Alternatively, the first beneficial agent and the second beneficial agent may be incompatible with each other. In particular, the first beneficial agent and the second beneficial agent can be undesirably chemically reactive or may have undesirably different release rates (or contrarily, undesirably similar release rates). Additionally, the first and second beneficial agents can simply be detrimental to each other, e.g., one of the beneficial agents may degrade the efficacy of the other beneficial agent. Thus, although loading the particular multiple beneficial agents onto the same surface of a prosthesis or balloon can be desired it often may be problematic due to some incompatibility when using a conventional loading technique. In accordance with the present invention, a method of loading such beneficial agents and an interventional device that combine a prosthesis and a balloon for the delivery of such beneficial agents is provided.

As noted above, the beneficial agent can include a drug and polymer mixture. In accordance with the method of the invention, the first and second beneficial agents can correspond to drug-polymer mixtures having different concentrations of polymer to effect different release rates of the particular drug in each beneficial agent. For example, the drug-polymer mixture having a higher concentration of polymer would have a slower release of the drug within the lumen than a drug-polymer mixture having a lower concentration. Alternatively, rather than providing drug-polymer mixtures having different polymer concentrations to provide different release rates, it is also possible to dispense beneficial agents using different polymers or other binders, wherein the specific polymer or binder has different diffusivity or affinity to assure delivery of the beneficial agents at different rates. Thus, in accordance with the invention, multiple beneficial agents can be released at rates appropriate for their activities, such that the prosthesis-balloon combination of the invention...
has multiple beneficial agents which elute off the prosthesis-balloon combination at desired rates.

For example, a cationic phosphorylcholine-linked polymer which has a higher affinity for anionic beneficial agents can be blended and dispersed as a first beneficial agent and lipophilic phosphorylcholine-linked polymer can be blended with lipophilic drugs as the second beneficial agent to effect different release rates respectively.

In yet another embodiment of the invention, one of the first and second beneficial agents loaded onto the prosthesis-balloon combination may be more hydrophobic than the other. Thus, in accordance with the invention is provided a prosthesis-balloon combination including first and second beneficial agents wherein one of the beneficial agents is more hydrophobic than the other. In this manner, the less hydrophobic beneficial agent is separated from the more hydrophobic beneficial agent, thereby not modifying the release rate of the more hydrophobic beneficial agent. For example and not limitation, the less hydrophobic beneficial agent may be ABT 620 \{1-Methyl-N-(3,4,5-trimethoxyphenyl)-1H-indole-5-sulfonamide\}, which is disclosed in US Patent No. 6,521,658, the disclosure of which is incorporated herein by reference; ABT 627, which is disclosed in US Patent No. 5,767,144, the disclosure of which is incorporated herein by reference; ABT 518 \{[(S - (R*,R*))]-N-[1-(2,2-dimethyl-1,3-dioxol-4-yl)-2-[(4-(trifluoro-methoxy)-phenoxy]phenyl]sulfonyl]ethyl]-N-hydroxyformamide \}, which is disclosed in US Patent No. 6,235,786, the disclosure of which is incorporated herein by reference; dexamethasone, and the like and the more hydrophobic beneficial agent may be Fenofibrate, Tricor™ or the rapamycin analog, ABT-578, i.e.,3S,6R,7E,9R,10R,12R,14S,15E,17E,19E,21S,23S,26R,27R,34aS)-9,10,12,13,14,21,22,23,24,25,26,27,32,33,34,34a-Hexadecahydro-9,27-dihydroxy-3-[(1R)-2-[(1S,3R,4R)-3-methoxy-4-tetrazol-1-yl)cyclohexyl]-1-methylethyl]-10,21-dimethoxy-6,8,12,14,20,26-hexamethyl-23,27-epoxy-3H-pyrido[2,1-c][1,4]oxaazacyclotriacontine-1,5,11,28,29(4H,6H,31H)-pentone; 23,27-Epoxy-3H-pyrido[2,1-c][1,4]oxaazacyclotriacontine-1,5,11,28,29(4H,6H,31H)-pentone, which is disclosed in US Patent No. 6,015,815, US Patent No. 6,329,386, WO 02/123505, and WO 03/129215, disclosures of which are incorporated herein by reference thereto.

Further in accordance with the invention, using the method and systems described above, a first beneficial agent loaded onto the prosthesis can have a first local areal density and a second beneficial agent loaded onto the balloon can have a second local areal density.
As used herein, "areal density" refers to the amount of beneficial agent per unit surface area of a selected portion of the prosthesis or balloon. "Local areal density" refers to the dosage of beneficial agent per local surface area of the prosthesis or balloon. The local areal density of the first beneficial agent and the local areal density of the second beneficial agent can be uniform across each respective portion to define stepped changes in local area density as depicted in Figure 5b or can be varied across a selected portion of the prosthesis or balloon to define gradients of local area density, as depicted in Figure 5c. Accordingly, an interventional device is provided having a prosthesis or balloon that is at least partially loaded with beneficial agent having a local areal density that is varied along a selected portion of the body of the prosthesis or balloon.

In another embodiment of the invention, the local areal density is varied as a continuous gradient along a selected portion of the prosthesis or balloon as shown in Fig. 5c. Accordingly, in one aspect of the invention the local areal density of beneficial agent is varied such as to provide a prosthesis or balloon having a local areal density of beneficial agent at the ends of the prosthesis or balloon that is different than the local areal density of beneficial agent at an intermediate section of the prosthesis or balloon. For purpose of illustration and not limitation, the local areal density of beneficial agent at the intermediate section of the prosthesis can be greater than that at the proximal and distal ends of the prosthesis as shown in Figure 5c. Alternatively, the proximal and distal ends of the prosthesis can have a greater local areal density of beneficial agent than that on the intermediate section of the prosthesis. In a preferred embodiment of the invention, the varied local areal density of beneficial agent corresponds to the location of a lesion when the prosthesis is deployed within a lumen. For example, the prosthesis or balloon can be loaded to have a greater local areal density of beneficial agent along a preselected portion of the prosthesis or balloon that corresponds to the location of the lesion when the prosthesis is deployed in a lumen. Thus, targeted therapy may be achieved with the interventional device of the present invention.

As noted above, the beneficial agent is at least partially loaded onto a surface of the prosthesis. Further in accordance with the invention the prosthesis includes a first surface and a second surface that are at least partially loaded with beneficial agent. In one embodiment of the invention, the first surface and the second surface each correspond to one of the inner surface and the outer surface of the prosthesis. Thus, according to this particular
embodiment, beneficial agent, as defined above, is loaded onto the inner or luminal surface of a prosthesis as well as the outer surface of the prosthesis. In this aspect of the invention, the interventional device can be designed to provide combination therapy of beneficial agents to targeted locations. For example and not limitation, the particular beneficial agent loaded on the balloon can be intended for systemic or down stream release, whereas the particular beneficial agent loaded onto the surface of the prosthesis is intended for release into the wall of the vessel. In accordance with one aspect of the invention, the beneficial agents loaded onto the balloon include, without limitation, antiplatelet agents, aspirin, cell adhesion promoters, agents that promote endothelial recovery, agents that promote migration, estradiol, anti-inflammatories, anti-proliferatives, smooth muscle inhibitors, cell adhesion promoters, and the rapamycin analog ABT-578, i.e.,


9,10,12,13,14,21,22,23,24,25,26,27,32,33,34a-Hexadecahydro-9,27-dihydroxy-3-[(1R)-2-[(1S,3R,4R)-3-methoxy-4-tetrazol-1-yl)cyclohexyl]-1-methylethyl]-10,21-dimethoxy-6,8,12,14,20,26-hexamethyl-23,27-epoxy-3H-pyrido[2,1-c][1,4]oxaazacyclohentriacontine-1,5,11,28,29(4H,6H,31H)-pentone; 23,27-Epoxy-3H-pyrido[2,1-c][1,4]oxaazacyclohentriacontine-1,5,11,28,29(4H,6H,31H)-pentone. The beneficial agents loaded onto the prosthesis include without limitation, antiplatelet agents, aspirin, cell adhesion promoters, agents that promote endothelial recovery, agents that promote migration, estradiol, anti-inflammatories, anti-proliferatives, smooth muscle inhibitors, cell adhesion promoters, angiotensin II receptor antagonists such as losartan, eposartan, valsartan and candesartan, antihypertensive agents such as carvedilol, and the rapamycin analog ABT-578, i.e., 3S,6R,7E,9R,10R,12R,14S,15E,17E,19E,21S,23S,26R,27R,34aS)-

9,10,12,13,14,21,22,23,24,25,26,27,32,33,34a-Hexadecahydro-9,27-dihydroxy-3-[(1R)-2-[(1S,3R,4R)-3-methoxy-4-tetrazol-1-yl)cyclohexyl]-1-methylethyl]-10,21-dimethoxy-6,8,12,14,20,26-hexamethyl-23,27-epoxy-3H-pyrido[2,1-c][1,4]oxaazacyclohentriacontine-1,5,11,28,29(4H,6H,31H)-pentone; 23,27-Epoxy-3H-pyrido[2,1-c][1,4]oxaazacyclohentriacontine-1,5,11,28,29(4H,6H,31H)-pentone. The beneficial agent is loaded onto the prosthesis to provide a controlled local areal density across a length of the interventional device. That is, it may be desirable to provide a greater concentration of beneficial agent at one portion of a prosthesis and a lower concentration, or perhaps no beneficial agent, at another portion of the prosthesis.
For example, in one preferred embodiment, a greater local areal density can be provided at a first portion, e.g., intermediate portion 10b, of a prosthesis or balloon 10, as shown in Fig. 5a, while providing a lower local areal density of beneficial agent to a second portion, e.g., one or both end portions (10a, 10c), of the prosthesis or balloon 10. In accordance with the present invention, each of the first and second portions of the prosthesis or balloon may be defined by any of a variety of patterns or selected portions of the prosthesis or balloon. For example, the first portion of the prosthesis can be defined by longitudinal connectors whereas the second portion of the prosthesis is defined by annular rings, or vice versa.

Alternatively, the beneficial agent distribution profile for the interventional device may be controlled to include any of a variety of desired patterns. For example, the prosthesis or balloon can have a decreased local areal density of beneficial agent on the distal and proximal ends, as noted above. This profile is highly desirable in preventing adverse dosing of beneficial agent if multiple prostheses are placed in combination with each other (for example overlapping prostheses or kissing prostheses at bifurcations) but still provides for decreased dosage of the extreme ends of the interventional device as a whole. Alternatively, as embodied herein, the beneficial agent distribution profile can provide a controlled local areal density that is uniform along the length of a first prosthesis and a second prosthesis in combination, or multiple prostheses in combination. Alternatively, in accordance with the invention, the beneficial agent distribution profile provides a controlled local areal density that is varied along the length of the first prosthesis and the second prosthesis in combination, or multiple prostheses in combination.

Another feature of the present invention includes applying a layer of base material on a selected portion of the prosthesis or balloon described above. The beneficial agent is loaded onto the base material layer according to the methods described above. The base material layer preferably defines a pattern for loading the beneficial agent onto the prosthesis or balloon.

The present invention also encompasses, for any of the embodiments disclosed, the application of a rate-controlling topcoat over the beneficial agent loaded prosthesis, balloon, or prosthesis-balloon combination for further controlling or sustaining the release of beneficial agent. The rate-controlling topcoat may be added by applying a coating layer posited over the beneficial agent loaded prosthesis, balloon, or prosthesis-balloon combination. The thickness of the layer is selected to provide such control. Preferably, the
overcoat is applied by spray coating or fluid-jet technology. Advantageously, fluid jetting an overcoat such as a polymer overcoat allows thinner and more uniform layers. However other conventional methods can be used such as other fluid-dispensers, vapor deposition, plasma deposition, spraying, or dipping, or any other coating technique known in the art.

The present invention also encompasses, for any of the embodiments disclosed, the application of polymer barriers, timing layers, top or capcoats, especially on the luminal side of the prosthesis, or the use of bare metal interfaces to be used to prevent drug transfer from the balloon surface into the delivery polymer of the prosthesis. Alternately, some of the beneficial agent from the balloon could be allowed to transfer to the stent creating a gradient of the two beneficial agents released from the stent into the tissue.

The present invention also provides a method for manufacturing an interventional device for delivery of beneficial agents. This method comprises the steps of providing a prosthesis to be deployed within a lumen; providing a balloon configured to be deployed in an overlapping relationship with the prosthesis, the prosthesis and the balloon in combination defining at least an overlapping segment; and loading the prosthesis with a first beneficial agent and the balloon with a second beneficial agent to provide a controlled local areal density along a length of the prosthesis and the balloon in combination. The method described in detail above is preferred for such loading step.

The present invention also provides a method of delivering beneficial agents. In accordance with this method, as described in detail in conjunction with the description of the interventional device of the present invention above, the method comprising the steps of providing a prosthesis having a tubular body when deployed in a lumen; providing a balloon capable of expanding in the lumen; loading the prosthesis with a first beneficial agent and the balloon with a second beneficial agent; deploying the prosthesis into a lumen with the beneficial agent coated balloon deploying the beneficial agent coated prosthesis into the lumen to define in combination at least one overlapping segment; wherein the beneficial agents are loaded onto the prosthesis and the balloon to provide a controlled local areal density of beneficial agent across a length of the prosthesis when deployed. The method described in detail above is preferred for such loading step.

For purposes of explanation and illustration, and not limitation, an exemplary embodiment of the interventional device in accordance with the invention is shown schematically in Figure 2 and 3. In accordance with one aspect of the invention, as shown
schematically in Figure 2 and 3, the interventional device generally includes a prosthesis loaded with beneficial agent (preferably ABT-578, rapamycin, or rapamycin analogies, alone or in combination with an additional drug such as dexamethasone or estradiol) to provide a local delivery of a first beneficial agent across a treatment zone and a balloon with a second beneficial agent (preferably paclitaxel, taxanes, or other taxane derivatives, alone or in combination with an additional drug) delivered across a second overlapping treatment zone. Alternatively, the prosthesis could be loaded with beneficial agent (preferably paclitaxel, taxanes, or other taxane derivatives alone or in combination with an additional drug such as dexamethasone or estradiol) to provide a local delivery of a first beneficial agent across a treatment zone and a balloon with a second beneficial agent (preferably ABT-578, rapamycin, or rapamycin analogies, alone or in combination with an additional drug) delivered across a second overlapping treatment zone. Particularly, as embodied herein the prosthesis may be a stent, a graft or a stent-graft, as previously noted, for intravascular or coronary delivery and implantation. However, the prosthesis may be any type of implantable member capable of being loaded with beneficial agent. The balloon may be any type of catheter based expandable entity that can act to expand the prosthesis, the local tissue, or push the second beneficial agent against the lumen wall.

The present invention will be further understood by the examples set forth below, which are provided for purpose of illustration and not limitation.

The following examples demonstrate how various embodiments of the present invention may be practiced. By "simultaneous" it is meant that a coated prosthesis (e.g., stent) is mounted on a coated balloon and the stent and balloon are delivered to the desired location at the same time. By "independent", it is meant that the coated balloon is delivered either before or after the coated stent is delivered. By "combined", it is meant that beneficial agent(s) are delivered from both the balloon and the prosthesis to the vessel tissue.

EXAMPLES

Example 1. Loading of stents with beneficial agents or multiple beneficial agents

I. Coating the Stents with PC1036
Prior to any experimentation, coated stents are prepared. These are 3.0 mm X 15 mm 316L electropolished stainless steel stents. Each stent is spray coated using a filtered 20-mg/mL solution of phosphorylcholine polymer PC1036 (product of Biocompatibles Ltd., Farnham, Surrey, UK) in ethanol. The stents are initially air dried and then cured at 70°C for 16 hours. They are then sent for gamma irradiation at <25KGY.

II. Loading the Stents with Drugs of interest

In these experiments, beneficial agents are loaded onto stents and elution profiles examined. In general, the procedure is as follows. Multiple PC-coated stents are loaded with each of several drugs or combinations thereof from solution. The solutions of the drugs are usually in the range of 2 – 20 mg/mL of ABT-578 and 10.0 mg/mL dexamethasone in 100% ethanol, with ~ 10% PC1036 added to the solution to enhance film formation.

The stents are weighed before loading with the drug solution. To load approximately 10 μg/mL of each drug, a solution with equal amounts of ABT-578 and dexamethasone is sprayed onto the stent in a controlled fashion. The stent is allowed to dry before the stents are re-weighted to determine total drug load. The loaded, dry stents are stored in a refrigerator and are protected from light.

III. Extracting Drugs from the Stents

For each drug, 3 stents are used to evaluate the total amount of drug loaded by the above procedure. The stents are immersed in 6 mL of 50% ethanol, 50% water solution and sonicated for 20 minutes. The concentration of the drug in the extraction solution is analyzed by HPLC.

Example 2. Loading of balloons with beneficial agents or multiple beneficial agents

I. Preparing the balloon for drug loading

Multiple balloons (Jomed 15mm X 3.0 mm) are rolled to minimize the final catheter crossing profile. If needed the balloons were washed in ethanol.

II. Loading the balloon with Drugs of interest
In these experiments, beneficial agents are loaded onto balloons. In general, the procedure is as follows. Multiple balloons (Jomed 15mm X 3.0 mm) are loaded with paclitaxel from a solution. The solutions of paclitaxel are usually in the range of 2 – 20 mg/mL of paclitaxel in 100% ethanol.

The balloons are weighed before loading with the drug solution. To load approximately 200 to 600 ug of paclitaxel, the balloons are dipped into a solution of paclitaxel. The balloon is removed in a controlled fashion to control drying. The stent is allowed to dry before the balloons are re-weighed to determine total drug load. The loaded, dry balloons are stored at room temperature and are protected from light.

III. Extracting Drugs from the Balloon

For each drug, 3 balloons are used to evaluate the total amount of drug loaded by the above procedure. The balloons are expanded and immersed in 6 mL of 50% ethanol, 50% water solution and sonicated for 20 minutes. The concentration of the drug in the extraction solution is analyzed by HPLC.

Example 3. Crimping of beneficial agent-coated stents onto beneficial agent-coated balloons.

Multiple stents loaded with ABT-578 and top coated with PC1036 are placed over the end of catheter balloons which have been coated with paclitaxel. The stent is centered over the radiopaque markers of the balloon and crimped onto the balloon using a Machine Solutions drug eluting stent crimper. The stent-ballooon final product is then leak-tested and visually inspected to ensure the quality of the final product. The catheter assembly is then packaged in Tyvek pouches, labeled, and ETO sterilized.

Example 4. Simultaneous combined delivery of a first beneficial agent on prosthesis and a second beneficial agent on Balloon

This example describes delivery of a stent containing at least one beneficial agent using a balloon coated with a second beneficial agent(s). In this example, a prosthesis will be coated with at least one beneficial agent and will be mounted on an angioplasty balloon which has been coated with a second beneficial agent(s). This complete system will be inserted into the body via a peripheral vessel, and advanced to the lesion targeted for treatment. After location at the lesion site, the angioplasty balloon containing the second beneficial agent(s) will be
expanded, simultaneously delivering said beneficial agent(s) as well as deploying the prosthesis containing the first beneficial agent(s). The simultaneous delivery will use a technique often described as direct stenting, in which no pre-dilatation of the vessel at the site of the lesion is involved and the delivery of each beneficial agent begins during the same time period. Alternatively, the simultaneous delivery can be completed after pre-dilatation with an uncoated balloon or with a coated balloon. When deployment of the prosthesis is complete, the balloon will be deflated and removed from the body, leaving the prosthetic device in place to continue delivering the first beneficial agent(s) over time. Beneficial agents on the prosthesis or the balloon can be the same or different.

Example 5. Independent combined delivery of first beneficial agent(s) on prosthesis and second beneficial agent(s) on Balloon

A balloon coated with one or more beneficial agents, but containing no prosthesis, will be inserted into the body, and advanced to the lesion site where it will be dilated to expand the vessel. This technique is commonly described as pre-dilatation. Delivery of a second beneficial agent(s) to the lesion site will proceed upon expansion of this balloon. The balloon will then be deflated and removed from the body. At that time, a second intervention, in which a second balloon without a beneficial agent, containing a prosthesis coated with one or more beneficial agents, will be introduced via the peripheral vessel. Upon expansion of the second balloon at the pre-dilated lesion site, the prosthesis will be expanded and will begin to deliver one or more beneficial agents to the lesion. The second balloon will then be removed from the body.

Example 6. Independent combined delivery of first beneficial agent(s) on prosthesis with a post-expansion delivery of a second beneficial agent(s) from a balloon.

This procedure involves the delivery of a prosthesis containing a first beneficial agent(s), using a balloon that has no beneficial agent. In this case, the balloon catheter, containing a drug-loaded prosthesis, is advanced to the lesion site, and expanded to deliver the device and initiate the delivery of the beneficial agent(s). The balloon is then deflated and removed from the body. At this time, a second balloon, coated with a second beneficial agent(s), is inserted into the peripheral vessel and advanced to the lesion site. A second balloon expansion is then conducted to further expand the previously placed stent or to deliver a second beneficial
agent or agents to the site of the lesion. Beneficial agents on the prosthesis or the balloon can be the same or different.

**Example 7. Delivery of a second beneficial agent on balloon to treat in-stent restenosis.**

This intervention involves the dilation of a vessel with a balloon that is coated with a second beneficial agent(s) at a restenosed lesion site where a stent or stents have been previously placed. In this way, restenosis of a vessel in which an intervention has previously failed can be adequately treated without placement of an additional prosthesis or prosthesis at the same site.

As will be recognized by those of ordinary skill, the examples can be adapted to address situations for which it is desired to deliver multiple stents, e.g., "kissing" stents or overlapping stents.
The claims defining the invention are as follows:

1. A system for delivering a beneficial agent, comprising:
   a) a balloon having an outer surface that is at least partially coated with a therapeutically effective amount of a first beneficial agent; and
   b) a prosthesis that can be delivered or implanted having a surface that is at least partially coated with a therapeutically effective amount of a second beneficial agent wherein the system is useful for the treatment and prevention of vascular diseases.

2. The system according to claim 1, wherein the balloon is an angioplasty balloon.

3. The system according to claim 1, wherein the prosthesis is a stent.

4. The system according to claim 3, wherein said prosthesis is crimped onto said balloon.

5. The system according to claim 1, wherein said prosthesis is at least partially retained on said balloon by said coating on said balloon.

6. The system according to claim 3, wherein the stent is a self-expanding stent.

7. The system according to claim 1, wherein the coating on the balloon further comprises a carrier for said first beneficial agent.

8. The system according to claim 7, wherein the carrier is selected from the group consisting of complex sugars (mannitol) starches (cellulose), collagens, and polymeric materials.

9. The system according to claim 1, wherein the coating on the prosthesis further comprises a carrier for said second beneficial agent.

10. The system according to claim 9, wherein said carrier for said second beneficial agent is selected from a group consisting of phosphorylcholine, polycaprolactone, poly-D,L-lactic acid, poly-L-lactic acid, poly(lactide-co-glycolide), poly(hydroxybutyrate), poly(hydroxybutyrate-co-valerate), polydioxanone, polyorthoester, polyanhydride, poly(glycolic acid), poly(glycolic acid-co-trimethylene carbonate), polyphosphoester, polyphosphoester urethane, poly(amo acid), cyanoacrylates, poly(trimethylene carbonate), poly(iminocarbonate), polyalkylene oxalates, polyphosphazenes, polyiminocarbonates, and aliphatic polycarbonates, fibrin, fibrinogen, cellulose, starch, collagen, one or more of a series of polymers synthesized by the pyrolitic, vapor-decomposition polymerization of p-xylylene dimers, or a plasma enhanced coating thereof, polyurethane, polycarbonate urethanes, polyethylene,
polyethylene terephthalate, ethylene vinyl acetate, ethylene vinyl alcohol, silicone polysiloxanes, substituted polysiloxanes, polyethylene oxide, polybutylene terephthalate-co-PEG, PCL-co-PEG, PLA-co-PEG, polyacrylates, polyvinyl pyrrolidone, polyacrylamide, thermoplastic elastomers, polyolefin elastomers, EPDM rubbers, polyamide elastomers, biostable plastic, acrylic polymers, nylon, polyesters, epoxies and derivatives or combinations thereof.

11. The system according to claim 1, further comprising a balloon which is substantially free of a beneficial agent.

12. The system according to claim 1, wherein the first beneficial agent is the same as the second beneficial agent.

13. The system according to claim 1, wherein a surface of said prosthesis is loaded with therapeutically effective amounts of more than one beneficial agent.

14. The system according to claim 1, wherein the first beneficial agent and the second beneficial agent are individually selected from the group consisting of antithrombotics, anticoagulants, antiplatelet agents, anti-lipid agents, thrombolitics, antiproliferatives, antiinflammatories, agents that inhibit hyperplasia, smooth muscle cell inhibitors, antibiotics, growth factor inhibitors, cell adhesion inhibitors, cell adhesion promoters, antimitotics, antifibrins, antioxidants, antineoplastics, agents that promote endothelial cell recovery, antiallergic substances, viral vectors, nucleic acids, monoclonal antibodies, antisense compounds, oligonucleotides, cell permeation enhancers, radiopaque agent markers, HMG CoA reductase inhibitors, indomethacin, phenyl salicylate, vinblastine, ABT-627 (atrasentan), ABT-578, testosterone, progesterone, paclitaxel, taxanes, cyclosporin A, vincristine, carvedilol, vindesine, dipryridamole, methotrexate, folic acid, thrombospondin mimetics, estradiol, dexamethasone, metrizamide, iopamidol, iohexol, iopromide, iobitridol, iomeprol, iopentol, ioversol, ioxilan, iodoxanol, iotrolan, rapamycin, rapamycin derivatives, pimecrolimus, everolimus, fenofibrate, carvedilol, taxotere, tacrolimus, peroxisomal proliferator activated receptor γ (PPAR-γ) agonists, paricalcitol, butylated hydroxytoluene, butylated hydroxyanisole, vitamin E, danazol, probucol, tocopherols, tocotrienols, colchicine, betamethasone, cortisone, budesonide, prednisolone, methylprednisolone, hydrocortisone, flurbiprofen, ibuprofen, ketoprofen, fenoprofen, naproxen, diclofenac, diflunisal, acetaminophen, sulindac, etodolac, ketorolac, meclofenamic acid, piroxicam, phenylbutazone, altretamine, bendamucine, carboptatin, carmustine, cisplatin, cyclophosphamide, fotemustine, ifosfamide, lomustine, nimustine, prednimustine, treosulfin, docetaxel, mercaptopurine, pentostatin, trimetrexate,
gemcitabine, azathioprine, fluorouracil, doxorubicin hydrochloride, mitomycin, RPR-101511A, rosiglitazone, pioglitazone, lovastatin, atorvastatin, simvastatin, pravastatin, cerivastatin, fluvastatin, ABT-518, permirolast, potassium nitroprusside, phosphodiesterase inhibitors, prostaglandin inhibitors, suramin, serotonin blockers, steroids, thiopeptase inhibitors, triazolopyrimidine, nitric oxide, dalteparin, enoxaparin, nadroparin, reviparin, aradoparin, certaparin, heparinoids, hirudin, argatroban, forskolin, vapriprost, prostacyclin, dextran, D-phe-pro-5-arg-chloromethylketonedipyridamole, glycoprotein IIb/IIIa, recombinant hirudin, thrombin inhibitors, urokinase, recombinant urokinase, pro-urokinase, tissue plasminogen activator, tenecteplase, angiopeptin, captopril, cilazapril, lisinopril, nifedipine, amlodipine, cilnidipine, lercanidipine, benidipine, trifluperazine, diltiazem, verapamil, fibroblast growth factor antagonists, fish oil (omega 3-fatty acid), histamine antagonists, etoposide, topotecan, tamoxifen and combinations thereof.

15. The system according to claim 14, wherein said HMG CoA reductase inhibitor is selected from the group consisting of atorvastatin, simvastatin, pravastatin, lovastatin, cerivastatin, fluvastatin.

16. The system according to claim 14, wherein the PPAR-γ agonist is a thiazolidinedione.

17. The system according to claim 1, wherein the beneficial agent is a nucleic acid that encodes a pharmaceutically useful peptide or an anti-sense oligonucleotide used to control a gene of interest in a cell of the patient.

18. The system according to claim 1, wherein the first beneficial agent has a varied local areal density as a measure of beneficial agent per unit local surface area of the balloon surface.

19. The system according to claim 1, wherein the coating on the balloon is discontinuous or forms a pattern.

20. The system according to claim 4, wherein said coating on the balloon extends beyond said prosthesis crimped on said balloon.

21. The system according to claim 6, wherein said coated balloon is delivered after said stent is implanted to accomplish final sizing of said stent.

22. The system according to claim 1, wherein said first beneficial agent is transferred from said coating on said balloon into said coating on said prosthesis.
23. The system according to claim 1, wherein said coating on said prosthesis comprises a barrier to transfer of said first beneficial agent from said coating on said balloon into said coating on said prosthesis or of said second beneficial agent from said coating on said prosthesis into said coating on said balloon.

24. The system according to claim 23, wherein said barrier is polymeric.

25. The system according to claim 24, wherein the polymeric component is selected from a group consisting of phosphorylcholine, polycaprolactone, poly-D,L-lactic acid, poly-L-lactic acid, poly(lactide-co-glycolide), poly(hydroxybutyrate), poly(hydroxybutyrate-co-valerate), polydioxanone, polyorthoester, polyanhydride, poly(glycolic acid), poly(glycolic acid-co-trimethylene carbonate), polyphosphoester, polyphosphoester urethane, poly(amine acids), cyanoacrylates, poly(trimethylene carbonate), poly(iminocarbonate), polyalkylene oxalates, polyphosphazenes, polyiminocarbonates, and aliphatic polycarbonates, fibrin, fibrinogen, cellulose, starch, collagen, one or more of a series of polymers synthesized by the pyrolytic, vapor-decomposition polymerization of p-xylylene dimers, or a plasma enhanced coating thereof, polyurethane, polycarbonate urethanes, polyethylene, polyethylene terephthalate, ethylene vinyl acetate, ethylene vinyl alcohol, silicone polysiloxanes, substituted polysiloxanes, polyethylene oxide, polybutylene terephthalate-co-PEG, PCL-co-PEG, PLA-co-PEG, polyacrylates, polyvinyl pyrrolidone, polyacrylamide, thermoplastic elastomers, polyolefin elastomers, EPDM rubbers, polyamide elastomers, biostable plastic, acrylic polymers, nylon, polyesters, epoxies and derivatives or combinations thereof.

26. The method of treating and preventing a vascular disease using the system of claim 1.

27. The system according to claim 22, wherein said first beneficial agent is delivered from said coating on said prosthesis into a target site.

28. The system according to claim 1, wherein at least one of said coating on said balloon and said coating on said prosthesis contains multiple beneficial agents.

29. A method of treating and preventing a vascular disease, comprising the steps of:
   a) delivering a balloon having an outer surface that is at least partially coated with a therapeutically effective amount of a first beneficial agent; and
b) delivering a prosthesis having a surface that is at least partially coated with a therapeutically effective amount of a second beneficial agent.

30. The method according to claim 29, wherein the first beneficial agent is not the same as the second beneficial agent.

31. The method according to claim 29, wherein step a and step b are performed simultaneously.

32. The method according to claim 29, wherein said balloon is adhered to said prosthesis by said coating on said balloon.

33. The method according to claim 29, wherein step a and step b are performed in sequence.

34. The method according to claim 29, further comprising a step of delivering an uncoated balloon.

35. The method according to claim 29, wherein said prosthesis is a self-expanding stent.

36. The method according to claim 35, wherein said coated balloon is used for final sizing of said coated stent.

37. A method of providing a device for treatment and prevention of vascular disease, comprising the steps of:

- providing a beneficial agent; and
- coating said beneficial agent onto a balloon, wherein the balloon includes a balloon surface having a working length, the working length of the balloon surface is at least partially coated with the beneficial agent, the beneficial agent having a varied local areal density as a measure of beneficial agent per unit local surface area on the working length of the balloon surface.

38. The method according to claim 37, wherein said coating step is accomplished by dipping, spraying, layering, painting or jetting.

39. The method according to claim 37, wherein said beneficial agent is coated from a solution.

40. The method according to claim 39, wherein said solution comprises at least one solvent.
41. The method according to claim 40, wherein said at least one solvent is selected from the group consisting of water, alcohols, dichloromethane, dimethylsulfoxide, ethanol, isopropyl alcohol, isobutyl alcohol, tetrahydrofuran, alkanes, dimethylformamide and mixtures thereof.

42. The method according to claim 37, further comprising a step of providing a prosthesis for use in association with said coated balloon.

43. The method according to claim 42, further comprising a step of enhancing retention of said prosthesis by mounting said prosthesis on said coated balloon prior to complete drying or curing of said coating on said balloon.

44. A method for treating vascular disease in a bifurcated vessel, comprising:

- deploying a balloon within a first branch of a bifurcated vessel, wherein the balloon includes a balloon surface having a working length, the working length of the balloon surface is at least partially coated with at least one beneficial agent, the at least one beneficial agent having a varied local areal density as a measure of beneficial agent per unit local surface area on the working length of the balloon surface; and
- delivering a therapeutically effective amount of the at least one beneficial agent to the first branch of the bifurcated vessel.

45. The method according to claim 44, further comprising: deploying a first stent device within the first branch of the bifurcated vessel.

46. A system for delivering a beneficial agent, substantially as hereinbefore described with reference to any one of the embodiments as that embodiment is shown in the accompanying drawings.

47. A method of treating and preventing a vascular disease, substantially as hereinbefore described with reference to any one of the embodiments as that embodiment is shown in the accompanying drawings.

48. A method of providing a device for treatment and prevention of vascular disease, substantially as hereinbefore described with reference to any one of the embodiments as that embodiment is shown in the accompanying drawings.