DELIVERY OF THERAPEUTIC COMPOSITIONS USING ULTRASOUND

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Related U.S. Application Data
Continuation of application No. 09/620,701, filed on Jul. 20, 2000, now Pat. No. 6,527,759, which is a division of application No. 09/158,316, filed on Sep. 21, 1998, now Pat. No. 6,176,842, which is a continuation-in-part of application No. 09/129,980, filed on Aug. 5, 1998, now Pat. No. 6,210,356, which is a continuation-in-part of application No. 08/972,846, filed on Nov. 18, 1997, now abandoned, which is a continuation of application No. 08/611,105, filed on Mar. 5, 1996, now abandoned.

Therapeutic compositions are delivered to a target site using a catheter which includes at least one ultrasound transducer coupled to an energy source. The therapeutic compositions include genetic material and the target site may be a DNA with affinity for the genetic material.
FIG. 17A

PORPHYRINA:
NATURALLY OCCURRING:

PROTOPORPHYRIA IX

SYNTHETIC:

FIG. 17C

CHLORINA
NATURALLY OCCURRING:

FIG. 17D

BPD MONO ACID RING A

FIG. 17B

VERDIA
FIG. 17K
NAPHTHALOCYANINE

FIG. 17L
PORPHYRINS:

FIG. 17M
SAPPYRRINS:

FIG. 17N
TEXAPHYRINAS:
FIG. 18E

OR

FIG. 18F
FIG. 20A

BPD - MA

AND

FIG. 20B

BPD - MB
FIG. 20C

BPD-DA

FIG. 20D

BPD-DB
Fig. 21A
FIG. 21B
HYDRAZINE HYDRATE, 10% Pd/C, EtOH

1.2 eq 12M HCl, MeOH
FIG. 31F

1. NHS, DCC, DMF

2. 5'-NH$_2$(CH$_2$)$_6$PO$_4$CTC GGC CAT AGC GAA TGT TC-3
   (0.4 M NaHCO$_3$, pH 8.6) SEQ. ID. NO. 1

FIG. 31G

FIG. 31H
FIG. 34
DELIVERY OF THERAPEUTIC COMPOSITIONS USING ULTRASOUND

RELATED APPLICATIONS

[0001] This application is a Continuation of U.S. application Ser. No. 09/620,701, filed Jul. 20, 2000 which is a Divisional of U.S. application Ser. No. 09/158,316, filed Sep. 21, 1998, now U.S. Pat. No. 6,176,842 which is a Continuation-in-Part of U.S. application Ser. No. 09/129,980; filed Aug. 5, 1998, and a Continuation-In-Part of Japanese application number HO9-255814, filed Sep. 19, 1997, entitled Drug Carrier and Method of Using Same and a Continuation-In-Part of U.S. application Ser. No. 08/972,846, filed Nov. 18, 1997, now abandoned, which is a Continuation of U.S. application Ser. No. 08/611,105, filed Mar. 5, 1996, now abandoned, which claims priority to Japanese application number 097-048710; filed Mar. 8, 1995. Each of the above applications are incorporated herein by reference in their entirety.

BACKGROUND OF THE INVENTION

[0002] 1. Field of the Invention

[0003] The present invention relates to a method that utilizes ultrasonic vibration to perform various types of therapies, and particularly to a therapy-accelerating substance that controls the administration of a drug by utilizing ultrasonic vibration at specific locations within the body to perform the release of the drug effectively. The drug may include a nucleic acid material such as a DNA, RNA, or one or more oligonucleotides.

[0004] 2. Description of the Related Art

[0005] Current methods of treating cancer can be broadly divided into two methods: the removal of the cancerous tissue by surgery or the killing of the cancerous cells with anti-cancer drugs. The removal of cancerous tissue can only be performed in cases in which the cancer occurs in limited small areas and no metastasis is found. On the other hand, cancer chemotherapy has extremely strong side effects such as nausea, impairment of kidney function and impairment of liver function which happens often with the administration of large doses of drugs. Moreover, cancer reacts only to drugs in high concentrations, so cancer chemotherapy has not demonstrated very good clinical results.

[0006] Many different approaches to solving this problem are currently being attempted. The method called “missile therapy” (a term used by Japanese physicians to refer to types of cancer therapy where a payload of drug or radioactive isotope is delivered accurately to the site of a cancer in the manner of a military “smart bomb” or missile) using anti-cancer drugs is a method that uses anti-cancer drugs bound to antibodies that selectively bind with the cancerous cells, so that the anti-cancer drugs act in a concentrated manner on the cancerous cells, but this method has not yet achieved adequate results.

[0007] On the other hand, a method has been proposed in which anti-cancer drugs enclosed in capsules or other drug carriers made of specific materials are injected into the body and the shells are made to rupture within blood vessels near the cancer so that a high concentration of anti-cancer drugs are administered to limited areas. Its effectiveness has been demonstrated experimentally.

[0008] However, methods of efficiently rupturing the capsule shells used in this method have not been established. Up until now, research has been conducted on embedding polymer-based temperature sensors, pH sensors and the like in the capsule shells so that the release of drugs is induced under certain temperature or pH conditions, but arbitrarily setting the temperature and pH conditions near the location of the tumor is extremely difficult.

[0009] In addition, methods have been proposed by which the drugs are released from the capsules when the capsule shells are ruptured by shock waves or ultrasonic energy applied from outside. For example, the specification of U.S. Pat. No. 5,580,575 recites a method wherein liposomes containing a gas and drugs are ruptured by ultrasound at a specific location in the body of the patient. However, rupturing a liposome or other capsule mechanically with the vibration of ultrasound alone in this manner requires irradiation with high-intensity ultrasound, and the resonance frequency is determined depending on the amount of gas within the capsule, so it is difficult to rupture the capsule with ultrasound using frequencies other than the resonance frequency.

[0010] In this manner, the use of acoustical energy requires considerable precision in the setting of the irradiation conditions, so it has not been easy to achieve the precise ultrasound frequency and intensity required in the location of the tumor.

[0011] The problem addressed by the present invention is to provide a drug carrier and method of using same that is able to achieve the delivery of high concentrations of anti-cancer drugs or other drugs to specific locations by means of capsules or other drug carriers carrying said anti-cancer drugs or other drugs that are reliably and simply ruptured using ultrasound at specific locations such as in blood vessels within cancerous tissue or on the surface of the skin. In order to solve the aforementioned problem, a drug carrier contains an ultrasound-sensitive substance.

[0012] It is frequently desirable to kill target biological tissues such as tumors and atheroma. One technique for causing targeted tissue death is called photodynamic therapy which requires the use of light activated drugs. Light activated drugs are inactive until exposed to light of particular wavelengths, however, upon exposure to light of the appropriate wavelength, light activated drugs can exhibit a cytotoxic effect on the tissues where they are localized. It has been postulated that the cytotoxic effect is a result of the formation of singlet oxygen on exposure to light.

[0013] Photodynamic therapy begins with the systemic administration of a selected light activated drug to a patient. At first, the drug disperses throughout the body and is taken up by most tissues within the body. After a period of time usually between 3 and 48 hours, the drug clears from most normal tissue and is retained to a greater degree in lipid rich regions such as the liver, kidney, tumor and atheroma. A light source, such as a fiber optic, is then directed to a targeted tissue site which includes the light activated drug. The tissues of the tissue site are then exposed to light from the light source in order to activate any light activated drugs within the tissue site. The activation of the light activated drug causes tissue death within the tissue site.

[0014] Several difficulties can be encountered during photodynamic therapy. For instance, since the light activated
drug is typically administered systemically, the concentration of the light activated drug within the targeted tissue site is limited by the quantity of light activated drug administered. The concentration of the light activated drug within a tissue site can also be limited by the degree of selective uptake of the light activated drug into the tissue site. Specifically, if the targeted tissue site does not selectively uptake the light activated drug, the concentration of light activated drug within the tissue site can be too low for an effective treatment.

[0015] An additional problem associated with photodynamic therapy concerns depth of treatment. Light cannot penetrate deeply into opaque tissues. As a result, the depth that light penetrates most tissue sites is limited. This limited depth can prevent photodynamic therapy from being used to treat tissues which are located deeply in the interior of a tissue site.

[0016] There is currently a need for a method and apparatus which can be used to cause death to tissues death deep within a tissue site. When the method and apparatus employ light activated drugs, the method and apparatus should be able to provide an appropriate concentration of light activated drug within the tissue site.

**SUMMARY OF THE INVENTION**

[0017] An object for an embodiment of the invention is causing tissue death within a tissue site.

[0018] Another object for an embodiment of the present invention is locally delivering a light activated drug to a tissue site and activating the light activated drug.

[0019] Yet another object for an embodiment of the present invention is locally delivering a light activated drug to a tissue site and delivering ultrasound energy to the delivered light activated drug to activate the light activated drug.

[0020] A further object for an embodiment of the present invention is using a catheter to locally deliver a light activated drug to a tissue site and delivering ultrasound energy from an ultrasound element on the catheter to activate the light activated drug.

[0021] Yet a further object for an embodiment of the present invention is including the light activated drug in an emulsion, locally delivering the emulsion to a tissue site and delivering ultrasound energy to the light activated drug within the tissue site to activate the light activated drug.

[0022] Even a further object for an embodiment of the present invention is including the light activated drug in a liposome, locally delivering the liposome to a tissue site and delivering ultrasound energy to the light activated drug within the tissue site to activate the light activated drug.

[0023] An additional object for an embodiment of the present invention is including the light activated drug in an aqueous solution, locally delivering the aqueous solution to a tissue site and delivering ultrasound energy to the light activated drug within the tissue site to activate the light activated drug.

[0024] Yet a further object for an embodiment of the present invention is including the light activated drug in an emulsion, systemically delivering the emulsion, providing the light activated drug sufficient time to localize within a tissue site and delivering ultrasound energy to the light activated drug within the tissue site to activate the light activated drug.

[0025] Even a further object for an embodiment of the present invention is including the light activated drug in liposomes, systemically delivering the liposomes, providing the light activated drug sufficient time to localize within a tissue site and delivering ultrasound energy to the light activated drug within the tissue site to activate the light activated drug.

[0026] An additional object for an embodiment of the present invention is including the light activated drug in an aqueous solution, systemically delivering the aqueous solution, providing the light activated drug sufficient time to localize within a tissue site and delivering ultrasound energy to the light activated drug within the tissue site to activate the light activated drug.

[0027] Yet another object for an embodiment of the present invention is coupling a site directing molecule to a light activated drug, locally delivering the light activated drug to a tissue site and activating the light activated drug within the tissue site.

[0028] Yet another object for an embodiment of the invention is providing a catheter for locally delivering a media including a light activated drug to a tissue site. The catheter including an ultrasound assembly configured to activate the light activated drug within the tissue site.

[0029] A further object for an embodiment of the invention is providing a catheter for delivering a media including a light activated drug to a tissue site. The catheter including an ultrasound assembly for reducing exposure of the light activated drug to ultrasound energy until the light activated drug has been delivered from within the catheter.

[0030] A kit for causing tissue death within a tissue site is disclosed. The kit includes a media with a light activated drug activatable upon exposure to a particular level of ultrasound energy. The kit also includes a catheter with a lumen coupled with a media delivery port through which the light activated drug can be locally delivered to the tissue site. The ultrasound transducer is configured to transmit the level of ultrasound energy which activates the light activated drug with sufficient power that the ultrasound energy can penetrate the tissue site.

[0031] A method for causing tissue death in a subdermal tissue site is also disclosed. The method includes providing a catheter for locally delivering a light activated drug to the subdermal tissue site, the catheter including an ultrasound transducer. The method also includes locally delivering the light activated drug to the tissue site; producing ultrasound energy from the ultrasound transducer; and directing the ultrasound energy to the subdermal tissue site following penetration of the light activated drug into the subdermal tissue site to activate at least a portion of the light activated drug within the subdermal tissue site.

[0032] A method for activating a light activated drug is also disclosed. The method includes providing a catheter with an ultrasound transducer. The method also includes introducing the light activated drug into a patient's body where a subdermal tissue site absorbs at least a portion of the
light activated drug; producing ultrasound energy; directing the ultrasound energy to the light activated containing subdermal tissue site including the light activated drug; and activating at least a portion of the light activated drug in the subdermal selected tissue site.

[0033] A method for releasing a therapeutic from a microbubble is also disclosed. The method includes providing a microbubble with a light activated drug activatable upon exposure to ultrasound energy; and delivering ultrasound energy to the microbubble at a frequency and intensity which activates the light activated drug to cause a rupture of the microbubble.

[0034] A microbubble is also disclosed. The microbubble includes a substrate defining a shell of the microbubble and having a thickness permitting hydraulic transport of the microbubble. The microbubble also includes a light activated drug activatable upon exposure to ultrasound energy. Activation of the light activated drug causes a disruption in the shell sufficient to cause a rupture of the microbubble. The microbubble further includes a therapeutic releasable from the microbubble upon rupture of the microbubble and yielding a therapeutic effect upon release from the microbubble.

**BRIEF DESCRIPTION OF THE DRAWINGS**

[0035] FIG. 1A is a side view of a catheter for locally delivering a media including a light activated drug to a tissue site.

[0036] FIG. 1B is an axial cross section of an ultrasound assembly for use with the catheter shown in FIG. 1A.

[0037] FIG. 1C is a lateral cross section of an ultrasound assembly for use with the catheter shown in FIG. 1A.

[0038] FIG. 2A is a side view of a catheter having an elongated body and an ultrasound assembly which is flush with the elongated body.

[0039] FIG. 2B is an axial cross section of the ultrasound assembly illustrated in FIG. 2A.

[0040] FIG. 2C is a lateral cross section of the ultrasound assembly illustrated in FIG. 2A.

[0041] FIG. 3A illustrates a catheter with a utility lumen and a second utility lumen.

[0042] FIG. 3B is an axial cross section of the ultrasound assembly illustrated in the catheter of FIG. 3A.

[0043] FIG. 4A is a side view of a catheter including a plurality of ultrasound assemblies.

[0044] FIG. 4B is a cross section of an ultrasound assembly included on a catheter with a plurality of utility lumens.

[0045] FIG. 4C is a cross section of an ultrasound assembly included on a catheter with a plurality of utility lumens.

[0046] FIG. 5A is a side view of a catheter including a balloon.

[0047] FIG. 5B is a cross section of a catheter with a balloon which include an ultrasound assembly.

[0048] FIG. 6A is a side view of a catheter with a balloon positioned distally relative to an ultrasound assembly.

[0049] FIG. 6B is a side view of a catheter with an ultrasound assembly positioned distally relative to a balloon.

[0050] FIG. 6C is a cross section of a catheter with an ultrasound assembly positioned at the distal end of the catheter.

[0051] FIG. 7A is a side view of a catheter with a media delivery port positioned between an ultrasound assembly and a balloon.

[0052] FIG. 7B is a side view of a catheter with an ultrasound assembly positioned between a media delivery port and a balloon.

[0053] FIG. 7C is a cross section of a catheter with an ultrasound assembly positioned at the distal end of the catheter.

[0054] FIG. 8A is a side view of a catheter including a media delivery port and an ultrasound assembly positioned between first and second balloons.

[0055] FIG. 8B is a side view of a catheter including a media delivery port and an ultrasound assembly positioned between first and second balloons.

[0056] FIG. 8C is a cross section of a balloon included on a catheter having a first and second balloon.

[0057] FIG. 9A illustrates an ultrasound assembly positioned adjacent to a tissue site and microbubbles delivered via a utility lumen.

[0058] FIG. 9B illustrates an ultrasound assembly positioned adjacent to a tissue site and a media delivered via a media delivery port.

[0059] FIG. 9C illustrates an ultrasound assembly positioned adjacent to a tissue site and a media delivered via a media delivery port while a guidewire is positioned in a utility lumen.

[0060] FIG. 9D illustrates a catheter including a balloon positioned adjacent to a tissue site.

[0061] FIG. 9E illustrates a catheter including a balloon expanded into contact with a tissue site.

[0062] FIG. 9F illustrates a catheter with an ultrasound assembly outside a balloon positioned at a tissue site.

[0063] FIG. 9G illustrates the balloon of FIG. 9F expanded into contact with a vessel so as to occlude the vessel.

[0064] FIG. 9H illustrates a catheter with an ultrasound assembly outside a first and second balloon positioned at a tissue site.

[0065] FIG. 9I illustrates the first and second balloon of FIG. 9H expanded into contact with a vessel so as to occlude the vessel.

[0066] FIG. 10A is a cross section of an ultrasound assembly according to the present invention.

[0067] FIG. 10B is a cross section of an ultrasound assembly according to the present invention.

[0068] FIG. 10C illustrates a support member with integral supports.

[0069] FIG. 10D illustrates a support member which is supported by an outer coating.
FIG. 1A is a cross section of an ultrasound assembly including two concentric ultrasound transducers in contact with one another.

FIG. 1B is a cross section of an ultrasound assembly including two separated and concentric ultrasound transducers.

FIG. 1C is a cross section of an ultrasound assembly including two ultrasound transducers where a chamber is defined between one of the ultrasound transducers and an elongated body.

FIG. 1D is a cross section of an ultrasound assembly including two longitudinally adjacent ultrasound transducers in physical contact with one another.

FIG. 1E is a cross section of an ultrasound assembly including two separated and longitudinally adjacent ultrasound transducers.

FIG. 1F is a cross section of an ultrasound assembly including two longitudinally adjacent ultrasound transducers with a single chamber positioned between both ultrasound transducers and an elongated body.

FIG. 1G is a cross section of an ultrasound assembly including two longitudinally adjacent ultrasound transducers with different chambers positioned between each ultrasound transducer and an elongated body.

FIG. 1H is a cross section of an ultrasound assembly including two longitudinally adjacent ultrasound transducers in contact with one another and having a single chamber positioned between each ultrasound transducer and an elongated body.

FIG. 12A is a cross section of a catheter which includes an ultrasound assembly module which is independent of a first catheter component and a second catheter component.

FIG. 12B illustrates the first and second catheter components coupled with the ultrasound assembly module.

FIG. 12C is a cross section of an ultrasound assembly which is integral with a catheter.

FIG. 13A is a cross section of an ultrasound assembly configured to radiate ultrasound energy in a radial direction. The lines which drive the ultrasound transducer pass through a utility lumen in the catheter.

FIG. 13B is a cross section of an ultrasound assembly configured to radiate ultrasound energy in a radial direction. The lines which drive the ultrasound transducer pass through line lumens in the catheter.

FIG. 13C is a cross section of an ultrasound assembly configured to longitudinally radiate ultrasound energy. The distal portion of one line travels proximally through the outer coating.

FIG. 13D is a cross section of an ultrasound assembly configured to longitudinally transmit ultrasound energy. The distal portion of one line travels proximally through a line lumen in the catheter.

FIG. 14A illustrates ultrasound transducers connected in parallel.

FIG. 14B illustrates ultrasound transducers connected in series.

FIG. 14C illustrates ultrasound transducers connected with a common line.

FIG. 15 illustrates a circuit for electrically coupling temperature sensors.

FIG. 16 illustrates a feedback control system for use with a catheter including an ultrasound assembly.

FIGS. 17A-N illustrate pyrrole-based macrocyclic classes of light emitting drugs.

FIG. 17B-2 illustrates possible texaphyrin derivation sites.

FIGS. 18A-F illustrate the formula of preferred light emitting drugs for use with media including microbubbles.

FIG. 19 illustrates a formula for a porphyrin group.

FIGS. 20A-D illustrate the formula of four preferred forms of the hydro-monobenzoporphyrin derivatives of the green porphyrins illustrated in formulae 3 and 4 of FIG. 18.

FIGS. 21A-B illustrate the formulae for specific examples of pyrrole-based macrocycle derivatives and xanthene derivatives which are preferred for inclusion in microbubbles to enhance rupture of the microbubbles upon activation.

FIG. 22 shows a sectional structural drawing of one embodiment of a drug carrier.

FIG. 23 shows a sectional structural drawing of another embodiment of a drug carrier.

FIG. 24 shows a sectional structural drawing of still another embodiment of a drug carrier.

FIG. 25 shows a sectional drawing showing the mounting of the ultrasound generator used in the preferred embodiment of the present invention.

FIG. 26 shows a sectional drawing showing one embodiment of a therapeutic ultrasound generator.

FIG. 27 shows an enlarged sectional drawing used to explain the case in which the drug carriers of the present invention are applied to thrombolytic therapy.

FIG. 28 shows an enlarged sectional drawing used to explain the case in which the drug carriers of the present invention are applied to blood vessel therapy.

FIG. 29 shows a sectional drawing of a transdermal administration apparatus that applies the drug carriers of the present invention.

FIGS. 30A-I schematically summarize the synthesis of an oligonucleotide conjugate of a texaphyrin metal complex.

FIGS. 31A-I illustrate the covalent coupling of texaphyrin metal complexes with amine, thiol, or hydroxy linked oligonucleotides.

FIGS. 32A-F illustrate the synthesis of diformyl monoic acid and oligonucleotide conjugate.
FIGS. 33A-J illustrate the synthesis of a texaphyrin based light activated drug.

FIG. 34 illustrates the formula for tin ethyl etiopurpurin (SnEt$_2$).

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

The present invention relates to a method and catheter for delivering a light activated drug to a tissue site and delivering ultrasound energy to the light activated drug within the tissue site. Since many light activated drugs are also activated by ultrasound energy, the delivery of ultrasound energy to the light activated drug activates the light activated drug within the tissue site. Similar to activation of a light activated drug by light, activation by ultrasound causes death of tissues within the tissue site. The tissue death is believed to result from the release of a singlet oxygen. Suitable tissue sites include, but are not limited to, atheroma, cancerous tumors, thrombi and potential restenosis sites. A potential restenosis site is a tissue site where restenosis is likely to occur such as the portion of vessels previously treated by balloon angioplasty. In contrast to light, ultrasound energy can be transmitted through opaque tissues. As a result, the ultrasound energy can be used to treat tissues which are deeper within a tissue site than could be treated via light activation.


Cavitation results when gas dissolved in a solution forms bubbles under certain types of acoustic vibration. Cavitation can also occur when small bubbles already present in the solution oscillate or repeatedly enlarge and contract to become bubbles. When the size of these cavitation bubbles reaches a size that cannot be maintained, they suddenly collapse and release various types of energy. The various types of energy include, but are not limited to, mechanical energy, visible light, ultraviolet light and other types of electromagnetic radiation. Heat, plasma, magnetic fields, shock waves, free radicals, heat and other forms of energy are also thought to be generated locally. The light activated drug is believed to be activated by at least one of the various forms of energy generated at the time of cavitation collapse.

The delivery of light activated drug to the tissue site can be through traditional systemic administration of a media including the light activated drug or can be performed through localized delivery of the media. Localized delivery can be achieved through injection into the tissue site or through other traditional localized delivery techniques. A preferred delivery technique is using a catheter which includes a media delivery lumen coupled with a media delivery port. The catheter can be positioned such that the media delivery port is within the tissue site or is adjacent to the tissue site via traditional over-the-guidewire techniques. The media can then be locally delivered to the tissue site through the media delivery port.

The localized delivery of the light activated drug to the tissue site serves to localize the light activated drug within the tissue site and can reduce the amount of light activated drug which concentrates in tissues outside the tissue site. Further, localized delivery of the light activated drug can serve to increase the concentration of the light activated drug within the tissue site above levels which would be achieved through systemic delivery of the light activated drug. Alternatively, the same concentration of light activated drug within the tissue site as would occur through systemic administration can be achieved by introducing smaller amounts of light activated drug into a patient’s body.

Localized delivery of the light activated drug also permits treatment of tissue sites which do not have selective uptake of the light activated drug. As discussed above, many light activated drugs, such as the texaphyrins, are taken up by most tissues within the body and later localize within lipid rich tissues. As a result, a non-lipid rich tissue site can be treated by delivering the ultrasound energy to the tissue site before the light activated drug has an opportunity to localize in lipid rich tissues.

Localized delivery is also advantageous when the tissue site is lipid rich such as in an atheroma or a tumor. The localized delivery of the light activated drug combined with the inherent affinity of the light activated drug for tissue site can result in a high degree of localization of the light activated drug within lipid rich tissue sites.

To increase localization of the light activated drug within the tissue site, the light activated drug can be coupled with a site directing molecule to form a light activated drug conjugate. The site directing molecule is chosen so the light activated drug conjugate specifically binds with the tissue site when the light activated drug conjugate is contacted with the tissue site under physiological conditions of temperature and pH. The specific binding may result from specific electrostatic, hydrophobic, entropic, or other interactions between certain residues on the conjugate and specific residues on the tissue site.

In one preferred embodiment, the light activated drug includes an oligonucleotide acting as a site specific molecule coupled with a texaphyrin. The oligonucleotide can have an affinity for a targeted site on a DNA strand. For instance, the oligonucleotide can be designed to have complementary Watson-Crick base pairing with the targeted DNA site. Activation of the light activated drug after the conjugate has bound the targeted DNA site can cause cleavage of the DNA strand at the targeted DNA site. As a result, the activated drug conjugate can be used for cleavage of targeted DNA sites. The light activated conjugate can be targeted to a site on viral DNA where activation of the light activated conjugate causes the virus to be killed. Similarly, the light activated conjugate can be targeted to oncogenes. Other applications of targeted DNA cleavage include, but are not limited to, antisense applications, specific cleavage and subsequent recombination of DNA; destruction of viral DNA; construction of probes for controlling gene expression at the cellular level and for diagnosis; and cleavage of DNA in footprinting analyses, DNA sequencing, chromosome analysis, gene isolation, recombinant DNA manipulations,
mapping of large genomes and chromosomes, in chemotherapy and in site directing mutagenesis.

[0118] In another preferred embodiment, the light activated drug includes a hormone. The hormone may be targeted to a particular biological receptor which is localized at the tissue site.

[0119] The light activated drug can be included within several media suitable for delivery into the body. Many light activated drugs are known to have low water solubilities of less than 100 mg/L. As a result, achieving the desired concentration of light activated drug in an aqueous solution media for systemic delivery can often be difficult. However, localized delivery of the light activated drug requires a lower concentration of light activated drug within the media. As a result, when the light activated drug is delivered locally, the light activated drug can be included in an aqueous solution.

[0120] The media can also be an emulsion which includes a lipid as a hydrophobic phase dispersed in a hydrophilic phase. These emulsions provide a media which is safe for delivery into the body with an effective concentration of light activated drug.

[0121] The media can also include microbubbles comprised from a substrate which forms a shell. Suitable substrates for the microbubble include, but are not limited to, biocompatible polymers, albumins, lipids, sugars or other substances. The light activated drug can be enclosed within the microbubble, coupled with the shell and/or distributed in the media outside the microbubble. A preferred microbubble comprises a lipid substrate such as liposome. Systemic administration of liposomes with light activated drug has been shown to result in an increased accumulation and more prolonged retention of light activated drugs within cultured malignant cells and within tumors in vivo. Jori et al., Br J. Cancer, 48:307-309 (1983); Cozzani et al., In Porphyrins in Tumor Phototherapy, 173-183, Plenum Press (Andreoni et al. eds. 1984).

As a result, inclusion of the light activated drug within a liposome combined with the localized delivery of the light activated drug can serve to enhance the localization of the light activated drug within the tissue site.

[0122] Including a light activated drug with the microbubbles has numerous advantages over microbubbles without light activated drug. After administration of microbubbles to a patient, the microbubbles often must be ruptured to achieve their therapeutic effects. One technique for rupturing microbubbles has been to expose the microbubbles to ultrasound energy. However, ultrasound energy of undesirably high intensity is frequently required to break the microbubbles. Further, the ultrasound energy frequency must be matched to the resonant frequency of the microbubbles. As a result, rupturing the microbubbles with ultrasound can present numerous challenges.

[0123] Activating a light activated drug within the microbubble and/or in the substrate of the microbubble can cause the microbubble to rupture. Activation of the light activated drug is believed to cause a disturbance which disrupts the shell of the microbubble enough to cause the microbubble to rupture. This disruption occurs when the light activated drug is coupled with the shell of the microbubble or is entirely within the microbubble. This disruption is also believed to occur when light activated drug located the media outside the microbubbles is activated in proximity of the microbubble. Accordingly, including a sufficient concentration light activated drug in the media outside the microbubble and activating a portion of that light activated drug can also cause rupture of the microbubbles. As a result, microbubbles can be ruptured by activating light activated drugs and without matching the ultrasound frequency to the resonant frequency of the microbubble. However, a more efficient rupturing of microbubbles can be achieved by delivering a level of ultrasound energy which is appropriate to activate the light activated drug and which is matched to the resonant frequency of the microbubble. Further, the cavitation threshold can require an ultrasound intensity which is lower than the intensity required to rupture microbubbles without light activated drugs. As a result, including light activated drug with microbubbles can reduce the intensity of ultrasound energy required to rupture the microbubble.

[0124] The threshold value of cavitation is also reduced in the proximity of many light activated drugs. As a result, the light activated drug encourages cavitation in the proximity of the light activated drug.

[0125] The interior of the microbubbles may include a gas or may be devoid of gas. When a gas is present, the gas can occupy any portion of the microbubble's volume but preferably occupies 0.01-50% of the volume of the microbubble interior, more preferably 5-30% and most preferably 10-20%. When the volume of gas is less than 0.01% of the volume, cavitation can be hindered and when the volume of gas is greater than 50% the structural integrity of the microbubble shell can become too weak for the microbubble to be transported to the tissue site. Suitable gasses for the interior of the microbubbles include, but are not limited to, biocompatible gasses such as air, nitrogen, carbon dioxide, oxygen, argon, fluorine, xenon, neon, helium, or combinations thereof. The presence of tiny bubbles is known to reduce the cavitation threshold. As a result, the presence of an appropriately sized gas bubble in the microbubble can enhance cavitation in the proximity of the light activated drug.

[0126] The microbubbles are preferably 0.01-100 μm in diameter. This size microbubble reduces excretion of the microbubble outside the body and also reduces interference of the microbubble with the flow of fluids within the body of the patient. Further, the microbubbles preferably have a shell thickness of 0.001-50 μm, 0.01-5 μm and 0.1-0.5 μm. This thickness provides the shells with sufficient thickness that the microbubble can withstand enough of the forces within the vasculature of a patient to be transported through at least a portion of the patient’s vasculature. Similarly, the thickness can permit the microbubbles to be transported through a lumen in an apparatus such as a catheter. However, this thickness is also sufficiently thin that alteration of the ultrasound activated substance upon activation is sufficient to disrupt the shell of the microbubble and cause the microbubble to rupture.

[0127] Activating the light activated drug to rupture microbubbles can cause the light activated drug to be released from the microbubble so the light activated drug can penetrate the tissue near the site of rupture. Further exposure of the light activated drug to ultrasound can activate the light activated drug within the tissue and cause death of the tissue as described above.
[0128] The microbubble can include a therapeutic in addition to the light activated drug. Activation of the light activated drug can serve to rupture the microbubble and release the therapeutic from the microbubble. As a result, the therapeutic is released in proximity to a tissue site by rupturing the microbubble in proximity to the tissue site. This is advantageous when the therapeutic can be detrimental when administered systemically. For instance, a therapeutic such as cisplatin is known to kill cancerous tissues but is also known to kill other tissues throughout the body. As a result, systemic administration of cisplatin can be detrimental. However, microbubbles can serve to protect tissues from the therapeutic agent until the therapeutic agent is released from the carrier. For instance, when the therapeutic is enclosed within the interior of the microbubble, contact between the therapeutic agent and tissues outside the carrier is reduced. As a result, the carrier increases protection of tissues outside the carrier are protected from the therapeutic agent until the microbubble is ruptured and the therapeutic released.

[0129] The therapeutics may be encapsulated in the microbubbles, included in the shell of the microbubbles or in the media outside the microbubbles. By therapeutic, as used herein, it is meant an agent having beneficial effect on the patient.

[0130] Examples of therapeutics which can be included with the microbubbles include, but are not limited to, hormone products such as, vasopressin and oxytocin and their derivatives, glucagon and thyroid agents as iodine products and anti-thyroid agents; cardiovascular products as chelating agents and mercurial diuretics and cardiac glycosides; respiratory products as xanthine derivatives (theophylline & aminophylline); anti-infectives as aminoglycosides, antifungals (amphotericin), penicillin and cephalosporin antibiotics, antiviral agents as Zidovudine, Ribavirin, Amantadine, Vidarabine, and Acyclovir, antiherbamines, antimarials, and antituberculous drugs; biologicals as immune serum, antioxidants and antiviruses, rabies prophylaxis products, bacterial vaccines, viral vaccines, toxoids; antineoplastics as antrazosin, nitrogen mustards, antimetabolites (flourouracil, hormones, asprogesins and estrogens and antitumestrogen; antibiotics as Dactinomycin; mitotic inhibitors as Etoposide and the Vinca alkaloids, Radiopharmaceuticals as radioactive iodine and phosphorus products; as well as Interferon, hydroxyurea, procarbazine, Dacarbazine, Mitotane, Asparaginase and cyclosporins.

[0131] Other suitable therapeutics include, but are not limited to: thrombolytic agents such as urokinase; coagulants such as thrombin; anticoagulant agents, such as platin compound (e.g., spiroplatin, cisplatin, and carboplatin), methotrexate, adriamycin, taxol, mitomycin, ansamitocin, bleomycin, cytosine arabinoside, arabinosyl adnine, mercaptopolylysine, vincristine, busulfan, chlorambucil, melphanal (e.g., PAM, L-PAM or phenylalanine mustard), mercaptouridine, mitotane, procarbazine hydrochloride dactinomycin (actinomycin D), daunorubicinhydrochloride, doxorubicin hydrochloride, mitomycin, plicamycin (mithramycin), aminoglutethimide, estramustine phosphate sodium, flutamide, leuprolide acetate, megestrol acetate, tamoxifen citrate, testolactone, triostane, amsacrine (m-AMS), asparaginase (L-asparaginase) Erwinasparaginase, etoposide (VP-16), interferon alpha-2a, interferon alpha-2b, teniposide (VM-26), vinblastine sulfate (VLB), vincristine sulfate, bleomycin, bloemycin sulfate, methotrexate, adriamycin, and arabinosyl; blood products such as parenteral iron, hemin; biological response modifiers such as muramyldecapeptide, muramyltripptide, microbial cell wall components, lymphokines (e.g., bacterial endotoxin such as lipopolysaccharide, macrophage activationfactor), sub-units of bacteria (such as Mycobacteria, Corynebacteria), the synthetic dipeptide N-acetyl-muramyl-L-alanyl-D-isoglutamine; anti-fungal agents such as ketoconazole, nystatin, griseofulvin, fluconysol (5-fluc), miconazole, amphoteric B, ricin, and beta-lactam antibiotics (e.g., sulfazecin); hormones such as growth hormone, melanocyte stimulating hormone, estradiol, beclomethasone dipropionate, betamethasone, betamethasone acetate and betamethasone sodium phosphate, betamethasone disodium phosphate, betamethasone sodium phosphate, cortisone acetate, dexamethasone, dexamethasone acetate, dexamethasone sodium phosphate, flumisolide, hydrocortisone, hydrocortisone acetate, hydrocortisone cypionate, hydrocortisone sodium phosphate, hydrocortisone sodium succinate, methylprednisolone, methylprednisolone acetate, methylprednisolone sodium succinate, paramethasone acetate, prednisolone, prednisoloneacetate, prednisolone sodium phosphate, prednisolone acetate, prednisone, triamcinolone, triamcinolone acetonide, triamcinolone diacetate, triamcinolone hexacetonide and fluadrocortisone acetate; vitamins such as ascorbic acid, vitamins B, C, and D; amino acids such as aminoacetic acid, isoniazid, capreomycin sulfate, cyclocaine, ethambutol hydrochloride ethionamide, pyrazinamide, rifampin, and streptomycin sulfate; antivirals such as acyclovir, amantadine azidothymidine (AZT or Zidovudine), Ribavirin and ribavirina monohydrate (adenine arabinoside, ara-A); antianginals such as lidocain, nifedipine, verapamil, erythritol, tetraniitrate, isosorbidinitrate, nitroglycerin (glyceryl trinitrate) and pentylaetitrotetranitrare; antigout such as phenprocoumon, heparin; antibiotics such as doxipam, chloramphenicol, neomycin, cefaclor, cefadroxil, cephalxin, cephradine erythromycin, clindamycin, lincomycin, amoxicillin, ampicillin, bacampicillin, carbencillin, dichloracillin, cefaclor, plicamycin, plicacillin, tetracyclic antibiotic, naltexerin, oxacillin, penicillin G, penicillin V, ticarcillin, rifampin and tetracycline; anti-inflammatory such as diflunisal, ibuprofen, indomethacin, meclofenamate, mafenamic acid, naproxen, oxyphenbutazone, phenylbutazone, piroxicam, sulindac, tolmetin, aspirin and salicylates; antiprotozoal such as chloroquine, hydroxychloroquine, metronidazole, quinine and meglumine antimonate; anti-tumestrogen such as tamoxifen; narcotics such as pargor; opiates such as codeine, heroin, methadone, morphine and opium; cardiac glycosides such as digitoxin, digoxin, digitoxin and digitalis; neuromuscular blockers such as atracurium besylate, gallamine triethiodide, hexafluorone bromide, metocurine iodide, pancuronium bromide, succinylcholine chloride (suxamethonium chloride), tubocurarine chloride and vecuronium bromide; sedatives (hypnotics) such as amobarbital, amobarbital sodium, apropobarbital, butabarbital sodium, chloral hydrate, ethchlorvynol, eluthan, flurazepam hydrochloride, glutethimide,
methotrimeprazine hydrochloride, methyprylon, midazolam hydrochloride, paraldehyde, pentobarbital, pentobarbital sodium, phenobarbital sodium, secobarbital sodium, talbutal, temazepam and triazolam; local anesthetics such as bupivacaine hydrochloride, chlorprocaine hydrochloride, etidocainehydrochloride, lidocaine hydrochloride, meperidine hydrochloride, procainehydrochloride and tetracaine hydrochloride; general anesthetics such as dospironidol, etomidate, lentanyl citrate with droperidol, ketamínhydrochloride, methohexital sodium and thiopental sodium; and radio-active particles or ions such as strontium, iodide, rhenium and yttrium.

[0132] In certain preferred embodiments, the therapeutic is a monoclonal antibody, such as a monoclonal antibody capable of binding to melanoma antigen.

[0133] Other preferred therapies include genetic material such as nucleic acids, RNA, and DNA, of either natural or synthetic origin, including recombinant RNA and DNA and antisense RNA and DNA. Types of genetic material that may be used include, for example, genes carried on expression vectors such as plasmids, phagemids, cosmids, yeast artificial chromosomes (YACs), and defective or “helper” viruses, antigenic nucleic acids, both single and double stranded RNA and DNA and analogs thereof, such as phosphorothioate and phosphorothioate oligodeoxynucleotides. Additionally, the genetic material may be combined, for example, with proteins or other polymers.

[0134] Examples of genetic therapies that may be included in the microbubbles include DNA encoding at least a portion of an HLA gene, DNA encoding at least a portion of dystrophin, DNA encoding at least a portion of CFTR, DNA encoding at least a portion of IL-2, DNA encoding at least a portion of INF, an antisense oligonucleotide capable of binding the DNA encoding at least a portion of Ras.

[0135] DNA encoding certain proteins may be used in the treatment of many different types of diseases. For example, adenosine deaminase may be provided to treat ADA deficiency; tumor necrosis factor and/or interleukin-2 may be provided to treat advanced cancers; HDL receptor may be provided to treat liver disease; thymidine kinase may be provided to treat ovarian cancer, brain tumors, or HIV infection; ILA-D7 may be provided to treat malignant melanoma; interleukin-2 may be provided to treat neuroblastoma, malignant melanoma, or kidney cancer; interleukin-4 may be provided to treat cancer; HIV-Env may be provided to treat HIV infection; antisense ras/p53 may be provided to treat lung cancer; and Factor VIII may be provided to treat Hemophilia B. See, for example, Science 258, 744-746.

[0136] If desired, more than one therapeutic may be included in the media. For example, a single microbubble may contain more than one therapeutic or microbubbles containing different therapeutics may be co-administered. By way of example, a monoclonal antibody capable of binding to melanoma antigen and an oligonucleotide encoding at least a portion of IL-2 may be administered in a single microbubble. The phrase “at least a portion of,” as used herein, means that the entire gene need not be represented by the oligonucleotide, so long as the portion of the gene represented provides an effective block to gene expression. Further, microbubbles including a therapeutic can be administered before, after, during or intermittently with the administration of microbubbles without a therapeutic. For instance, microbubbles without a therapeutic and microbubbles including a coagulant such as thrombin can be administered to a patient having liver cancer. Activating the light activated drug with the microbubbles serves to rupture the microbubbles and release the light activated drug and thrombin from the microbubbles. Further activation of the light activated drug can cause tissue death and the thrombin can cause coagulation in and around the damaged tissues.

[0137] Prudrgs may be included in the microbubbles, and are included within the ambit of the term therapeutic, as used herein. Prudrgs are well known in the art and include inactive drug precursors which, when exposed to high temperature, metabolizing enzymes, cavitation and/or pressure, in the presence of oxygen or otherwise, or when released from the microbubbles, will form active drugs. Such prudrgs can be activated via the application of ultrasound to the prudrg-containing microbubbles with the resultant cavitation, heating, pressure, and/or release from the microbubbles. Suitable prudrgs will be apparent to those skilled in the art, and are described, for example, in Sinkula et al., J. Pharm. Sci. 1975 64, 181-210, the disclosure of which is hereby incorporated herein by reference in its entirety. Prudrgs, for example, may comprise inactive forms of the active drugs wherein a chemical group is present on the prudrg which renders it inactive and/or confers solubility or some other property to the drug. In this form, the prudrgs are generally inactive, but once the chemical group has been cleaved from the prudrg, by heat, cavitation, pressure, and/or by enzymes in the surrounding environment or otherwise, the active drug is generated. Such prudrgs are well described in the art, and comprise a wide variety of drugs bound to chemical groups through bonds such as esters to short, medium or long chain aliphatic carboxates, hemiesters of organic phosphate, pyrophosphate, sulfate, amides, amino acids, azo bonds, carbamate, phosphamide, glucosiduronate, N-acetylglucosamine and beta-glucoside. Examples of drugs with the parent molecule and the reversible modification or linkage are as follows: convallatoxin with ketals, hydantoin with alkyl esters, chlorphenesin with glycine or alanins esters, acetaminophen with caffeine complex, acetalsaliclyc acid with THAM salt, acetylsalicylic acid with acetamidophenyl ester, nalozone with sulfateester, 15-methylprostaglandin F 2 with methyl ester, procaine with polyethylene glycol, erythromycin with alkyl esters, eilindamycin with allylesters or phosphate esters, tetracyclines with betaines salts, 7-acylaminocephalosporins with ring-substituted acyloxybenzyl esters, mandrolone with phenylproprionate decanoate esters, estradiol with enolether acetal, methylprednisolone with acetate esters, testosterone with N-acetylglucosaminide glucosiduronate (trimethylsilyl) ether, cortisol or prednisone or dexamethasone with 21-phosphate esters. Prudrgs may also be designed as reversible drug derivatives and utilized as modifiers to enhance drug transport to site-specific tissues. Examples of parent molecules with reversible modifications or linkages to influence transport to a site specific tissue and for enhanced therapeutic effect include isocyanate with halooalkyl nitrosurea, testosterone with propionateester, metrotrexate (3-5-dichloromethotrexate) with diakyl esters, cytosine arabinoside with 5-acylate, nitrogen mustard (2,2′- dichloro-N-methyldiethylamine), nitrogen mustard with aminomethyltetraacycline, nitrogen mustard with
cholesterol or estradiol or dehydroepiandrosterone esters and nitrogen mustard with azobenzene. As one skilled in the art would recognize, a particular chemical group to modify a given drug may be selected to influence the partitioning of the drug into either the shell or the interior of the microbubbles. The bond selected to link the chemical group to the drug may be selected to have the desired rate of metabolism, e.g., hydrolysis in the case of ester bonds in the presence of serum esterases after release from the microbubbles. Additionally, the particular chemical group may be selected to influence the biodistribution of the drug employed in the microbubbles, e.g., N,N-bis(2-chloroethyl)-phosphorodiamidic acid with cyclic phosphoramidate for ovarian adenocarcinoma. Additionally, the prodrugs employed within the microbubbles may be designed to contain reversible derivatives which are utilized as modifiers of duration of activity to provide, prolong or depot action effects. For example, nicotinic acid may be modified with dextran and carboxymethyl dextran esters, streptomyacin with alginic acid salt, dihydrostreptomycin with pamoate salt, cytarabine (ara-C) with 5'-adamantano ester, ara-adenosine (ara-A) with 5-palmitate and 5-benzoate esters, amphetamine B with methyl esters, testosterone with 17-beta-alkyl esters, estradiol with formate ester, proglandin with 2-(4-imidazolyl) ethylamine salt, dopamine with amino acid amides, chloramphenicol with mono- and bis(trimethylsilyl) ethers, and cycloguanil with pamoate salt. In this form, a depot or reservoir of long-acting drug may be released in vivo from the prodrug bearing microbubbles. In addition, compounds which are generally thermally labile may be utilized to create toxic free radical compounds. Compounds with azolinkages, peroxides and disulfide linkages which decompose with high temperature are preferred. With this form of prodrug, azo, peroxide or disulfide bond containing compounds are activated by cavitaton and/or increased heating caused by the interaction of ultrasound with the microbubbles to create cascades of free radicals from these prodrugs entrapped therein. A wide variety of drugs or chemicals may constitute these prodrugs, such asazo compounds, the general structure of such compounds being R—N=N—R, wherein R is a hydrocarbon chain, where the double bond between the two nitrogen atoms may react to create free radical products in vivo. Exemplary drugs or compounds which may be used to create free radical products include azo containing compounds such as azobenzene, 2,2'-azo bisobutyronitrile, azodicarbonamide, azoliminate, azomycin, azosemide, azosulfamide, azoxybenzene, aztreonam, sudan III, sulfachrysoidine, sulfamidochrysoidine and sulfasalazine, compounds containing disulfide bonds such as sulfuric acid, thiamine disulfide, thioluln, thiram, compounds containing peroxides such as hydrogen peroxide and benzoyl peroxide, 2,2'-azo bisobutyronitrile, 2,2'-azo bis(2-amidopropane) dihydrochloride, and 2,2'-azo bis(2,4-dimethylbenzenonitrile). A microbubble having oxygen gas on its interior should create extensive free radicals with cavitaton. Also, metal ions from the transition series, especially manganese, iron and copper can increase the rate of formation of reactive oxygen intermediates from oxygen. By including metal ions within the microbubbles, the formation of free radicals in vivo can be increased. These metal ions may be incorporated into the microbubbles as freestals, as complexes, e.g., with EDTA, DTPA, DOTA or desferrioxamine, or as oxides of the metal ions. Additionally, derivatized complexes of the metal ions may be bound to lipid head groups, or lipophilic complexes of the ions may be incorporated into a lipid bilayer, for example. When exposed to thermal stimulation, e.g., cavitaton, these metal ions then will increase the rate of formation of reactive oxygen intermediates. Further, radiosensitizers such as metronidazole and misonidazole may be incorporated into the gas-filled liposomes to create free radicals on thermal stimulation. By way of an example of the use of prodrugs, an acylated chemical group may be bound to a drug via an ester linkage which would readily cleave in vivo by enzymatic action in serum. The acylated prodrug can be included in the microbubble. When the microbubble is ruptured, the prodrug will then be exposed to the serum. The ester linkage is then cleaved by esterases in the serum, thereby generating the drug. Similarly, ultrasound may be utilized not only to activate the light activated drug so as to burst the gas-filled liposome, but also to cause thermal effects which may increase the rate of the chemical cleavage and the release of the active drug from the prodrug. The microbubbles may also be designed so that there is a symmetric or an asymmetric distribution of the therapeutic both inside and outside of the microbubble. The particular chemical structure of the therapeutics may be selected or modified to achieve desired solubility such that the therapeutic may either be encapsulated within the interior of the microbubble or couple with the shell of the microbubble. The shell-bound therapeutic may bear one or more acyl chains such that, when the microbubble is popped or heated or ruptured via cavitation, the acylated therapeutic may then leave the surface and/or the therapeutic may be cleaved from the acyl chains chemical group. Similarly, other therapeutics may be formulated with a hydrophobic group which is aromatic or sterol in structure to incorporate into the surface of the microbubble. When the microbubble is a liposome, the liposome can be “fast breaking”. In fast breaking liposomes, the light activated drug-liposome combination is stable in vitro but, when administered in vivo, the light activated drug is rapidly released into the bloodstream where it can associate with serum lipoproteins. As a result, the localized delivery of liposomes combined with the fast breaking nature of the liposomes can result in localization of the light activated drug and/or the therapeutic in the tissues near the catheter. Further, the fast breaking liposomes can prevent the liposomes from leaving the vicinity of the catheter intact and then concentrating in non-targeted tissues such as the liver. Delivery of ultrasound energy from the catheter can also serve to break apart the liposomes after they have been delivered from the catheter. A catheter for locally delivering a media including a light activated drug includes an elongated body with at least one utility lumen extending through the elongated body. The utility lumens can be used to deliver the media including the light activated drug locally to a tissue site and/or to receive a guidewire so the catheter can be guided to the tissue site. The ultrasound assembly can include an ultrasound transducer designed to transmit ultrasound energy which activates the light activated drug. A support member can support the ultrasound transducer adjacent to an outer surface of the elongated body so as to define a chamber between the ultrasound transducer and the elongated body. The chamber can be filled with a material which creates a low acoustic impedance to reduce the exposure of at least one utility lumen within the elong-
gated body to ultrasound energy delivered from the ultrasound transducer. For instance, the chamber can be filled with a material which absorbs, reflects or prevents transmission of ultrasound energy through the chamber. Alternatively, the chamber can be evacuated to reduce transmission of ultrasound energy through the chamber. Reducing the exposure of at least one lumen to the ultrasound energy reduces exposure of media delivered through the at least one lumen to the ultrasound energy. As a result, the effect of the ultrasound energy on the light activated drug is reduced until the light activated drug has been delivered out of the catheter. Further, ultrasound energy is known to rupture microbubbles. As a result, when the media includes microbubbles, the chamber reduces the opportunity for the ultrasound energy to rupture the microbubbles within the catheter.

[0141] The support member can have ends which extend beyond the ultrasound transducer. As a result, the chamber can be positioned adjacent to the entire longitudinal length of the ultrasound transducer and can extend beyond the ends of the ultrasound transducer. This configuration maximizes the portion of the ultrasound transducer which is adjacent to the chamber. Increasing the portion of ultrasound transducer adjacent to the chamber can reduce the amount of ultrasound energy transmitted to the utility lumens. The ultrasound assembly can include an outer coating over the ultrasound transducer. Temperature sensors can be positioned in the outer coating adjacent to ultrasound transducer. This position of the temperature sensors feedback regarding the temperature adjacent to the ultrasound transducers where the thermal energy has a reduced opportunity to dissipate. As a result, the temperature sensors provide a measure of the temperature on the exterior surface of the transducer.

[0142] FIGS. 1A-1B illustrates a catheter 10 for delivering a media including a light activated drug to a tissue site. The catheter 10 includes an ultrasound assembly 12 for delivering ultrasound energy to light activated drug within the tissue site. The catheter 10 includes an elongated body 14 with a utility lumen 16 extending through the elongated body 14. The utility lumen 16 can receive a guidewire (not shown) so the catheter 10 can be threaded along the guidewire. The utility lumen 16 can also be used for the delivering media which include a light activated drug. A fiber optic can also be positioned in the utility lumen 16 to provide a view of the tissue site or to provide light to the tissue site. As a result, the catheter can also be used as an endoscope.

[0143] The ultrasound assembly 12 can also include an outer coating 18. Suitable outer coatings 18 include, but are not limited to, polyimide, parylene and polyester. An ultrasound transducer 20 is positioned within the outer coating 18. Suitable ultrasound transducers 20 include, but are not limited to, PZT-4D, PZT-4, PZT-8 and cylindrically shaped piezoelectrics. When the ultrasound transducer 20 has a cylindrical shape, the ultrasound transducer 20 can encircle the elongated body 14 as illustrated in FIG. 1C. One or more temperature sensors 22 can be positioned in the outer coating 18. The temperature sensors 22 can be positioned adjacent to the ultrasound transducer 20 to provide feedback regarding the temperature adjacent to the ultrasound transducer 20. The temperature sensors can be in electrical communication with an electrical coupling 24. The electrical coupling 24 can be coupled with a feedback control system (not shown) which adjusts the level of the ultrasound energy delivered from the ultrasound transducer 20 in response to the temperature at the temperature sensors 22.

[0144] The catheter 10 can include a perfusion lumen 25. The perfusion lumen 25 allows fluid to flow from outside the catheter into the utility lumen 16. Once a guidewire has been removed from the utility lumen 16, fluid flow which is obstructed by the ultrasound assembly can continue through the perfusion lumen 25 and the utility lumen. As illustrated in FIGS. 2A-2B, the ultrasound assembly 12 can be flush with the elongated body 14. Further, the ultrasound transducer 20 and the temperature sensors 22 can be positioned within the elongated body 14. This configuration of elongated body 14 and ultrasound transducer 20 can eliminate the need for the outer coating 18 illustrated in FIGS. 1A-1C.

[0145] As illustrated in FIG. 3A, the catheter 10 can also include a media delivery port 26, a media inlet port 28 and a second utility lumen 16A. The media inlet port 28 is designed to be coupled with a media source (not shown). Media can be transported from the media source and through the media delivery port 26 via the second utility lumen 16A. As a result, a guidewire can be left within the utility lumen 16 while media is delivered via the second utility lumen 16A.

[0146] FIG. 4A illustrates a catheter 10 including a plurality of ultrasound assemblies 12. FIGS. 4B-4C are cross sections of a catheter 10 with second utility lumen 16A coupled with the media delivery ports 26. The second utility lumen 16A can also be coupled with the media inlet port 28 illustrated in FIG. 4A. The media inlet port 28 is designed to be coupled with a media source (not shown). Media can be transported from the media source and through the media delivery ports 26 via the second utility lumen 16A.

[0147] The catheter 10 can include a balloon 30 as illustrated in FIG. 5A. The balloon 30 can be constructed from an impermeable material or a permeable membrane or a selectively permeable membrane which allows certain media to flow through the membrane while preventing other media from flowing through the membrane. Suitable membranous materials for the balloon 30 include, but are not limited to cellulose, cellulose acetate, polyvinylchloride, polyolein, polyurethane and polysulphone. When the balloon 30 is constructed from a permeable membrane or a selectively permeable membrane, the membrane pore sizes are preferably 5 A-2 μm, more preferably 50 A-900 A and most preferably 100 A-300 A in diameter.

[0148] As illustrated in FIG. 5B, an ultrasound assembly 12, a first media delivery port 26A and a second media delivery port 26B can be positioned within the balloon 30. The first and second media delivery ports 26A, 26B are coupled with a second utility lumen 16A and third utility lumen 16B. The second and third utility lumens 16A, 16B can be coupled with the same media inlet port 28 or with independent media inlet ports 28. When the first and second media delivery ports 26A, 26B are coupled with different media inlet ports 28, different media can be delivered via the second and third media delivery ports 26A, 26B. For instance, a medication media can be delivered via the third utility lumen 16B and an expansion media can be delivered via the second utility lumen 16A. The medication media can include drugs or other medicaments which can provide a therapeutic effect. The expansion media can serve to expand...
the balloon 30 or wet the membrane comprising the balloon 30. Wetting the membrane comprising the balloon 30 can cause a minimally permeable membrane to become permeable.

[0149] The ultrasound assembly 12 can be positioned outside the balloon 30 as illustrated in FIGS. 6A-6C. In FIG. 6A the balloon 30 is positioned distally of the ultrasound assembly 12 and in FIG. 6B the ultrasound assembly 12 is positioned distally of the balloon 30. FIG. 6C is a cross section a catheter 10 with an ultrasound assembly 12 positioned outside the balloon 30. The catheter includes a second utility lumen 16A coupled with a first media delivery port 26A. The second utility lumen 16A can be used to deliver an expansion media and/or a medication media to the balloon 30. When the balloon 30 is constructed from a permeable membrane, the medication media and/or the expansion media can pass through the balloon 30. Similarly, when the balloon 30 is constructed from a selectively permeable membrane, particular components of the medication media and/or the expansion media can pass through the balloon 30. Pressure can be used to drive the media or components of the media across the balloon 30. Other means such as phoresis can also be used to drive the media or components of the media across the balloon 30.

[0150] As illustrated in FIG. 6C, the ultrasound assembly 12 may be positioned at the distal end of the catheter 10. The second utility lumen 16A can be used to deliver an expansion media and/or a medication media to the balloon 30. The utility lumen 16 can be used to deliver a medication media as well as to guide the catheter 10 along a guidewire.

[0151] As illustrated in FIGS. 7A-7C, the catheter 10 can include a second media delivery port 26B positioned outside the balloon. In FIGS. 7A-7C the ultrasound assembly 12 and the second media delivery port 26B are positioned distally relative to a balloon 30, however, the balloon 30 can be positioned distally relative to the ultrasound assembly 12 and the second media delivery port 26B. In FIG. 7A the ultrasound assembly 12 is positioned distally of the second media delivery port 26B and in FIG. 7B the second media delivery port 26B is positioned distally of the ultrasound assembly 12.

[0152] FIG. 7C is a cross section of the catheter 10 illustrated in FIG. 7A. The catheter 10 includes first and second media delivery ports 26A, 26B coupled with a second utility lumen 16A and third utility lumen 16B. The second and third utility lumens 16A, 16B can be coupled with independent media inlet ports 28 (not shown). The second utility lumen 16A can be used to deliver an expansion media and/or a medication media to the balloon 30 while the third utility lumen 16B can be used to deliver a medication media through the second media delivery port 26B.

[0153] As illustrated in FIGS. 8A-8B, the catheter 10 can include a first balloon 30A and a second balloon 30B. The ultrasound assembly 12 can be positioned between the first and second balloons 30A, 30B. A second media delivery port 26A can optionally be positioned between the first and second balloons 30A, 30B. In FIG. 8A the second media delivery port 26B is positioned distally relative to the ultrasound assembly and in FIG. 8B the ultrasound assembly is positioned distally relative to the second media delivery port 26B.

[0154] FIG. 8C is a cross section of the first balloon 30A illustrated in FIG. 8B. The catheter includes a second, third and fourth utility lumens 16A, 16B, 16C. The second utility lumen 16A is coupled with a first media delivery port 26A within the first balloon. The third utility lumen 16B is coupled with the second media delivery port 26B and the fourth utility lumen 16C is coupled with a third media delivery port 26C in the second balloon 30B (not shown). The second and fourth utility lumens 16A, 16C can be used to deliver expansion media and/or medication media to the first and second balloon 30A, 30B. The second and fourth utility lumens 16A, 16C can be coupled with the same media inlet port or with independent media inlet ports (not shown). When the second and fourth utility lumens are coupled with the same media inlet port, the pressure within the first and second balloons 30A, 30B will be similar. When the second and fourth utility lumens are coupled with independent media inlet ports, different pressures can be created within the first and second balloons 30A, 30B. The third utility lumen 16B can be coupled with an independent media inlet port and can be used to deliver a medication media via the second media delivery port 26B.

[0155] FIGS. 9A-91 illustrate operation of various embodiments of catheters 10 for delivering ultrasound energy to a light activated drug within a tissue site. FIGS. 9A-91 illustrate the tissue site 32 as an atheroma in a vessel 34, however, it is contemplated that the catheter 10 can be used with other tissue sites 32 such as a tumor and that the catheter 10 can be positioned within the vasculature of the tumor. In each of FIGS. 9A-91, the catheter 10 is illustrated as being within a vessel 34. The catheter 10 can be positioned within the vessel 34 by applying conventional over-the-guidewire techniques and can be verified by including radiopaque markers upon the catheter 10.

[0156] In FIG. 9A, the catheter 10 is positioned so the ultrasound assembly 12 is adjacent to a tissue site 32 within a vessel 34. When the catheter 10 is in position, the guidewire is removed from the utility lumen 16 and media can be delivered via the utility lumen 16 as illustrated by the arrows 36. In FIG. 9A, the media includes microparticles 38 but can alternatively be an emulsion. The media is delivered to the tissue site 32 via the utility lumen 16 and ultrasound energy 40 is delivered from the ultrasound assembly 12. Suitable periods for delivering the ultrasound energy include, but are not limited to, 1 minute to three hours, 2 minutes to one hour and 10-30 minutes.

[0157] Suitable intensities for the ultrasound energy include, but are not limited to, 0.1-1000 W/cm², 1-100 W/cm² and 10-50 W/cm². Suitable frequencies for the ultrasound energy include, but are not limited to, 10 kHz-100 MHz and 10 kHz-50 MHz but is preferably 20 kHz-10 MHz. Suitable ultrasound energies also include, but are not limited to 0.02 to 10 W/cm² at a frequency of about 20 kHz to about 10 MHz and more preferably about 0.3 W/cm² at a frequency of about 1.3 MHz. The ultrasound energy can be intermittently switched between a first and second frequency to increase the efficiency of microparticle rupture and to increase activation of the light activated drug. For instance, the ultrasound energy can be switched between about 100 kHz and about 270 kHz in short pulses of approximately 0.001-10 seconds duration. Similarly, the ultrasound energy can be switched between first and second intensities. When the catheter includes a plurality of ultrasound transducers as
will be discussed below, the first and second frequencies can be provided by different ultrasound transducers. Similarly, the first and second intensities can be provided by different ultrasound transducers. Further, when the catheter includes a plurality of ultrasound transducers each transducer can simultaneously transmit ultrasound energy with different intensity and/or frequency.

[0158] The delivery of ultrasound energy 40 can be before, after, during or intermittently with the delivery of the microbubbles 38. As discussed above, the microbubbles 38 can be "fast breaking" so they rupture upon exiting the utility lumen and being exposed to the vessel 34. As described above, the ultrasound energy from the ultrasound assembly 12 can cause the microbubbles 38 within the delivered media to rupture. As will be described in more detail below, the ultrasound assembly can be designed to reduce the exposure of media within the catheter 10 to the ultrasound energy from the ultrasound assembly 12. When the catheter 10 is so designed, the number of microbubbles 38 which rupture within the catheter is reduced and the number of microbubbles 38 which rupture outside the catheter is increased.

[0159] Delivery of the ultrasound energy before delivery of the light activated drug can enhance absorption of the light activated drug into the tissue site. Delivery of the ultrasound energy a pre-determined time after delivery of the light activated drug can provide the light activated drug time to penetrate the tissue site. The pre-determined time can be of sufficient duration that at least a portion of the light activated drug penetrates into the tissue site. The pre-determined time can also be of sufficient duration that the light activated drug localizes within the lipid rich tissue of the atheroma. Sufficient time between delivery of the media and the ultrasound energy include but are not limited to, 1 minute to 48 hours, 1 minute to 3 hours, 1 to 15 minutes and 1 to 2 minutes. Once the light activated drug has penetrated the tissue site 32, the ultrasound energy from the ultrasound assembly 12 can activate the light activated drug within the tissue site 32 so as to cause tissue death within the tissue site 32.

[0160] In FIG. 9B, ultrasound energy 40 is delivered from the ultrasound transducer 20 and a media is delivered through the media delivery port 26 as illustrated by the arrows 36. The delivery of ultrasound energy 40 can be before, after, during or intermittently with the delivery of the media via the media delivery port 26. As illustrated in FIG. 9C, the guidewire 104 can remain in the utility lumen 16 during the delivery of the media via the media delivery ports 26. As will be discussed in further detail below, the ultrasound assembly can be designed to reduce the transmission of the ultrasound energy into the utility lumen. Because the transmission of ultrasound energy 40 into the utility lumen 16 is reduced, the change in the frequency of the ultrasound transducer 20 which is due to the presence of the guidewire in the utility lumen 16 is also reduced.

[0161] In FIG. 9D, a catheter 10 including a balloon 30 is positioned with the balloon adjacent to the tissue site 32. In FIG. 9E, the balloon 30 is expanded into contact with the tissue site 32. As discussed above, the catheter 10 can include a perfusion lumen which permits a continuous flow of fluid from the vessel through the utility lumen during the partial or full obstruction of the vessel by the balloon. When the balloon 30 is constructed from a membrane or a selectively permeable membrane a media can be delivered to the tissue site 32 via the balloon 30. The media can serve to wet the membrane or can include a drug or other medicament which provides a therapeutic effect. Ultrasound energy 40 can be delivered from the ultrasound assembly 12 before, after, during or intermittently with the delivery of the media. The ultrasound energy 40 can serve to drive the media across the membrane via phonophoresis or can enhance the therapeutic effect of the media.

[0162] In FIG. 9F a catheter 10 with an ultrasound assembly 12 outside a balloon 30 is positioned at the tissue site 32 so the ultrasound assembly 12 is adjacent to the tissue site 32. A fluid within the vessel flows past the balloon as indicated by the arrow 42. In FIG. 9G, the balloon 30 is expanded into contact with the vessel 34. The balloon 30 can be constructed from an impermeable material so the vessel 34 is occluded. As a result, the fluid flow through the vessel 34 is reduced or stopped. A medication media is delivered through the utility lumen 16 and ultrasound energy 40 is delivered from the ultrasound assembly 12. In embodiments of the catheter 10 including a media delivery port 26 outside of the balloon 30 (i.e. FIGS. 7A-7C), the medication media can be delivered via the media delivery port 26. Further, a first medication media can be delivered via the media delivery port 26 while a second medication media can be delivered via the utility lumen 16 or while a guidewire is positioned within the utility lumen 16. The ultrasound energy 40 can be delivered from the ultrasound assembly 12 before, after, during or intermittently with the delivery of the media. The occlusion of the vessel 34 before the delivery of the media can serve to prevent the media from being swept from the tissue site 32 by the fluid flow. Although the balloon 30 illustrated in FIGS. 9F-9G is positioned proximally relative to the ultrasound assembly 12, the fluid flow through the vessel 34 can also be reduced by expanding a single balloon 30 which is positioned distally relative to the ultrasound assembly 12.

[0163] In FIG. 9H a catheter 10 including a first balloon 30A and a second balloon 30B is positioned at a tissue site 32 so the ultrasound assembly 12 is positioned adjacent to the tissue site 32. A fluid within the vessel 34 flows past the balloon 30 as indicated by the arrow 42. In FIG. 9I, the first and second balloons 30A, 30B are expanded into contact with the vessel 34. The first and second balloons 30A, 30B can be constructed from an impermeable material so the vessel 34 is occluded proximally and distally of the ultrasound assembly 12. As a result, the fluid flow adjacent to the tissue site 32 is reduced or stopped. A medication media is delivered through the media delivery port 26 and ultrasound energy 40 is delivered from the ultrasound assembly 12. The ultrasound energy 40 can be delivered from the ultrasound assembly 12 before, after, during or intermittently with the delivery of the media. The occlusion of the vessel 34 before the delivery of the media can serve to prevent the media from being swept from the tissue site 32 by the fluid flow.

[0164] In each of the FIGS. 9A-91 illustrated above, the media can be systemically delivered. The catheter 10 is positioned adjacent to the tissue site before, after or during the systemic administration of the media. When the media includes microbubbles which must be burst before their therapeutic effect can be obtained, the ultrasound energy can be delivered after the microbubbles have had sufficient time
to reach the desired tissue site in sufficient concentrations. A level of ultrasound which ruptures the microbubbles is then delivered from the ultrasound assembly. After rupture of the microbubbles, the delivery of ultrasound energy can be stopped to provide the light activated drug or other therapeutic time to penetrate the tissue site. The delivery of the ultrasound energy can also be continuous to maximize the number of microbubbles which are burst.

[0165] When the media is systemically delivered and the light activated drug is included in media which does not require an ultrasound activated release, the behavior of the light activated drug within the patient must be taken into consideration. As described above, many light drugs such as the macrocycles, initially disperse throughout the body and where they are taken up by most tissues. After a period of time, usually between 3 and 48 hours, the drug clears from most normal tissue and is retained to a greater degree in lipid rich regions such as the liver, kidney, tumor and atheroma. As a result, when the tissue site is not a lipid rich region, the ultrasound energy should be delivered to the tissue site within 3 to 48 hours of systemically administering the media. However, when the tissue site is lipid rich, improved results can be achieved by waiting 3 to 48 hours after systemic administration of the media before delivering the ultrasound energy.

[0166] FIG. 10A provides a cross section of an ultrasound assembly which reduces transmission of ultrasound energy from the ultrasound transducer into the catheter. The ultrasound assembly 12 includes a support member 44. Suitable support members 44 include, but are not limited to, polyimide, polyester and nylon. The support member 44 can be attached to the ultrasound transducer 20. Suitable means for attaching the ultrasound transducer 20 to the support member 44 include, but are not limited to, adhesive bonding and thermal bonding.

[0167] The support member 44 supports the ultrasound member 44 at an external surface 46 of the elongated body 14 such that a chamber 48 is defined between the ultrasound transducer 20 and the external surface 46 of the elongated body 14. The chamber 48 preferably has a height from 0.25-1 mm, more preferably from 0.50-5 μm and most preferably from 0.0-1.5 μm. The support member 44 can be supported by supports 50 positioned at the ends 52 of the support member 44 as illustrated in FIG. 10B. The supports 50 can be integral with the support member 44 as illustrated in FIG. 10C. The outer coating 18 can serve as the supports as illustrated in FIG. 10D.

[0168] The ends 52 of the support member 44 can extend beyond the ends 54 of the ultrasound transducer 20. The supports 50 can be positioned beyond the ends 54 of the ultrasound transducer 20. As a result, the chamber 48 can extend along the longitudinal length 56 of the ultrasound transducer 20, maximizing the portion of the ultrasound transducer 20 which is adjacent to the chamber 48. The chamber 48 can be filled with a medium which absorbs ultrasound energy or which prevents transmission of ultrasound energy. Suitable gaseous media for filling the chamber 48 include, but are not limited to, helium, argon, air and nitrogen. Suitable liquid media for filling the chamber 48 include, but are not limited to, silicone and rubber. The chamber 48 can also be evacuated. Suitable pressures for an evacuated chamber 48 include, but are not limited to, negative pressures to -760 mm Hg.

[0169] The ultrasound assembly can include a second ultrasound transducer 20A as illustrated in FIGS. 11A-11H. In FIGS. 11A-11C one ultrasound transducer encircles the other and in FIGS. 11D-11H the ultrasound transducers are longitudinally adjacent to one another. The ultrasound transducers 20, 20A can be in contact with one another as illustrated in FIGS. 11A, 11E and 11H or separated from one another as illustrated in FIGS. 11B-11D, 11F and FIG. A single chamber 54 can be defined between the ultrasound transducers 20, 20A and the external surface 46 of the elongated body 14 as illustrated in FIGS. 11C, 11F and 11G or a different chamber can be defined between each of the ultrasound transducers 20, 20A and the external surface 46. Although the ultrasound transducers 20, 20A in FIGS. 11A-11C are illustrated as having the same longitudinal length, the longitudinal length may be different.

[0170] In FIGS. 11A-11H, the different temperature sensors can be positioned adjacent to different ultrasound transducers 20, 20A. As a result, the temperature adjacent to different ultrasound transducers 20, 20A can be detected and the level of ultrasound energy produced by each ultrasound transducer adjusted in response to the detected temperature.

[0171] When the ultrasound assembly includes a second transducer 20A, the transducers 20, 20A may be constructed from the same or different materials. Both transducers 20, 20A may be configured to radiate ultrasound energy in the same direction. Further, one transducer may be configured to transmit ultrasound energy in a radial direction and the other in a longitudinal direction in order to increase the angular spectrum over which ultrasound energy can be simultaneously transmitted. The ultrasound transducers can be configured to transmit ultrasound energy having the same or different characteristics. The transmission of ultrasound energy with different characteristics allows the same ultrasound assemblies to be used to perform different functions. For instance, one ultrasound transducer can transmit a frequency which is appropriate for activating a light activated drug while the second ultrasound transducer transmits a frequency appropriate for enhancing penetration of a therapeutic agent into the treatment site. The transducers can be operated independently or simultaneously. When the transducers are operated simultaneously, the ultrasound assembly produces a waveform which is more complex than a single ultrasound transducer. More complex waveforms can provide advantages such as more efficient rupture of microbubbles. It is also contemplated that the ultrasound assembly can include three or more ultrasound transducers arranged similar to the transducers illustrated in FIGS. 11A-11H.

[0172] The ultrasound assembly 12 can be a separate module 58 as illustrated in FIGS. 12A-12B. In FIG. 12A, the catheter 10 includes a first catheter component 60 and a second catheter component 62 and an ultrasound assembly module 58. The first and second catheter components 60, 62 include component ends 64 which are complementary to the ultrasound assembly module ends 66. The component ends 64 can be coupled with the ultrasound assembly module ends 66 as illustrated in FIG. 12B. Suitable means for coupling the component ends 64 and the ultrasound assembly module ends 66 include, but are not limited to, adhesive, mechanical and thermal methods. The ultrasound assembly 12 can be integral with the catheter 10 as illustrated in FIG. 12C. Further, the outer coating 18 can have a diameter which
is larger than the diameter of the elongated body 14 as illustrated in FIG. 10A or can be flush with the external surface 46 of the elongated body 14 as illustrated in FIGS. 12A-12C.

[0173] The ultrasound assembly 12 can be electrically coupled to produce radial vibrations of the ultrasound transducer 20 as illustrated in FIGS. 13A-13B. A first line 68 is coupled with an outer surface 70 of the ultrasound transducer 20 while a second line 72 is coupled with an inner surface 74 of the ultrasound transducer 20. The first and second lines 68, 72 can pass proximally through the utility lumen 16 as illustrated in FIG. 13A. Alternatively, the first and second lines 68, 72 can pass proximally through line lumens 76 within the catheter 10 as illustrated in FIG. 13B. Suitable lines for the ultrasound transducer 20 include, but are not limited to, copper, gold and aluminum. Suitable frequencies for the ultrasound energy delivered by the ultrasound transducer 20 include, but are not limited to, 20 KHz to 2 MHz.

[0174] The ultrasound assembly 12 can be electrically coupled to produce longitudinal vibrations of the ultrasound transducer 20 as illustrated in FIGS. 13C-13D. A first line 68 is coupled with a first end 78 of the ultrasound transducer 20 while a second line 72 is coupled with a second end 80 of the ultrasound transducer 20. The distal portion 82 of the second line 72 can pass through the outer coating 18 as illustrated in FIG. 13C. Alternatively, the distal portion 82 of the second line 72 can pass through line lumens 76 in the catheter 10 as illustrated in FIG. 13D. As discussed above, the first and second lines 68, 72 can pass proximally through the utility lumen 16.

[0175] As discussed above, the catheter 10 can include a plurality of ultrasound assemblies. When the catheter 10 includes a plurality of ultrasound assemblies, each ultrasound transducer 20 can be individually powered. When the elongated body 14 includes N ultrasound transducers 20, the elongated body 14 must include 2N lines to individually power N ultrasound transducers 20. The individual ultrasound transducers 20 can also be electrically coupled in serial or in parallel as illustrated in FIGS. 14A-14B. These arrangements permit maximum flexibility as they require only 2 lines. Each of the ultrasound transducers 20 receive power simultaneously whether the ultrasound transducers 20 are in series or in parallel. When the ultrasound transducers 20 are in series, less current is required to produce the same power from each ultrasound transducer 20 than when the ultrasound transducers 20 are connected in parallel. The reduced current allows smaller lines to be used to provide power to the ultrasound transducers 20 and accordingly increases the flexibility of the elongated body 14. When the ultrasound transducers 20 are connected in parallel, an ultrasound transducer 20 can break down and the remaining ultrasound transducers 20 will continue to operate.

[0176] As illustrated in FIG. 14C, a common line 84 can provide power to each ultrasound transducer 20 while each ultrasound transducer 20 has its own return line 86. A particular ultrasound transducer 20 can be individually activated by closing a switch 88 to complete a circuit between the common line 84 and the particular ultrasound transducer’s 20 return line 86. Once a switch 88 corresponding to a particular ultrasound transducer 20 has been closed, the amount of power supplied to the ultrasound transducer 20 can be adjusted with the corresponding potentiometer 90. Accordingly, an catheter 10 with N ultrasound transducers 20 requires only N+1 lines and still permits independent control of the ultrasound transducers 20. This reduced number of lines increases the flexibility of the catheter 10. To improve the flexibility of the catheter 10, the individual return lines 86 can have diameters which are smaller than the common line 84 diameter. For instance, in an embodiment where N ultrasound transducers 20 will be powered simultaneously, the diameter of the individual return lines 86 can be the square root of N times smaller than the diameter of the common line 84.

[0177] As discussed above, the ultrasound assembly 12 can include at least one temperature sensor 22. Suitable temperature sensors 22 include, but are not limited to, thermistors, thermocouples, resistance temperature detectors (RTD)s, and fiber optic temperature sensors 22 which use thermochromic liquid crystals. Suitable temperature sensor geometries include, but are not limited to, a point, patch, stripe and a band encircling the ultrasound transducer 20.

[0178] When the ultrasound assembly 12 includes a plurality of temperature sensors 22, the temperature sensors 22 can be electrically connected as illustrated in FIG. 15. Each temperature sensor 22 can be coupled with a common line 84 and then include its own return line 86. Accordingly, N+1 lines can be used to independently sense the temperature at the temperature sensors 22 when N temperature sensors 22 are employed. A suitable common line 84 can be constructed from Constantine and suitable return lines 86 can be constructed from copper. The temperature at a particular temperature sensor 22 can be determined by closing a switch 88 to complete a circuit between the thermocouple’s return line 86 and the common line 84. When the temperature sensors 22 are thermocouples, the temperature can be calculated from the voltage in the circuit. To improve the flexibility of the catheter 10, the individual return lines 86 can have diameters which are smaller than the common line 84 diameter.

[0179] Each temperature sensor 22 can also be independently electrically coupled. Employing N independently electrically coupled temperature sensors 22 requires 2N lines to pass the length of the catheter 10.

[0180] The catheter 10 flexibility can also be improved by using fiber optic based temperature sensors 22. The flexibility can be improved because only N fiber optics need to be employed sense the temperature at N temperature sensors 22.

[0181] The catheter 10 can be coupled with a feedback control system as illustrated in FIG. 16. The temperature at each temperature sensor 22 is monitored and the output power of an energy source adjusted accordingly. The physician can, if desired, override the closed or open loop system.

[0182] The feedback control system includes an energy source 92, power circuits 94 and a power calculation device 96 coupled with each ultrasound transducer 20. A temperature measurement device 98 is coupled with the temperature sensors 22 on the catheter 10. A processing unit 100 is coupled with the power calculation device 96, the power circuits 94 and a user interface and display 102.
In operation, the temperature at each temperature sensor 22 is determined at the temperature measurement device 98. The processing unit 100 receives signals indicating the determined temperatures from the temperature measurement device 98. The determined temperatures can then be displayed to the user at the user interface and display 102.

The processing unit 100 includes logic for generating a temperature control signal. The temperature control signal is proportional to the difference between the measured temperature and a desired temperature. The desired temperature can be determined by the user. The user can set the predetermined temperature at the user interface and display 102.

The temperature control signal is received by the power circuits 94. The power circuits 94 adjust the power level of the energy supplied to the ultrasound transducers 20 from the energy source 92. For instance, when the temperature control signal is above a particular level, the power supplied to a particular ultrasound transducer 20 is reduced in proportion to the magnitude of the temperature control signal. Similarly, when the temperature control signal is below a particular level, the power supplied to a particular ultrasound transducer 20 is increased in proportion to the magnitude of the temperature control signal. After each power adjustment, the processing unit 100 monitors the temperature sensors 22 and produces another temperature control signal which is received by the power circuits 94.

The processing unit 100 can also include safety control logic. The safety control logic detects when the temperature at a temperature sensor 22 has exceeded a safety threshold. The processing unit 100 can then provide a temperature control signal which causes the power circuits 94 to stop the delivery of energy from the energy source 92 to the ultrasound transducers 20.

The processing unit 100 also receives a power signal from the power calculation device 96. The power signal can be used to determine the power being received by each ultrasound transducer 20. The determined power can then be displayed to the user on the user interface and display 102.

The feedback control system can maintain the tissue adjacent to the ultrasound transducers 20 within a desired temperature range for a selected period of time. As described above, the ultrasound transducers 20 can be electrically connected so each ultrasound transducer 20 can generate an independent output. The output maintains a selected energy at each ultrasound transducer 20 for a selected length of time.

The processing unit 100 can be a digital or analog controller, or a computer with software. When the processing unit 100 is a computer it can include a CPU coupled through a system bus. The user interface and display 102 can be a mouse, keyboard, a disk drive, or other non-volatile memory systems, a display monitor, and other peripherals, as are known in the art. Also coupled to the bus is a program memory and a data memory.

In lieu of the series of power adjustments described above, a profile of the power delivered to each ultrasound transducer 20 can be incorporated in the processing unit 100 and a preset amount of energy to be delivered may also be profiled. The power delivered to each ultrasound transducer 20 can then be adjusted according to the profiles.

The above catheters are suitable for locally delivering a media including a light activated drug. Suitable light activated drugs include, but are not limited to, fluorescein, merocyanin. However, preferred light activated drugs include xanthene and its derivatives and the photoactive pyrrole-derived macrocycles and their derivatives due to a reduced toxicity and an increased biological affinity. Suitable photoactive pyrrole-derived macrocycles include, but are not limited to, naturally occurring or synthetic porphyrins, naturally occurring or synthetic chlorins, naturally occurring or synthetic bacteriochlorins, synthetic isobacteriochlorins, phthalocyanines, naphthalocyanines, and expanded pyrrole-based macrocyclic systems such as porphyrazines, sapphyrins, and texaphyrins. Examples of suitable pyrrole-based macrocyclic classes are illustrated in FIG. 17A-17N.

As described above, the derivative of the pyrrole-based macrocycle classes can be used. For the purposes of illustrating some of the derivatives a macrocycle class, FIG. 17B-2 illustrates a formula for the derivatives of texaphyrin where M is H, CH₃, a divalent metal cation selected from the group consisting of Cu(II), Mn(II), Zn(II), Cd(II), Hg(II), Fe(III), Sm(II), and U(VI) or a trivalent metal cation selected from the group consisting of Mn(III), Co(II), Ni(II), Fe(III), Ho(III), Ce(III), Y(III), Ln(III), Pr(III), Nd(III), Sm(III), Eu(III), Gd(III), Tb(III), Dy(III), Er(III), Tm(III), Yb(III), Lu(III), La(III), and U(III). Preferred metals include Lu(III), Dy(III), Eu(III), or Gd(III). M may be H or CH₃ in a non-metalated form of texaphyrin. R₁, R₂, R₃, R₄, R₅ and R₆ can independently be hydrogen, hydroxyl, alkyl, hydroxyalkyl, alkoxy, hydroxalkoxy, saccharide, carboxyalkyl, carboxyamidealkyl, a site-directing molecule, or a linker to a site-directing molecule where at least one of R₁, R₂, R₃, R₄, R₅ and R₆ is hydrogen, hydroxyalkyl, saccharide, alkoxy, carboxyalkyl, carboxyamidalkyl, hydroxyalkyl, a site-directing molecule- or a couple to a site-directing molecule; and N is an integer less than or equal to 2.


R₁, R₂, R₃, R₄, R₅ and R₆ may also independently be amino, carboxy, carboxamide, ester, amide sulfonato, aminoalkyl, sulfonatoalkyl, amidealkyl, aryl, etheramide or equivalent formulae conferring the desired properties. In a preferred embodiment, at least one of R₁, R₂, R₃, R₄, R₅ and R₆ is a site-directing molecule or is a couple to a site-directing molecule. For bulky R groups on the benzene ring portion of the molecule such as oligonucleotides, one skilled in the art would realize that derivatization at one position on the benzene potion is more preferred.
Hydroxyalkyl means alkyl groups having hydroxyl groups attached. Alkoxy means alkyl groups attached to an oxygen. Hydroxyalkoxy means alkyl groups having ether or ester linkages, hydroxy groups, substituted hydroxyl groups, carboxyl groups, substituted carboxyl groups or the like. Saccharide includes oxidized, reduced or substituted saccharide; hexoses such as D-glucose, D-mannose or D-galactose; pentoses such as D-ribose or D-arabinose; ketoses such as D-ribulose or D-fructose; disaccharides such as sucrose, lactose, or maltose; derivatives such as acetals, amides, and phosphorylated sugars; oligosaccharides, as well as examples of amine-derivatized sugars are galactosamine, glucosamine, and sialic acid. Carboxyamidoalkyl means alkyl groups with secondary or tertiary amide linkages or the like. Carboxyalkyl means alkyl groups having hydroxyl groups, carboxyl or amide substituted esters, ester linkages, tertiary amide linkages removed from the ether or the like.

For the above-described texaphyrins, hydroxyalkoxy may be alkyl having independently hydroxy substituents and ether branches or may be C,H,O,O,R or OC(R)xH(OH)H(OH)H(OH)O,R where n is a positive integer from 1 to 10, x is zero or a positive integer less than or equal to n, and y is zero or a positive integer less than or equal to (2n+1)-2x. The hydroxyalkoxy or saccharide may be C,H,O,O,R or OC(R)xH(OH)H(OH)H(OH)O,R where n is a positive integer from 1 to 10, y is zero or a positive integer less than or equal to 2n+1 and R is independently H, alkyl, hydroxyalkyl, saccharide, C,H,O,R or OC(R)xH(OH)H(OH)H(OH)O,R. In this case, m is a positive integer from 1 to 10, z is zero or a positive integer less than or equal to m, R is H, alkyl, hydroxyalkyl, or C,H,O,R, where m is a positive integer from 1 to 10, z is zero or a positive integer less than or equal to m and m is independently H, alkyl, hydroxyalkyl, or saccharide.

Carboxyamidoalkyl may be alkyl having secondary or tertiary amide linkages or (CH)xCONH,R, O(CH)xCONH,R, (CH)xCONH,R or O(CH)xCONH,R where n is a positive integer from 1 to 10, and R is independently H, alkyl, hydroxyalkyl, saccharide, C,H,O,R or OC(R)xH(OH)H(OH)H(OH)O,R where m is a positive integer from 1 to 10, z is zero or a positive integer less than or equal to (2m+1)-2w, and R is H, alkyl, hydroxyalkyl, or C,H,O,R. In this case, m is a positive integer from 1 to 10, z is zero or a positive integer less than or equal to 2m+1 and R is independently H, alkyl, hydroxyalkyl, or saccharide. In a preferred embodiment, R is an oligonucleotide.

Carboxyalkyl may be alkyl having a carboxyl substituted ether, an amide substituted ether or a tertiary amide removed from an ether or C,H,O,R or OC(R)xH(OH)H(OH)H(OH)O,R where n is a positive integer from 1 to 10, y is zero or a positive integer less than or equal to 2n+1 and R is independently H, alkyl, hydroxyalkyl, saccharide, C,H,O,R or OC(R)xH(OH)H(OH)H(OH)O,R. In this case, m is a positive integer from 1 to 10, z is zero or a positive integer less than or equal to m, R is H, alkyl, hydroxyalkyl, or C,H,O,R, where m is a positive integer from 1 to 10, z is zero or a positive integer less than or equal to m and R is independently H, alkyl, hydroxyalkyl, or saccharide. In a preferred embodiment, R is an oligonucleotide.

Exemplary texaphyrin are listed in Table 1.

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<td>„</td>
<td>„</td>
<td>saccharide</td>
<td>„</td>
</tr>
</tbody>
</table>

Representative Substituents for Texaphyrin Macrocycles.
<table>
<thead>
<tr>
<th>TXP</th>
<th>R₁</th>
<th>R₂</th>
<th>R₃</th>
<th>R₄</th>
<th>R₅</th>
</tr>
</thead>
<tbody>
<tr>
<td>A15</td>
<td>CH₂CH₂CON(CH₂CH₂OH)₂</td>
<td>CH₂CH₃</td>
<td>CH₂CH₃</td>
<td>&quot;</td>
<td>CH₂CON(CH₂CH₂OH)₂</td>
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<tr>
<td>A16</td>
<td>CH₃CH(ON(CH₂CH₂OH) CH₂—</td>
<td>&quot;</td>
<td>&quot;</td>
<td>OCH₃</td>
<td>OCH₃</td>
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<tr>
<td>A17</td>
<td>CH₂(CH₂)₂OH</td>
<td>&quot;</td>
<td>&quot;</td>
<td>O(CH₂)₄COOH, n = 1-7</td>
<td>H</td>
</tr>
<tr>
<td>A18</td>
<td>&quot;</td>
<td>&quot;</td>
<td>&quot;</td>
<td>(CH₂)ₙ—CON-linker-site-directing molecule, n = 1-7</td>
<td>&quot;</td>
</tr>
<tr>
<td>A19</td>
<td>&quot;</td>
<td>&quot;</td>
<td>&quot;</td>
<td>YOCH₃—linker-site-directing molecule - Y = NH, O</td>
<td>&quot;</td>
</tr>
<tr>
<td>A20</td>
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<td>CH₂CH₂CON-</td>
<td>&quot;</td>
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<tr>
<td>A21</td>
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<td>&quot;</td>
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<tr>
<td>A22</td>
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<td>CH₂CH₃</td>
<td>CH₂CH₂CON-</td>
<td>O(CH₂)₄CO-histamine</td>
<td>H</td>
</tr>
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Continued

[0201] Preferred pyrrole-based macrocycles include, but are not limited to the hydro-monomobenzoporphyrins (the so-called “Gp” compounds) disclosed in U.S. Pat. Nos. 4,920,143 and 4,883,790 which are incorporated herein by reference. Typically, these compounds are poorly water-soluble (less than 1 mg/ml) or water-insoluble. Gp is preferably selected from the group consisting of those compounds having one of the formulae A-F set forth in FIGS. 18A-F, mixtures thereof, and the metalated and labeled forms thereof.

[0202] In FIGS. 18A-F, R¹ and R² can be independently selected from the group consisting of carboxalkylox (2-6C), alkyl (1-6C) sulfonyl, aryl (6-10C), sulfonylethyl (6-10C), cyano, and —CONR²CO— wherein R² is aryl (6-10C) or alkyl (1-6C). Preferably, however, each of R¹ and R² is carboxalkylox (2-6C). R³ can be independently carboxalkylox (2-6C) or a salt, amide, ester or acylhydrazo thereof, or is alkyl (1-6C). Preferably R³ is =CH₂COOH or a salt, amide, ester or acylhydrazo thereof.

[0203] R⁴ is =CH₂CH₃ or CHOR² wherein R² is H or alkyl (1-6C), optionally substituted with a hydrophilic substituent: —CHO; —COOR²; CH(OH)₂CH₂; CH₂OCOR²; CH₂OR²; —CH(SR²)CH₂; —CHNR²CH₂; —CHO(NH)₂CH₂; —CH(COOR²)CH₂; —CH(OOCR²)CH₂; —CH(halo)CH₂; —CH(halo)CH₂(halo); an organic group of <12C resulting from direct or indirect derivatization of a vinyl group; or R⁴ consists of 1-3 tetrapyrole-type nuclei of the formula -L-P, wherein -L- is selected from the group consisting of

-CH=CH— and
-CH=C—,

and

[0204] and P is a second Gp, which is one of the formulae A-F (FIG. 18) but lacks R’, or another porphyrin group. When P is another porphyrin group, P preferably has the formula illustrated in FIG. 19: wherein each R is independently H or lower alkyl (1-4C); two of the four bonds shown as unoccupied on adjacent rings are joined to R³; one of the remaining bonds shown as unoccupied is joined to R²; and the other is joined to L₁ with the proviso that, if R³ is =CH₂CH₃, both R³ groups cannot be carboxalkyloxyl. The preparation and use of such compounds is disclosed in U.S. Pat. Nos. 4,920,143 and 4,883,790, which are hereby incorporated by reference.

[0205] Even more preferred for including in liposomes are light activated drugs that are designated as benzoporphyrin derivatives (“BPD’s”). BPD’s are hydrolyzed forms, or partially hydrolyzed forms, of the rearranged products of formula A-C or formula A-D, where one or both of the protected carboxyl groups of R² are hydrolyzed. Particularly preferred is the compound referred to as BPD-MA in FIGS. 20A-D, which has two equally active regiosomers.

[0206] As described above, activating a light activated drug included in a microbubble can enhance rupture of the microbubble. Preferred light activated drugs for including in a microbubble to enhance rupture of the microbubble include Hematoporphyrin, Rose Bengal, Eosin Y, Erythrocruin, Rhodamine B, and PHOTOFRIN. The formulae for these preferred light activated drugs are illustrated in FIGS.
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21A-B where Rose Bengal, Eosin Y, Erythrocyan and Rhodamine B are xanthene derivatives.

[0207] The present invention has a first characteristic in that by adopting ultrasound-sensitive substance-containing drug carriers that can carry a therapeutic drug and transport it to a target location, when said drug carriers carrying drugs are irradiated with ultrasound at the target location, a chemical change or a physical change occurs in said ultrasound-sensitive substance so that said drug carriers rupture to release the drugs.

[0208] In comparison to the method in which the capsule shells are simply ruptured by the action of vibration due to ultrasound in order to release the drugs inside, the rupture of capsules according to the present invention is not as greatly affected by the ultrasound irradiation conditions, so it is possible to adopt a relatively wide range of ultrasound energies, namely 0.1-1000 watts/cm². Therefore, there is no setting of extremely difficult ultrasound irradiation conditions, as the drug carriers can be ruptured and the drugs released efficiently at the target locations within the body even with ultrasound of frequencies other than the resonance frequency.

[0209] In addition, the present invention has a second characteristic in that hollow areas in the drug carriers are formed by shell walls that have a prescribed thickness, and said shell walls contain or are coated with an ultrasound-sensitive substance. To wit, the drug carriers are designed so that while they are sufficiently resistant to pressure and other types of mechanical energy, their structure is such that they are ruptured easily by the chemical changes or physical changes in said ultrasound-sensitive substance.

[0210] Specifically, the drug carrier takes the form of a capsule or the like, and the shell walls that make up the capsule contain or are coated with an ultrasound-sensitive substance in a laminar manner or in lumps. In the case that the aforementioned ultrasound-sensitive substance is disposed in a laminar manner, then the effects of modification of said ultrasound-sensitive substance may extend over the entire aforementioned shell walls. On the other hand, in the case that the aforementioned ultrasound-sensitive substance is disposed in lumps, the said ultrasound-sensitive substance is present locally in said shell walls, making it possible for said shell walls to be ruptured reliably. Note that a uniform mixture of the ultrasound-sensitive substance in a substrate to be described later may be used as the substance that makes up the drug carrier itself. In this case, not only will the shell walls be ruptured uniformly, but the fabrication of the drug carrier will be simplified.

[0211] Moreover, the present invention has a third characteristic in that a drug is carried in the hollow areas formed by the shell walls, but a prescribed amount of gas may also be present in said hollow areas together with said drug. As described later, the type and amount of said gas is arbitrary, but this should be preferably set in the range of 0.01-50% of the volume of said hollow areas. There follows a description of the composition and structure of the drug carriers.

[0212] Here, a “drug carrier” is defined to be a carrier that can carry a therapeutic drug and transport it to a target location. While its form is not particularly limited, in consideration of the case of manufacture, manufacturing costs and other considerations, a capsule form having a hollow area which is isolated from the outside by a shell wall is preferable.

[0213] The size of the drug carrier is normally set appropriately in the range of 0.01-100 μm. If less than 0.01 μm, then it will be excreted outside of the body and be inadequately effective. If greater than 100 μm, then there is a risk of interfering with the flow of blood within the blood vessels.

[0214] The substrate material that is used for the drug carrier may be one of various biocompatible polymers, albumins, liposomes, sugars or other substances.

[0215] In addition, by using a drug carrier that has been modified to the prodrug form, the selective movement to the locations of tagged tissue can be improved, aqueous solubility can be increased, absorption can be promoted and side effects can be lessened. In this case, the prodrug reverts to the original drug carrier in an enzymatic or non-enzymatic manner in the body after it reaches the target of the aforementioned modification, so it is possible for the sensitivity to ultrasound to be restored. A drug carrier that has been modified to the prodrug form is included in the scope of the present invention. Note that it is also appropriate for the drug carried by the drug carrier to be modified to the prodrug form.

[0216] An ultrasound-sensitive substance is defined to be a substance that is activated by ultrasound of a prescribed frequency and intensity by means of a mechanism to be described later, and which causes chemical changes to occur in itself or another substance, or changes its own structure or causes some sort of changes to occur. Examples of ultrasound-sensitive substances include fluorescein, merocyanin, and the like, but from the standpoint of toxicity and affinity with respect to the body, porphyrin derivatives or xanthene derivatives are preferable. Specific examples of the aforementioned porphyrin derivatives or xanthene derivatives are discussed in more detail later.

[0217] Moreover, with regard to the structure of the drug carrier, as shown in FIG. 22, lumps of the drug 108 and ultrasound-sensitive substance 110 may be distributed at appropriate intervals within a drug carrier 106 of indefinite shape, but it is preferable to shape the drug carrier itself so that it has shell walls having a prescribed thickness to form hollow areas therein, and have said shell walls contain or be covered with the aforementioned ultrasound-sensitive substance to permit said drug carrier to be ruptured efficiently.

[0218] If the aforementioned hollow areas are formed on the interior of the drug carrier, there are no particular limits to the numbers thereof, as there may be one or more. In addition, there are no particular limits to their shapes or locations, but they are preferably formed in the surface layers of the drug carrier in order for the drug release to be performed well. Note that making the shape of the drug carrier itself as that of a capsule consisting of layers of shell walls that have the aforementioned prescribed thickness is also contained in the scope of the present invention.

[0219] Here, the thickness of the aforementioned shell walls is normally determined within the range of 0.001-50 μm. If the thickness of the shell walls is less than 0.001 μm, then the shell walls become easily ruptured by shock and so there is a risk of the aforementioned shell walls being
ruptured and drug inside leaking out before the target location within the body is reached. On the other hand, if 50 \( \mu \text{m} \) or thicker, rupturing the aforementioned shell walls becomes difficult even with the action of the ultrasound-sensitive substance, and even if a portion of the shell walls is ruptured, there is a risk that unrupturable shell wall portions remain, blocking the release of drugs.

[0220] At the time that an ultrasound-sensitive substance is contained in the drug carrier, if said drug carrier is equipped with a shell wall structure, it is preferable that the aforementioned ultrasound-sensitive substance have said shell walls contain or be covered with the aforementioned ultrasound-sensitive substance.[0221] Specifically, the outside surface of shell wall 112 may be covered with a layer of the ultrasound-sensitive substance 110 as shown in FIG. 23(a), or the inside surface of the shell wall 112 may be coated with a layer of the substance as shown in FIG. 23(b), or a layer of the substance may be present entirely within the interior of shell wall 112 in a laminar manner as shown in FIG. 23(c). Note that the layer of the ultrasound-sensitive substance 110 need not have a continuous layer structure as shown in FIG. 23, and moreover the layer need not be concentric with the shell wall 112 as shown in FIG. 23.

[0222] In addition, the shell wall 112 may contain or be covered with the ultrasound-sensitive substance 110 in a manner such that it is distributed in lumps. To wit, as shown in FIGS. 24(a) and 24(b), lumps of the ultrasound-sensitive substance 110 may be adhered to the outside surface of inside surface of said shell wall 112, or as shown in FIG. 24(c), lumps may be distributed appropriately within the interior of said shell wall 112. In this case, portions of the individual lumps may protrude from the shell wall 112 or they need not protrude.

[0223] Moreover, as shown in FIG. 24(d), the lumps of the ultrasound-sensitive substance 110 may penetrate through the shell wall 112 in a structure such that portions of the lumps are exposed to the outside and the interior space.

[0224] In the cases shown in FIG. 24, lumps of the ultrasound-sensitive substance 110 may be distributed uniformly within the aforementioned shell wall 112, or the lumps may be distributed non-uniformly. In addition, there are no particular limitations to the shapes of the individual lumps, as shapes other than spherical are also possible.

[0225] Note that in FIGS. 23 and 24, the numeral 114 indicates the hollow area formed by the shell wall 112 so that various types of drugs are carried in this hollow area.

[0226] The method of using the drug carrier described above will now be explained with reference to FIG. 25.

[0227] A drug of a prescribed type carried in the aforementioned drug carrier is administered orally or administered hypodermically using a syringe or other drug administration apparatus, or using a special method, injected into the body by means of the iontophoresis method using ionized drug carriers.

[0228] Moreover, in the case that the use of a catheter or endoscope is possible, a therapeutic ultrasound generator is attached to the tip of a catheter or endoscope and introduced into the interior of the body of the patient and allowed to reach the affected area.

[0229] FIGS. 25(a) and 25(b) are both sectional drawings showing the method of attaching the ultrasound generator used in the working of the present invention, where FIG. 25(a) shows the structure in the case in which the ultrasound generator is attached to the tip of an endoscope, while FIG. 25(b) shows the structure in the case in which it is attached to the tip of a catheter.

[0230] In the manner shown in FIG. 25(a), a miniature center tube 118 containing an optical fiber (not shown) and wiring used to operate ultrasonic oscillators (to be described later) is laid in the interior of a fine tube 116 constituting the endoscope. A cylindrical first ultrasonic oscillator 124 and second ultrasonic oscillator 126 are disposed concentrically on the tip of the fine tube 116. Examples of the ultrasonic oscillators include one made by mounting electrodes on either side of a piezoelectric element, and in this case, ultrasound is generated by applying an electrical signal with an ultrasonic frequency between said electrodes. Numerical 122 is a core used to transmit an image of the outside to the optical fiber (not shown) embedded in the aforementioned hollow area. In addition, the gap between the fine tube 116 and the miniature center tube 118 becomes the drug delivery path 120 connected to penetration holes 128 opened at appropriate intervals in the circumferential surface of the tip side of the fine tube 116.

[0231] Moreover, the respective frequency characteristics of the aforementioned first ultrasonic oscillator 124 and second ultrasonic oscillator 126 are different, so by controlling the operation of both of these, a mixture of two different frequencies can be generated in a direction (the direction indicated by arrows) perpendicular to the axial direction of the endoscope. Radiating a combination of different ultrasound frequencies in this manner is done because a more complex ultrasound waveform improves the efficiency of rupturing the drug carriers. Note that the first and second ultrasonic oscillators having the structure described above may also be mounted on the tip of a catheter.

[0232] On the other hand, the first ultrasonic oscillator 124 and second ultrasonic oscillator 126 with different frequency characteristics may both have a solid cylindrical shape and be laminated in the direction of the axis of the catheter in the manner shown in FIG. 25(b).

[0233] Therefore, by controlling the operation of both of these, ultrasound having two different frequencies can be generated in the axial direction of the endoscope (the direction indicated by arrows).

[0234] Note that the first and second ultrasonic oscillators having the structure described above may also be mounted on the tip of an endoscope.

[0235] Now, after confirming that the tip of the endoscope or catheter has reached the affected tissue, the drug carriers are released toward the affected part along the drug delivery path 120 through the penetration holes 128. At the same time or after a prescribed amount of time has elapsed, the first and second ultrasonic oscillators are operated thereby irradiating the affected area with ultrasound of different frequencies, thus rupturing the drug carriers present in the affected tissue. Therefore, the drug within the drug carrier is administered over a range limited to the vicinity of the location of the affected tissue.

[0236] Considering the location of the affected area relative to the aforementioned endoscope or catheter, it is...
preferable to select the equipment such that suitable irradiation of ultrasound is obtained. In addition, the diameter of the aforementioned endoscope or catheter may be selected appropriately as one in the range from 1 mm to 5 cm.

Note that three or more types of the aforementioned ultrasonic oscillators may also be mounted. In this case, an even more complex ultrasound waveform can be generated, so the efficiency of rupturing the drug carriers can be increased even further. A single ultrasonic oscillator may be used, for example, if the drug carriers have a shape that is easily ruptured.

On the other hand, in the event that a catheter or endoscope cannot be used, as shown in FIG. 26, a first ultrasonic oscillator 124 and a second ultrasonic oscillator 126 having the same characteristics as those described above may be laminated and disposed in a hollow area of a base 130 consisting of flexible synthetic resin or the like. One or a plurality of such therapeutic ultrasound generators are placed upon the skin in areas corresponding to the affected areas and irradiate ultrasound toward the affected areas (in the direction indicated by arrows). As shown in the figure; it is preferable to provide a plurality of laminates of ultrasonic oscillators in the base 130. The base 130 can flex to match the shape of the patient’s body so the ultrasound can be concentrated in the affected area. The diameter of the aforementioned first and second ultrasonic oscillators is normally set appropriately in the range of 5-10 cm. Note that it is preferable for the aforementioned first and second ultrasonic oscillators to be made of flexible oscillating material such as a fluorine compound in order to maintain the overall flexibility of the apparatus and keep its tight fit to the skin or the like.

Moreover, the ultrasound energy concentrated in the affected area ruptures the drug carriers present there and releases the drug into the interior of the affected area.

In this manner, according to the present invention, by including an ultrasound-sensitive substance in the drug carrier, a drug carrier that is difficult to rupture with ultrasound only can be easily ruptured and the release of the drug into a desired location in the body can be controlled.

Note that in the case that substances such as antibodies that selectively bind to cancerous cells, thrombi, organs, blood vessels, hardened arteries or the like are included in the substrate material of the drug carrier, the drug carriers can be concentrated in the tissue of the affected area, so it is possible to use ultrasound to rupture the drug carriers concentrated in said tissue of the affected area to administer the drug locally in high concentration.

Here follows a description of the conditions for the ultrasound used in the present invention.

The power of ultrasound irradiated in order to rupture the drug carriers is set appropriately in the range of 0.1-1000 watts/cm². If the power of ultrasound is below 0.1 W/cm², there is insufficient energy to activate the ultrasound-sensitive substance, and above 1000 W/cm², excessive amounts of heat are generated, causing damage to the body.

In addition, the frequency of the ultrasound is set appropriately in the range 10 kHz-100 MHz, but the range 20 kHz-10 MHz is particularly preferable. With ultrasound in this frequency band, a relatively low energy is able to cause the cavitation to be described later and efficiently rupture the drug carriers.

Thus, by combining multiple frequencies as described above, it is possible to activate the ultrasound-sensitive substance and efficiently rupture the drug carriers. For example, during irradiation with ultrasound of a fixed frequency, intermittently switching between this and a different frequency can be performed intentionally to enhance or suppress the generation of the cavitation to be described later and thus rupture or disintegrate the drug carriers.

Thus, by combining multiple frequencies as described above, it is possible to activate the ultrasound-sensitive substance and efficiently rupture the drug carriers. For example, during irradiation with ultrasound of a fixed frequency, intermittently switching between this and a different frequency can be performed intentionally to enhance or suppress the generation of the cavitation to be described later and thus rupture or disintegrate the drug carriers.

To present an actual example, while the affected area is continuously irradiated with ultrasound at 100 kHz, switching to a frequency of 270 kHz in short (0.001-10 sec) pulses have even better effectiveness of rupturing. The same effect is expected if the frequency of the ultrasound is continuously varied within a fixed range. This phenomenon is thought to increase the rupture forces by temporarily halting the resonance movement of the drug carrier.

Here follows an explanation of the mechanism of rupturing of the drug carriers.


Cavitation is a phenomenon in which gas dissolved in an aqueous solution forms bubbles under certain types of acoustic vibration, or in which extremely small bubbles already present oscillate or repeatedly enlarge and contract, becoming bubbles. Then, when the size of these cavitation bubbles reaches a size that cannot be maintained, they collapse and this collapse occurs suddenly so it is known that various types of energy occur locally at this time.

To wit, at the time of the aforementioned cavitation collapse, hot spots of 6000-7000 degrees are formed at the center, so in addition to vibration and other types of mechanical energy, visible light, ultraviolet light and other types of electromagnetic radiation, heat, plasma, magnetic fields, shock waves, free radicals, heat and other forms of energy are thought to be generated locally.

The ultrasound-sensitive substance according to the present invention is activated by the aforementioned various forms of energy generated at the time of cavitation collapse, and also chemical changes are thought to occur along with structural changes.
For example, one of the ultrasound-sensitive substances according to the present invention, Rose Bengal, is excited and activated by light of wavelength 530 nm or ultraviolet light. Therefore, the activation of Rose Bengal is thought to be caused by the ultraviolet light generated at the time of cavitation collapse.

The threshold value of ultrasonic energy for cavitation to occur is known to become lower in a liquid in which an ultrasound-sensitive substance is present. Therefore, the ultrasound-sensitive substance contained in the drug carrier exhibits the effects of selectively inducing the occurrence of cavitation in the vicinity of the drug carrier, and also itself being activated by the energy generated by the collapse of cavitation to rupture the drug carrier.

On the other hand, it is known that the threshold of ultrasonic energy in order to generate cavitation also becomes lower if minute bubbles are present in the liquid. Therefore, cavitation can be effectively caused to occur during ultrasound irradiation by causing a prescribed amount of gas to be present within the drug carrier, and the energy generated at the time of this cavitation collapse can be used effectively to rupture the drug carrier. The type and amount of said gas is arbitrary, but the amount is preferably set in the range of 0.01-50% of the volume of the hollow area, then the generation of cavitation cannot be effectively induced, and if the amount of same gas exceeds 50% of the volume of said hollow area, the drug carrier may not have sufficient strength to reach the target location, and the amount of drug transported is limited.

Note that the visible light, ultraviolet light and other types of electromagnetic radiation, heat, plasma, magnetic fields, shock waves, free radicals, heat and other forms of energy generated at the time of collapse of cavitation can be used directly in the treatment of affected tissue.

For example, by generating ultrasound in the vicinity of the affected tissue, ultraviolet light that is normally absorbed by the skin and does not reach the interior of the body can be generated in the vicinity of affected tissue within the body by means of the collapse of cavitation derived from ultrasound, so it is possible to treat affected areas by its disinfectant action.

To wit, cavitation can be induced within the body by means of ultrasound and the energy generated at the time of the collapse can be used in the treatment of affected areas. By means of this method, ultraviolet light and other forms of energy can be generated at will at any location within the body to perform treatment of affected areas, so there would be no need to consider the side effects specific to drug therapy.

The present invention can be worked in the manner described below:

Cancer Therapy

The anti-cancer drug cisplatin is enclosed in a polymer capsule, and then the outside surface of this capsule is coated with an ultrasound-sensitive substance and the resulting capsule is injected arterially. Since the cisplatin is covered with a polymer, these capsules have no toxicity (side effects) even when injected arterially, as they merely flow along in the blood.

However, when these capsules flow along inside the blood vessels inside cancerous tissue and are irradiated with ultrasound, the ultrasound-sensitive substance on the surface of the capsule is activated so the capsule enclosing the cisplatin disintegrates and the cisplatin is released inside this tissue in high concentrations. Therefore, the administration of anti-cancer drugs in high concentrations is possible in limited areas where cancer is located, yet normal cellular tissue can be spared the strong toxicity of cisplatin.

This method is particularly effective in cancer of the liver, brain tumors and other diseases in which large numbers of blood vessels are present. The method of irradiation with ultrasound may be irradiation of the tumor portions from the surface of the skin, or irradiation of the cancerous tissue with ultrasound directly during a laparotomy.

In addition, it is also possible to attach ultrasound generators to an endoscope and irradiate the interior of the body with ultrasound, and in this case, the interior of the stomach and colon cancers and the like can be directly irradiated with ultrasound from bodily cavities in the inside of the colon.

The route of administration of these capsules may be arterial injection or absorption from the intestines after oral administration, direct injection into the affected area via lymphatic vessels and other routes. The appropriate route can be selected depending on the case.

However, if Photofrin® is used as the aforementioned ultrasound-sensitive substance, since Photofrin® itself has an affinity for cancer cells, the aforementioned cisplatin-containing capsules concentrate in the cancerous tissue and accumulate in high concentration.

Moreover, cisplatin can be administered in cancerous tissue at even higher concentrations by irradiating with ultrasound in this state. Note that when Photofrin® is activated by ultrasound, it also has the character of causing a cell-kill action, so in this case, the anti-cancer action is augmented in a synergistic manner with cisplatin.

In addition, in the case of cancer that has metastasized within the abdomen or bladder cancer, a conceivable therapy method would be to inject capsules containing anticancer drugs directly into the interior of the abdomen and then irradiate the entire abdomen with ultrasound from the surface of the skin. In the case of bladder cancer, the interior of the bladder can be filled with these capsules from the urethra and the bladder cancer can be treated by irradiating the lower abdomen with ultrasound from the surface of the skin. In these cases, there is an added advantage in that the state of release of drugs can be observed using diagnostic ultrasound equipment during irradiation with therapeutic ultrasound.

Thrombolytic Therapy

Thrombolytic agents are used as drugs for the treatment of myocardial infarction and cerebral infarction. However, if large doses of drugs are administered in order to dissolve the thrombi as fast as possible, there is a risk of the blood conversely becoming less easily clotted, resulting in excessive hemorrhaging.

To solve this problem, urokinase or another thrombolytic agent is enclosed in capsules made of biocompatible
polymer or albumin which contains an ultrasound-sensitive substance and the resulting capsules are injected into a blood vessel. Since these capsules are not ruptured in the normal state, they do not induce thrombolytic action within the blood vessels.

[0272] In the case of a myocardial infarction, for example, when these capsules reach the peripheral blood vessels of the coronary artery or other locations where thrombi are present, an apparatus such as that described above can be used to irradiate these locations with ultrasound from outside the body or inside the body, thus rupturing the capsules and releasing the drug locally in high concentrations.

[0273] Specifically, as shown in FIG. 27, the catheter or endoscope 132 equipped with ultrasound generators as shown in FIG. 25 is inserted into the interior of the blood vessel 134 up until near the thrombus 136 and the thrombolysis agent-containing capsules 138 are released immediately upstream of the thrombus 136 while ultrasound is simultaneously generated. These capsules are ruptured by the action of the ultrasound-sensitive substance which is activated by ultrasound, so the thrombolysis agent inside is released to the location of the thrombus.

[0274] Note that the capsules that were not ruptured by ultrasound the first time are ruptured and used when they again flow back to the affected areas. These capsules are not ruptured during the period before they again reach the affected area so they will not become the cause of hemorrhaging.

[0275] In addition, if a substance or antibody with particular affinity to thrombi is attached to the outside surface of the capsules, these capsules will collect at a thrombus in high concentrations. If the location of the thrombus is irradiated with ultrasound in this state, these capsules can be ruptured to administer the thrombolysis agent effectively in the vicinity of the thrombus. A substance that has particular affinity to thrombi has been reported by Lanza, et al. (Circulation, 1995, 92, Suppl. I: 1-260). By doing such, thrombolytic therapy can be performed efficiently without causing hemorrhaging or other side effects.

[0276] Blood Vessel Therapy

[0277] In the case that angioinvasion due to arteriosclerosis or the like occurs, a surgical treatment in which a balloon catheter is used to enlarge the cavity within the blood vessel to reopen the flow of blood has become common in recent years. In addition, a procedure in which a metal stent is used to fix the cavity within the blood vessel in the enlarged state so that the narrowing of blood vessels does not recur after the above surgery has also become common. However, in either case, the inside walls of the blood vessel are damaged to a certain extent. In order to restore the damaged blood vessel tissue to the original state of the blood vessel, repair, including the grafting of blood vessel tissue is performed, but excessive repair in the repair process occurs in more than 50% of cases, leading to recurrence of angioinvasion. This is a drawback of this therapy.

[0278] To solve this problem, as shown in FIG. 28, drug carriers 140 that carry Photofrin® internally are mixed into the material for the balloon 146, and a balloon catheter with an ultrasound generator 132 positioned in its center 142 is inserted into the blood vessel 144. The balloon 146 is inflated at the target location and brought into close contact with the blood vessel walls 144. When ultrasound is generated in the direction perpendicular to the axis of the catheter, the drug carriers 140 located in areas of the balloon 146 in contact with the blood vessel walls 144 are ruptured and the Photofrin® inside is injected directly into the blood vessel walls 144.

[0279] When ultrasound is generated in the direction perpendicular to the axis of the catheter, the drug carriers located in areas of the balloon in contact with the blood vessel walls are ruptured and the Photofrin® inside is injected directly into the blood vessel walls. When activated with ultrasound, Photofrin® has the character of blocking the blood vessel tissue repair process to a certain degree so it is possible to suppress excessive repair of the blood vessel walls that were damaged by balloon, so that recurrence of angioinvasion can be prevented.

[0280] In this embodiment, Photofrin® is carried within the drug carrier as a drug for treatment of affected areas, and the ultrasound-sensitive substance contained in the drug carriers used to rupture these carriers may be selected arbitrarily, but Photofrin® may also be used as this ultrasound-sensitive substance.

[0281] Note that examples of drugs for treatment of affected areas that can be used in blood vessel therapy include genes, heparin, radioactive substances and the like which are administered to the inside wall of the blood vessels.

[0282] Use as a Hemostatic Agent

[0283] Current therapies for liver cancer include injecting ethanol into the blood vessels that supply nutrition to the cancerous tissue, causing damage to the inside walls of these blood vessels and thus artificially create thrombi that block these blood vessels, injecting a specific type of fluid to block the blood vessels and other methods to prevent the propagation of cancerous cells.

[0284] An alternative method involves preparing Rose Bengal-containing drug carriers that exhibit action as a blood vessel inside wall damaging agent when activated by ultrasound, and also preparing drug carriers that contain thrombin which acts as a blood coagulating agent, and injecting both of these simultaneously into the blood vessel and irradiating the affected area with ultrasound.

[0285] In this embodiment, Rose Bengal is carried within the drug carriers as the drug for treatment of affected areas. Moreover, the ultrasound-sensitive substance contained in the drug carriers used to rupture these carriers may be selected arbitrarily, but Rose Bengal may also be used.

[0286] In this method, the drug carriers are ruptured by ultrasound to release the Rose Bengal inside into the affected area, and at this time, the Rose Bengal itself is activated by ultrasound and damages the blood vessel walls to form thrombi. At the same time the thrombin released from individual drug carriers causes the blood to coagulate. Therefore, blood flow is halted in the affected area. By combining these two types of drug carriers in this manner, a synergistic effect as a hemostatic agent is obtained.

[0287] The method described above can be applied not only to the treatment of liver cancer but also to the stanching of hemorrhage from organs due to traffic accidents and the like.
Transdermal Administration of Medication

The transdermal administration of medication with the additional use of ultrasound is already known. For example, as recited in the specification of Japanese Patent Application No. 9-166334, a treatment apparatus consisting of a disc-shaped plate which has a large number of fine permeation holes and which contains fluid in the interior was developed. In this apparatus, the cavitation generation phenomenon can be used to open fine holes in the surface of the skin, so the administration of medication or collection of bodily fluids can be performed effectively without any accompanying pain.

In this embodiment, drug carriers containing an ultrasound-sensitive substance are disposed in the permeation holes in the treatment apparatus described above. Here follows a description in reference to FIG. 29.

In this preferred embodiment, the transdermal medication administration apparatus 140 consists of a film of synthetic resin material or the like which is relatively thin with a thickness in the range of 1 µm-1 cm, where a circular space 142 is formed in the interior.

Formed in the bottom-side film 144 of the transdermal administration apparatus 140 is a plurality of permeation holes 146 that connect the circular space 142 to the outside. The diameter of the permeation holes 144 may be set in the range from 0.1 µm to 3 mm. Note that in the example shown in FIG. 29, the permeation holes 146 are distributed uniformly, but they may be provided with non-uniform density if necessary. In addition, the sectional shape of the permeation holes 146 is not limited to circular, as they may also be star-shaped, polygonal-shaped or irregularly shaped. The density of permeation holes 146 can be set in the range from 1 to 1 million per square centimeter.

An ultrasonic oscillator 150 is attached to the top-side film 148 of the transdermal administration apparatus 140. This ultrasonic oscillator 150 may be formed as a unit with the transdermal administration apparatus 140, or a separate ultrasonic oscillator may be prepared independent of the apparatus and pressed against the top-side film 148 of the transdermal administration apparatus 140.

When the transdermal medication administration apparatus 140 is in use, drug carriers 152 are disposed within the permeation holes 146. The circular space 142 and carrying space 154 in the interior of the drug carriers 152 are filled with liquid medication 156.

The bottom-side film 144 of the transdermal administration apparatus 140 is pressed against the surface of the skin and a driving signal is supplied to the ultrasonic oscillator 150 to generate ultrasound. Upon doing this, the drug carriers 152 are ruptured and the permeation holes 146 are opened and at the same time, cavitation occurs in the liquid medication 156 within the circular space 142, so a fast flow of fluid that occurs at the time of cavitation collapse passes through the permeation holes 146 to reach the skin, forming fine holes in its surface. The liquid medication 156 passes through these holes and is absorbed into the body.

In this manner, the liquid medicine does not flow out from the permeation holes 146 while the transdermal administration apparatus 140 is in storage, even when the permeation holes 146 are relatively large. On the other hand, the drug carriers 152 are ruptured by ultrasound so the permeation holes 146 can be opened reliably at the time of use.

In addition, the ultrasound-sensitive substance inside the drug carriers 152 lowers the threshold value for the occurrence of cavitation, so cavitation can be caused to occur in the liquid medication 156 at a low ultrasonic energy. Thereby the ultrasonic energy absorbed by the skin can be reduced and the risk of causing deleterious effects on the skin is reduced.

The medications and substances that can be administered transdermally using this apparatus include anti-allergic drugs, insulin, various hormones, anti-cancer drugs, anti-inflammatory drugs, anesthetics, anticoagulant factors (heparin, urokinase), antibiotics, various vitamins, steroids, antihypertensive agents, vasopressor drugs, tranquilizers, hair restoratives depilatories and others.

Treatment of Infectious Disease

Although the disinfectant effects of UV light are well known, the ability of UV light to pass through liquids is extremely poor, so its intensity attenuates immediately, and thus it is used exclusively for sterilizing the surfaces of medical equipment and the like.

Now the ultrasound-sensitive substance Rose Bengal is known also to have the action of reducing the threshold for generation of cavitation by ultrasound. Taking advantage of this characteristic, ultrasound can be used for the treatment of infectious disease both inside and outside the body, since ultrasound can be used for the treatment of infectious disease inside the body.

To wit, in the treatment of infectious disease inside the body, carriers containing Rose Bengal are infused deeply into the affected area by injection or other method, and when the affected area is irradiated with ultrasound in this state, cavitation is generated in the vicinity of these carriers with a relatively low ultrasonic energy, and UV light is generated at the time of its collapse as described previously.

Therefore, affected areas within the body can be disinfected by irradiating the affected areas with UV light from a very close distance, so this method can be applied to the treatment of infectious disease. In addition, this method has an additional advantage in that there is no need to use various types of antibiotics, so no drug-resistant bacteria are created.

Next, in the case of an infectious disease of the skin, drug carriers that carry a skin absorption accelerant and have their surface covered with Rose Bengal are applied to the surface of the skin. Rose Bengal permeates the skin relatively easily so the drug carriers permeate slightly into the surface portion of the skin. If the skin is irradiated with ultrasound in this state, the skin absorption accelerant is released into the skin and the skin's barrier function is lowered or vanishes, so insulin or other drugs that are normally not easily absorbed by the skin are absorbed into the skin.

Note that the above method can be applied not only to the treatment of infectious diseases of the skin, but also to the treatment of athlete's foot, viral blisters, psoriasis scabies, skin cancer, AIDS-related Kaposi's sarcoma and the like.
The treatment of diabetes can be performed by intravenously injecting drug carriers containing insulin and irradiating the interior of the body with ultrasound when needed so that these drug carriers are ruptured to release the insulin inside into the body. In this case, by adjusting the time and intensity of the irradiation of ultrasound, insulin can be administered regularly with a simple operation.

In addition, it is possible to use red blood cells in the blood as the drug carriers described above. For example, red blood cells can be separated from blood and insulin injected into the individual red blood cells and then the ultrasound-sensitive substance Photofrin® made to adhere to the surface of the red blood cell membranes.

By supplying red blood cells treated in this manner into the body of the patient through a transfusion or the like, the red blood cells have a lifetime of approximately 10-100 days, so they will not be ruptured during this period unless radiated with ultrasound, but the insulin can be released when necessary by irradiation with ultrasound from outside the body. In this case, since the drug carriers consist of materials that are very compatible with the body, namely red blood cells, rejection from the body can be suppressed.

As discussed above, the light activated drug can be coupled with a site directing molecule to form a light activated drug conjugate. Suitable site-directing molecules include, but are not limited to: polyoxyethylorcouleotides, oligodeoxycytidinucleotides, polyribonucleotide analogs, oligoribonucleotide analogs; polyamides including peptides having an affinity for a biological receptor and proteins such as antibodies; steroids and steroid derivatives; hormones such as estradiol or histamine; hormone mimics such as morphine and further macrocycles such as sapphirins and rubyrins. It is understood that the terms “nucleotide”, “poly-nucleotide”, and “oligonucleotide”, as used herein and in the appended claims, refer to both naturally occurring and synthetic nucleotides, poly- and oligonucleotides and to analogs and derivatives thereof such as methylphosphonates, phosphothriesters, phosphorothioates, and phosphoramidates and the like. Deoxycytidinucleotides and ribonucleotide analogs are contemplated as site-directing molecules.

When the site-directing molecule is an oligonucleotide, the oligonucleotide may be derivatized at the bases, the sugars, the end of the chains, or at the phosphate groups of the backbone to promote in vivo stability. Modifications of the phosphate groups are preferred in one embodiment since phosphate linkages are sensitive to nuclease activity. Preferred derivatives are the methylphosphonates, phosphothriesters, phosphorothioates, and phosphoramidates. Additionally, the phosphate linkages may be completely substituted with non-phosphate linkages such as amide linkages. Appendages to the ends of the oligonucleotide chains also provide exonuclease resistance. Sugar modifications may include alkyl groups attached to an oxygen of a ribose moiety in a ribonucleotide. In particular, the alkyl group is preferably a methyl group and the methyl group is attached to the 2’ oxygen of the ribose. Other alkyl groups may be ethyl or propyl.

A linker may be used to couple the light activated drug with the site directing molecule. Exemplary linkers include, but are not limited to, amides, amine, thioether, ether, or phosphate covalent bonds as described in the examples for attachment of oligonucleotides. In a preferred embodiment, an oligonucleotide or other site-directing molecules is covalently bonded to a texaphyrin or other light activated drugs via a carbon-nitrogen, carbon-sulfur, or a carbon-oxygen bond.

As described above, the media can be an emulsion which includes a light activated drug. The emulsions described below are suitable for delivery into a body since they avoid pharmaceutically undesirable organic solvents, solubilizers, oils or emulsifiers. A wide range of light activated drug concentrations can be used in the emulsion. Suitable concentrations of light activated drug within the emulsion include, but are not limited to, approximately 0.01 to 1 gram/100 ml, preferably about 0.05 to about 0.5 gram/100 ml, and approximately 0.1 g/100 ml.

The emulsion includes a lipoid as a hydrophobic component dispersed in a hydrophilic phase. The hydrophobic component of the emulsion comprises a pharmaceutically acceptable triglyceride, such as an oil or fat of a vegetable or animal nature, and preferably is selected from the group consisting of soybean oil, safflower oil, marine oil, black current seed oil, borage oil, palm kernel oil, cotton seed oil, corn oil, sunflower seed oil, olive oil or coconut oil. Physical mixtures of oils and/or interesterified mixtures can be employed. The preferred oils are medium chain length triglycerides having C12-C17 chain length and more preferably a saturated. The preferred triglyceride is a distillate obtained from coconut oil. The hydrophobic content of the emulsion is preferably approximately 5 to 50 g/100 ml, more preferably about 10 to about 30 g/100 ml and approximately 20 g/100 ml of the emulsion.

The emulsion can also contain a stabilizer such as phosphatidic, soybean phospholipids, nonionic block copolymers of polyox yethylene and polyoxpropylene (e.g., poloxamers), synthetic or semi-synthetic phospholipids, and the like. The preferred stabilizer is purified egg yolk phospholipid. The stabilizer is usually present in the composition in amounts of about 0.1 to about 10, and preferably about 0.3 to about 3 grams/100 ml, a typical example being about 1.5 grams/100 ml.

The emulsion can also include one or more bile acids salts as a costabilizer. The salts are pharmacologically acceptable salts of bile acids selected from the group of cholic acid, deoxycholic acid and glycodeoxycholic acid, and preferably of cholic acid. The salts are typically alkaline metal or alkaline earth metal salts and preferably sodium, potassium, calcium or magnesium salts, and most preferably, sodium salts. Mixtures of bile acid salts can be employed if desired. The amount of bile acid salt employed is usually about 0.01 to about 1.0 and preferably about 0.05 to about 0.4 grams/100 ml, a typical example being about 0.2 grams/100 ml.

Suitable pH for the emulsion includes, but is not limited to approximately 7.5 to 9.5, and preferably approximately 8.5. The pH can be adjusted to the desired value, if necessary, by adding a pharmaceutically acceptable base, such as sodium hydroxide, potassium hydroxide, calcium hydroxide, magnesium hydroxide and ammonium hydroxide.

Water can be added to the emulsion to achieve the desired concentration of various components within the
emulsion. Further, the emulsion can include auxiliary ingredients for regulating the osmotic pressure to make the emulsion isotonic with the blood. Suitable auxiliary ingredients include, but are not limited to, auxiliary surfactants, isotonic agents, antioxidants, nutritive agents, trace elements and vitamins. Suitable isotonic agents include, but are not limited to, glycerin, amino acids, such as alanine, histidine, glycine, and/or sugar alcohols, such as xylitol, sorbitol and/or mannitol. Suitable concentrations for isotonic agents within the emulsions include, but are not limited to, approximately 0.2 to about 8.0 grams/100 ml and preferably about 0.4 to about 4 grams/100 ml and most preferably 1.5 to 2.5 grams/100 ml.

[0319] Antioxidants can be used to enhance the stability of the emulsion, a typical example being α-tocopherol. Suitable concentrations for the antioxidants include, but are not limited to, approximately 0.005 to 0.5 grams/100 ml, approximately 0.02 to about 0.2 grams/100 ml and most preferably approximately 0.05 to 0.15 grams/100 ml.

[0320] The emulsions can also contain auxiliary solvents, such as an alcohol, such as ethyl alcohol or benzyl alcohol, with ethyl alcohol being preferred. When employed, such is typically present in amounts of about 0.1 to about 4.0, and preferably about 0.2 to about 2 grams/100 ml, a typical example being about 1 gram/100 ml. The ethanol is advantageous since it facilitates dissolution of poorly water-soluble light activated drugs and especially those that form crystals which may be very difficult to dissolve in the hydrophobic phase. Accordingly, the ethanol must be added directly to the hydrophobic phase during preparation to be effective. For maximum effectiveness, the ethanol should constitute about 5% to 15% by weight of the hydrophobic phase. In particular, if ethanol constitutes less than 5% by weight of the hydrophobic phase, dissolution of the light activated drug can become unacceptably slow. When the ethanol concentration exceeds 15%, large (>5 μm diameter) and poorly emulsified oil droplets can form in the emulsion. The particles in the emulsion are preferably less than about 5.0 μm in diameter, more preferably less than 2.0 μm in diameter and most preferably less than 0.5 μm or below.

[0321] A typical emulsion is prepared using the following technique. The triglyceride oil is heated to 50°-70° C. while sparging with nitrogen gas. The required amounts of stabilizer (e.g., egg yolk phospholipids), bile acid salt, alcohol (e.g., ethanol), antioxidant (e.g., α-tocopherol) and light activated drug are added to the triglyceride while processing for about 5 to about 20 minutes with a high speed blender or overhead mixer to ensure complete dissolution or uniform suspension.

[0322] In a separate vessel, the required amounts of water and isotonic agent (e.g., -glycerin) are heated to the above temperature (e.g., 50°-70°) while sparging with nitrogen gas. Next, the aqueous phase is transferred into the prepared hydrophobic phase and high speed blending is continued for another 5 to 10 minutes to produce a uniform but coarse premulsion (or premix). This premix is then transferred to a conventional high pressure homogenizer (APV Gaulin) for emulsification at about 8,000-10,000 psi. The diameter of the dispersed oil droplets in the finished emulsion will be less than 5 μm, with a large proportion less than 1 μm. The mean diameter of these oil droplets will be less than 1 μm, preferably from 0.2 to 0.5 μm. The emulsion product is then filled into borosilicate (Type I) glass vials which are stoppered, capped and terminally heat sterilized in a rotating steam autoclave at about 121° C.

[0323] These emulsions can withstand autoclaving as well as freezing at about 0° to −20° C. Such can be stored for a relatively long time with minimal physical and chemical breakdown, i.e. at least 12-18 months at 4°-8° C. The vehicle composition employed is chemically inert with respect to the incorporated pharmacologically active light activated drug.

[0324] The emulsions can exhibit very low toxicity following intravenous administration and exhibit no venous irritation and no pain on injection. The emulsions exhibit minimal physical and chemical changes (e.g., formation of non-emulsified surface oil) during controlled shake-testing on a horizontal platform. Moreover, the oil-in-water emulsions promote desirable pharmacokinetics and tissue distribution of the light activated drug in vivo.

[0325] As discussed above, the light activated drug can also be delivered to the body in a matrix which includes microbubbles. Suitable substrates for the microbubble include, but are not limited to, biocompatible polymers, albumins, lipids, sugars or other substances. U.S. Pat. Nos. 5,701,899 and 5,578,291 teach a method for synthesizing microbubbles with a sugar and protein substrate and is incorporated herein by reference. U.S. Pat. Nos. 5,665,383 and 5,665,382 teaches a method for synthesizing microbubbles with a polymeric substrate and is incorporated herein by reference. U.S. Pat. Nos. 5,626,833 and 5,798,091 teach methods for synthesizing microbubbles with a surfactant substrate and are incorporated herein by reference. A preferred microbubble has a lipid substrate. U.S. Pat. Nos. 5,772,929 teaches methods for synthesizing microbubbles with a lipid substrate. U.S. Pat. Nos. 5,776,429, 5,715,824 and 5,770,222 teach preferred methods for synthesizing microbubbles with a lipid substrate and a gas interior and are incorporated herein by reference.

[0326] Suitable microbubbles with a lipid substrate can be liposomes. The liposomes can be unilamellar vesicles having a single membrane bilayer or multilamellar vesicles having multiple membrane bilayers, each bilayer being separated from the next by an aqueous layer. A lipid bilayer is composed of two lipid monolayers having a hydrophobic “tail” region and a hydrophilic “head” region. The formula of the membrane bilayer is such that the hydrophobic (nonpolar) “tails” of the lipid monolayers orient themselves towards the center of the bilayer, while the hydrophilic “heads” orient themselves toward the aqueous phase. Either unilamellar or multilamellar or other types of liposomes may be used.

[0327] A hydrophilic light activated drug can be entrapped in the aqueous phase of the liposome before the drug is delivered into the patient. Alternatively, if the light activated drug is lipophilic, it may associate with the lipid bilayer. Liposomes may be used to help “target” the light activated drug to an active site or to solublize hydrophobic light activated drugs. Light activated drugs are typically hydrophobic and form stable drug-lipid complexes.

[0328] As discussed above, many light activated drugs have low solubility in water at physiological pH’s, but are also insoluble in (1) pharmaceutically acceptable aqueous-

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organic co-solvents, (2) aqueous polymeric solutions and (3) surfactant/micellar solutions. However, such light activated drugs can still be “solubilized” in a form suitable for delivery into a body by using a liposome composition. For example, one example of a light activated drug BPD-MA (See Formula A of FIG. 20) can be “solubilized” at a concentration of about 2.0 mg/ml in aqueous solution using an appropriate mixture of phospholipids to form encapsulating liposomes.

[0329] Although the light activated drug can be included in many different types of liposomes, the following description discloses particular liposome compositions and methods for making the liposomes which are known to be “fast breaking”. In fast breaking liposomes, the light activated drug-liposome combination is stable in vitro but, when administered in vivo, the light activated drug is rapidly released into the bloodstream where it can associate with serum lipoproteins. As a result, the localized delivery of liposomes combined with the fast breaking nature of the liposomes can result in localization of the light activated drug in the tissues near the catheter. Further, the fast breaking liposomes can prevent the liposomes from leaving the vicinity of the catheter intact and then concentrating in non-targeted tissues such as the liver. Delivery of ultrasound energy from the catheter can also serve to break apart the liposomes after they have been delivered from the catheter.

[0330] Liposomes are typically formed spontaneously by adding water to a dry lipid film. Liposomes which include light activated drugs can include a mixture of the commonly encountered lipids dimyristoyl phosphatidyl choline (“DMPC”) and egg phosphatidyl glycerol (“EPG”). The presence of DMPC is important because DMPC is the major component in the composition to form liposomes which can solubilize and encapsulate insoluble light activated drugs into a lipid bilayer. The presence of EPG is important because the negatively charged, polar head group of this lipid can prevent aggregation of the liposomes.

[0331] Other phospholipids, in addition to DMPC and EPG, may also be present. Examples of suitable additional phospholipids that may also be incorporated into the liposomes include phosphatidyl cholines (PCs), including mixtures of dipalmitoyl phosphatidyl choline (DPPC) and distearoyl phosphatidyl choline (DSPC). Examples of suitable phosphatidyl glycerols (PGs) include dimyristoyl phosphatidyl glycerol (DMPG), DLPG and the like.

[0332] Other types of suitable lipids that may be included are phosphatidyl ethanolamines (PEs), phosphatidic acids (PAs), phosphatidyl serines, and phosphatidyl inositol.

[0333] The molar ratio of the light activated drug to the DMPC/EPG phospholipid mixture can be as low as 1:7.0 or may contain a higher proportion of phospholipid, such as 1:7.5. Preferably, this molar ratio is 1:8 or more phospholipid, such as 1:10, 1:15, or 1:20. This molar ratio depends upon the exact light activated drug being used, but will assure the presence of a sufficient number of DMPC and EPG lipid molecules to form a stable complex with many light activated drugs. When the number of lipid molecules is not sufficient to form a stable complex, the lipophilic phase of the lipid bilayer becomes saturated with light activated drug molecules. Then, any slight change in the process conditions can force some of the previously encapsulated light activated drug to leak out of the vesicle, onto the surface of the lipid bilayer, or even out into the aqueous phase.

[0334] If the concentration of light activated drug is high enough, it can actually precipitate out from the aqueous layer and promote aggregation of the liposomes. The more unencapsulated light activated drug that is present, the higher the degree of aggregation. The more aggregation, the larger the mean particle size will be, and the more difficult aseptic or sterile filtration will be. As a result, small changes in the molar ratio can be important in achieving proper filterability of the liposome composition.

[0335] Accordingly, slight increases in the lipid content can increase significantly the filterability of the liposome composition by increasing the ability to form and maintain small particles. This is particularly advantageous when working with significant volumes of 500 ml, a liter, five liters, 40 liters, or more, as opposed to smaller batches of about 100-500 ml or less. This volume effect is thought to occur because larger homogenizing devices tend to provide less efficient agitation than can be accomplished easily on a small scale. For example, a large size Microfluidizer™ has a less efficient interaction chamber than that one of a smaller size.

[0336] A molar ratio of 1.05:3.5 BPD-MA:EPG:DMPC (i.e., slightly less phospholipid than 1:8.0 light activated drug-phospholipid) may provide marginally acceptable filterability in small batches of up to 500 ml. However, when larger volumes of the composition are being made, a higher molar ratio of phospholipid provides more assurance of reliable aseptic filterability. Moreover, the substantial potency losses that are common in scale-up batches, due at least in part to filterability problems, can thus be avoided.

[0337] Any cryoprotective agent known to be useful in the art of preparing freeze-dried formulations, such as di- or poly saccharides or other bulking agents such as lysine, may be used. Further, isotonic agents typically added to maintain isomolarity with body fluids may be used. In a preferred embodiment, a disaccharide or polysaccharide is used and functions both as a cryoprotective agent and as an isotonic agent.

[0338] In a particular embodiment, the particular combination of the phospholipids, DMPC and EPG, and a disaccharide or polysaccharide form a liposomal composition having liposomes of a particularly narrow particle size distribution. When the process of hydrating a lipid film is prolonged, larger liposomes tend to be formed, or the light activated drug can even begin to precipitate. The addition of a disaccharide or polysaccharide provides instantaneous hydration and the large surface area for depositing a thin film of the drug-phospholipid complex. This thin film provides for faster hydration so that, when the liposome is initially formed by adding the aqueous phase, the liposomes formed are of a smaller and more uniform particle size. This provides significant advantages in terms of manufacturing case.

[0339] However, it is also possible that, when a saccharide is present in the composition, it is added after dry lipid film formation, as a part of the aqueous solution used in hydration. In a particularly preferred embodiment, a saccharide is added to the dry lipid film during hydration.
Disaccharides or polysaccharides are preferred to monosaccharides for this purpose. To keep the osmotic pressure of the liposome composition similar to that of blood, no more than 4-5% monosaccharides could be added. In contrast, about 9-10% of a disaccharide can be used without generating an unacceptable osmotic pressure. The higher amount of disaccharide provides for a larger surface area, which results in smaller particle sizes being formed during hydration of the lipid film.

Accordingly, the preferred liposomal composition comprises a disaccharide or polysaccharide, in addition to the light activated drug and the mixture of DMPC and EPG phospholipids. When present, the disaccharide or polysaccharide is preferably chosen from among the group consisting of lactose, trehalose, maltose, maltotriose, palatinose, lactulose or sucrose, with lactose or trehalose being preferred. Even more preferably, the liposomes comprise lactose or trehalose.

Also, when present, the disaccharide or polysaccharide is formulated in a preferred ratio of about 10-20 saccharide to 0.5-6.0 DMPC/EPG phospholipid mixture, respectively, even more preferably at a ratio from about 10 to 1.5-4.0. In one embodiment, a preferred but not limiting formulation is lactose or trehalose and a mixture of DMPC and EPG in a concentration ratio of about 10 to 0.94-1.88 to about 0.65-1.30, respectively.

The presence of the disaccharide or polysaccharide in the composition not only tends to yield liposomes having extremely small and narrow particle size ranges, but also provides a liposome composition in which light activated drugs, in a particular, may be stably incorporated in an efficient manner, i.e., with an encapsulation efficiency approaching 80-100%. Moreover, liposomes made with a saccharide typically exhibit improved physical and chemical stability, such that they can retain an incorporated light activated drug without leakage upon prolonged storage, either as a reconstituted liposomal or a cryodesiccated powder.

Other optional ingredients include minor amounts of nontoxic, auxiliary substances in the liposomal composition, such as antioxidants, e.g., butylated hydroxytoluene, alphatocopherol and ascorbyl palmitate; pH buffering agents, e.g., phosphates, glycine, and the like.

Liposomes containing a light activated drug may be prepared by combining the light activated drug and the DMPC and EPG phospholipids (and any other optional phospholipids or excipients, such as antioxidants) in the presence of an organic solvent. Suitable organic solvents include any volatile organic solvent, such as diethyl ether, acetone, methylene chloride, chloroform, piperidine, piperidine-water mixtures, methanol, tert-butanol, dimethyl sulfoxide, N-methyl-2-pyrrolidone, and mixtures thereof. Preferably, the organic solvent is water-immiscible, such as methylene chloride, but water immiscibility is not required. In any event, the solvent chosen should not only be able to dissolve all of the components of the lipid film, but should also not react with, or otherwise deleteriously affect, these components to any significant degree.

The organic solvent is then removed from the resulting solution to form a dry lipid film by any known laboratory technique that is not deleterious to the dry lipid film and the light activated drug. Preferably, the solvent is removed by placing the solution under a vacuum until the organic solvent is evaporated. The solid residue is the dry lipid film. The thickness of the lipid film is not critical, but usually varies from about 30 to about 45 mg/cm² depending upon the amount of solid residual and the total area of the glass wall of the flask. Once formed, the film may be stored for an extended period of time, preferably not more than 4 to 21 days, prior to hydration. While the temperature during a lipid film storage period is also not an important factor, it is preferably below room temperature, most preferably in the range from about −20 to about 4°C.

The dry lipid film is then dispersed in an aqueous solution, preferably containing a disaccharide or polysaccharide, and homogenized to form the desired particle size. Examples of useful aqueous solutions used during the hydration step include sterile water, a calcium- and magnesium-free, phosphate-buffered (pH 7.2-7.4) sodium chloride solution; a 9.75% w/v lactose solution; a lactose-saline solution; 5% dextrose solution; or any other physiologically acceptable aqueous solution of one or more electrolytes. Preferably, however, the aqueous solution is sterile. The volume of aqueous solution used during hydration can vary greatly, but should not be so great as about 98% nor so small as about 30-40%. A typical range of useful volumes would be from about 75% to about 95%, preferably about 85% to about 90%.

Upon hydration, coarse liposomes are formed that incorporate a therapeutically effective amount of the light activated drugs-phospholipid complex. The “therapeutically effective amount” can vary widely, depending on the tissue to be treated and whether it is coupled to a target-specific ligand, such as an antibody or an immunologically active fragment. It should be noted that the various parameters used for selective photodynamic therapy are interrelated. Therefore, the therapeutically effective amount should also be adjusted with respect to other parameters, for example, fluence, irradiance, duration of the light used in photodynamic therapy, and the time interval between administration of the light activated drug and the therapeutic irradiation. Generally, all of these parameters are adjusted to produce significant damage to tissue deemed undesirable, such as neovascular or tumor tissue, without significant damage to the surrounding tissue, or to enable the observation of such undesirable tissue without significant damage to the surrounding tissue.

Typically, the therapeutically effective amount is such to produce a dose of light activated drug within a range of from about 0.1 to about 20 mg/kg, preferably from about 0.15-2.0 mg/kg and even more preferably, from about 0.25 to about 0.75 mg/kg. Preferably, the w/v concentration of the light activated drug in the composition ranges from about 0.1 to about 8.0-10.0 g/L. Most preferably, the concentration is about 2.0 to 2.5 g/L.

The hydration step should take place at a temperature that does not exceed about 30°C, preferably below the glass transition temperature of the light activated drug-phospholipid complex formed, even more preferably at room temperature or lower, e.g., 15°-20°C. The glass transition temperature of the light activated drug-lipid complex can be measured by using a differential scanning microcalorimeter.
In accordance with the usual expectation that the aqueous solubility of a substance should increase as higher temperatures are used, at a temperature around the transition temperature of the complex, the lipid membrane tends to undergo phase transition from a “solid” gel state to a pre-transition state and, finally, to a more “fluid” liquid crystal state. At these higher temperatures, however, not only does fluidity increase, but the degree of phase separation and the proportion of membrane defects also increases. This results in an increasing degree of leakage of the light activated drug from inside the membrane to the interface and even out into the aqueous phase. Once a significant amount of liposome leakage has occurred, even slight changes in the conditions such as a small drop in temperature, can shift the equilibrium away from aqueous “solubility” in favor of precipitation of the light activated drug. Moreover, once the typically water-insoluble light activated drug begins to precipitate, it is not possible to re-encapsulate it when the lipid bilayer. The precipitate is thought to contribute significantly to filterability problems.

In addition, the usual thickness of a lipid bilayer in the “solid” gel state (about 47 Å) decreases in the transition to the “liquid” crystalline state to about 37 Å, thus shrinking the entrapped volume available for the light activated drugs to occupy. The smaller “room” is not capable of containing as great a volume of light activated drug, which can then be squeezed out of the saturated lipid bilayer interstices. Any two or more liposomes exuding light activated drug may aggregate together, introducing further difficulties with respect to particle size reduction and ease of sterile filtration. Moreover, the use of higher hydration temperatures, such as, for example, about 35° to 45° C, can also result in losses of light activated drug potency as the light activated drug either precipitates or aggregates during aseptic filtration.

The particle sizes of the coarse liposomes first formed in hydration are then homogenized to a more uniform size, reduced to a smaller size range, or both, to about 150 to 300 nm, preferably also at a temperature that does not exceed about 30° C, preferably below the glass transition temperature of the light activated phospholipid complex formed in the hydration step, and even more preferably below room temperature of about 25° C. Various high-speed agitation devices may be used during the homogenization step, such as a Microfluidizer™ model 110F; a sonicator; a high-shear mixer; a homogenizer; or a standard laboratory shaker.

It has been found that the homogenization temperature should be at room temperature or lower, e.g., 15°-20° C. At higher homogenization temperatures, such as about 32°-42° C, the relative filterability of the liposome composition may improve initially due to increased fluidity as expected, but then, unexpectedly, tends to decrease with continuing agitation due to increasing particle size.

Preferably, a high pressure device such as a Microfluidizer™ is used for agitation. In microfluidization, a great amount of heat is generated during the short-period of time during which the fluid passes through a high pressure interaction chamber. In the interaction chamber, two streams of fluid at a high speed collide with each other at a 90° angle. As the microfluidization temperature increases, the fluidity of the membrane also increases, which initially makes particle size reduction easier, as expected. For example, filterability can increase by as much as four times with the initial few passes through a Microfluidizer™ device. The increase in the fluidity of the bilayer membrane promotes particle size reduction, which makes filtration of the final composition easier. In the initial several passes, this increased fluidity mechanism advantageously dominates the process.

However, as the number of passes and the temperature both increase, more of the hydrophobic light activated drug molecules are squeezed out of the liposomes, increasing the tendency of the liposomes to aggregate into larger particles. At the point at which the aggregation of vesicles begins to dominate the process, the sizes cannot be reduced any further. Surprisingly, particle sizes actually then tend to grow through aggregation.

For this reason, the homogenization temperature is cooled down to and maintained at a temperature no greater than room temperature after the composition passes through the zone of maximum agitation, e.g., the interaction chamber of a Microfluidizer™ device. An appropriate cooling system can easily be provided for any standard agitation device in which homogenization is to take place, e.g., a Microfluidizer™, such as by circulating cold water into an appropriate cooling jacket around the mixing chamber or other zone of maximum turbulence. While the pressure used in such high pressure devices is not critical, pressures from about 10,000 to about 16,000 psi are not uncommon.

As a last step, the compositions are preferably aseptically filtered through a filter having an extremely small pore size, i.e., 0.22 μm. Filter pressures used during sterile filtration can vary widely, depending on the volume of the composition, the density, the temperature, the type of filter, the filter pore size, and the particle size of the liposomes. However, as a guide, a typical set of filtration conditions would be as follows: filtration pressure of 15-25 psi; filtration load of 0.8 to 1.5 ml/cm²; and filtration temperature of about 25° C.

A typical general procedure is described below with additional exemplary detail:

(1) Sterile filtration of organic solvent through a hydrophobic, 0.22 μm filter.

(2) Addition of EPG, DMPC, light activated drug, and excipients to the filtered organic solvent, dissolving both the excipients and the light activated drug.

(3) Filtration of the resulting solution through a 0.22 μm hydrophobic filter.

(4) Transfer of the filtrate to a rotary evaporator apparatus, such as that commercially available under the name Rotovap™.

(5) Removal of the organic solvent to form a dry lipid film.

(6) Analysis of the lipid film to determine the level of organic solvent concentration.

(7) Preparation of a 10% lactose solution.

(8) Filtration of the lactose solution through a 0.22 μm hydrophilic filter.
(0368) Hydration of the lipid film with a 10% lactose solution to form coarse liposomes.

(0369) Reduction of the particle sizes of the coarse liposomes by passing them through a Microfluidizer\textsuperscript{TM} three times.

(0370) Determination of the reduced particle size distribution of liposomes.

(0371) Aseptic filtration of the liposome composition through a 0.22 μm hydrophilic filter. (Optionally, the solution may first be pre-filtered with a 5.0 μm prefiter.)

(0372) Analysis of light activated drug potency.

(0373) Filling of vials with the liposome composition.

(0374) Freeze-drying.

Once formulated, the liposome composition may be freeze-dried for long-term storage if desired. For example, BPD-MA, a preferred light activated drug, has maintained its potency in a cryodesiccated liposome composition for a period of at least nine months at room temperature, and a shelf life of at least two years has been projected. If the composition is freeze-dried, it may be packed in vials for subsequent reconstitution with a suitable aqueous solution, such as sterile water or sterile water containing a saccharide and/or other suitable excipients, prior to administration, for example, by injection.

Preferably, liposomes that are to be freeze-dried are formed upon the addition of an aqueous vehicle containing a disaccharide or polysaccharide during hydration. The composition is then collected, placed into vials, freeze-dried, and stored, ideally under refrigeration. The freeze-dried composition can then be reconstituted by simply adding water for injection just prior to administration.

The liposomal composition provides liposomes of a sufficiently small and narrow particle size that the aseptic filtration of the composition through a 0.22 μm hydrophilic filter can be accomplished efficiently and with large volumes of 500 ml to a liter or more without significant clogging of the filter. A particularly preferred particle size range is below about 300 nm, more preferably below from about 250 nm. Most preferably, the particle size is below about 220 nm.

Generally speaking, the concentration of the light activated drugs in the liposome depends upon the nature of the light activated drug used. When BPD-MA is used for example, the light activated drug is generally incorporated in the liposomes at a concentration of about 0.10% up to 0.5% w/v. If freeze-dried and reconstituted, this would typically yield a reconstituted solution of up to about 5.0 mg/ml light activated drug.

For diagnosis, the light activated drugs incorporated into liposomes may be used along with, or may be labeled with, a radioisotope or other detecting means. If this is the case, the detection means depends on the nature of the label. Scintigraphic labels such as technetium or indium can be detected using ex vivo scanners. Specific fluorescent labels can also be used but, like detection based on fluorescence of the light activated drugs themselves, these labels can require prior irradiation.

The methods of preparing various light activated drugs, light activated drug conjugates, emulsions and microbubbles are described in greater detail in the examples below. These examples are readily adapted to preparing analogous light activated drugs, light activated drug conjugates, emulsions and microbubbles by substitutions of appropriate light activated drugs, site directing molecule, phospholipids, and other analogous components. The following examples are being presented to describe the preferred components, embodiments, utilities and attributes of the media. For example, although BPD-MA is used as the light activated drugs in the microbubble (liposome) examples, the invention is not intended to be limited to this particular light activated drug.

Example 1 describes the synthesis of a preferred texaphyrin derivative. Examples 2-4 describe different light activated drugs conjugated with oligonucleotides as site directing molecules. Examples 5 and 6 describes a synthesis of an emulsion including a light activated drug. Example 7 describes preparation of microbubbles which include a light activated drug.

**EXAMPLE 1**

Synthesis of Texaphyrin T2BET Metal Complexes

The synthesis of texaphyrins is provided in U.S. Pat. Nos. 4,935,408, 5,162,509 and 5,252,720, all incorporated by reference herein. The present example provides the synthesis of a preferred texaphyrin, named T2BET, having substituents containing ethoxy groups.

Lutetium(III) acetate hydrate can be purchased from Strem Chemicals, Inc. (Newburyport, Mass.), gadolinium(III) acetate tetrahydrate can be purchased from Acros/Johnson Matthey (Wheel Hill, Mass.) and LZY-54 zeolite can be purchased from UOP (Des Plaines, Ill.). Acetone, glacial acetic acid, methanol, ethanol, isopropyl alcohol, and n-heptanes can be purchased from J.T. Baker (Phillipsburg, N.J.). Triethyamine and Amberlite 904 anion exchange resin can be purchased from Aldrich (Milwaukee, Wisc.). All chemicals should be ACS grade and used without further purification.

**FIGS. 30A-I** illustrate the synthesis of the gadolinium (III) complex of 4,5-diethyl-10,23-dimethyl-9,24-bis(3-hydroxypropyl)-16,17-bis [2-[2-(2-methoxyethoxy) ethoxy]ethoxy]-pentazapentacyclo [20.2.1.1\textsuperscript{3,6},1\textsuperscript{10,11},0\textsuperscript{14,19}] heptacosa-1,3,5,7,9,11(27),12,14,16,18,20,22(25),23-tridecaene which is illustrated as Formula I of FIG. 30. The critical intermediate 1,2-bis[2-[2-(2-methoxyethoxy)ethoxy]ethoxy]-4,5-dinitrobenzene (Formula E) can be prepared according to a three-step synthetic process outlined in FIGS. 30A-I. (Note: References to “Formula A,” “Formula B,” etc. relate to FIGS. 30A, 30B, etc.).

**SYNTHESIS OF TRISTYLYNE GLYCOL MONOMETHYL ETHER MONOTOSYLATED, FORMULA B:** In an oven dried 12 L three-necked round-bottom flask, equipped with a magnetic stir bar and a 1000 mL pressure-equalizing dropping funnel, a solution of NaOH (440.0 g, 11.0 mol) is added to 1800 mL water, and the mixture is cooled to approximately 0° C. A solution of tristylyne glycol monomethyl ether, Formula A, (656.8 g, 4.0 mol) in THF (1000 mL) is added. The clear solution is stirred vigorously at 0° C. for 15 min and a solution of tosyl chloride (915.12, 4.8 mol) in THF (2.0 L)
added dropwise over a 1 h period. The reaction mixture is stirred for an additional 1 h at 0°C, and 10% HCl (5.0 L) is added to quench the reaction (to pH 5-7). The two-phase mixture is transferred to a 4 L separatory funnel, the organic layer removed, and the aqueous layer extracted with t-butyldimethyl ether (3×250 mL). The combined organic extracts are washed with brine (2×300 mL), dried (MgSO₄), and evaporated under reduced pressure to afford Formula B, 1217.6 g (95%) as a light colored oil. This material is taken to the next step without further purification.

[0386] Synthesis of 1,2-bis[2-(2-methoxyethoxy)ethoxy]benzene, Formula D: In a dry 5 L round-bottom flask equipped with an overhead stirrer, reflux condenser, and a gas line, K₂CO₃ (439.47 g, 3.18 mol) and MeOH (1800 mL) are combined under an argon atmosphere.

To this well-stirred suspension, catechol, Formula C, (140.24 g, 1.21 mol) is added, and the mixture heated to reflux. Formula B (1012.68 g, 3.18 mol) is then added in one portion. The suspension is stirred at reflux for 24 h, cooled to room temperature, and filtered through Celite. The pad is rinsed with 500 mL of methanol and the combined filtrates are evaporated under reduced pressure. The resulting brown residue is taken up in 10% NaOH (800 mL), and methylene chloride (800 mL) added with stirring. The mixture is transferred to a 2 L separatory funnel, the organic layer removed and the aqueous layer extracted with methylene chloride (3×350 mL). The organic extracts are combined, washed with brine (350 mL), dried (MgSO₄), evaporated under reduced pressure, and the residue dried in vacuo for several hours to yield 485.6 (95%) of 1,2-bis[2-(2-methoxyethoxy)ethoxy]benzene (Formula D). For Formula D: bp 165°-220°C. (0.2-0.5 mm Hg); FAB MS, m/z 402; HRMS, M+: 402.2258 (calcd. for C₂₅H₂₃O₃S, 402.2253).

[0387] Synthesis of 1,2-bis[2-(2-methoxyethoxy)ethoxy]hexyl]-4,5-dinitrobenzene, Formula E: In an oven dried 500 mL round-bottom flask Formula D (104 g, 0.26 mol) and glacial acetic acid (120 mL) are combined and cooled to 5°C. To this well stirred solution, concentrated nitric acid (80 mL) is added dropwise over 15-20 min. The temperature of the mixture is held below 40°C by cooling and proper regulation of the rate of addition of the acid. After addition, the reaction is allowed to stir for an additional 10-15 min and is then cooled to 0°C. Fuming nitric acid (260 mL) is added dropwise over 30 min while the temperature of the solution is held below 30°C. After the addition is complete, the red colored solution is allowed to stir at room temperature until the reaction is complete (ca. 5 h, TLC: 95/5: CH₂Cl₂/MeOH) and then poured into well stirred ice water (1500 mL). Methylene chloride (400 mL) is added, the two-phase mixture transferred to a 2 L separatory funnel and the organic layer removed. The aqueous layer is extracted with CH₂Cl₂ (2×150 mL) and the combined organic extracts washed with 10% NaOH (2×250 mL) and brine (250 mL), dried (MgSO₄), and concentrated under reduced pressure. The resulting orange oil is dissolved in acetone (100 mL), and the solution layered with n-hexanes (500 mL), and stored in the freezer. The resulting precipitate is collected by filtration yielding 101.69 g (80%) of Formula E as a yellow solid. For Formula E: mp 43°-45°C; FAB MS, (M+H): m/z 493; HRMS, (M+H): 493; HRMS, (M+H): 493.2030 (calcd. for C₂₉H₂₃N₂O₂S, 493.2035).

[0388] Synthesis of 1,2-diamino-4,5-bis[2-(2-methoxyethoxy)ethoxy]hexyl], Formula F: In an oven dried 500 mL round bottom flask, equipped with a Claisen adapter, pressure equalizing dropping funnel, and reflux condenser, 1,2-bis[2-(2-methoxyethoxy)ethoxy]hexylj, 4,5-dinitrobenzene (Formula E) (20 g, 0.04 mol) is dissolved in absolute ethanol (200 mL). To this clear solution, 10% palladium on carbon (4 g) is added and the dark black suspension is heated to reflux under an argon atmosphere. Hydrazine hydrate (20 mL) in EtOH (20 mL) is added dropwise over 10 min to avoid bumping. The resulting brown suspension is heated at reflux for 1.5 h at which time the reaction mixture is colorless and TLC analysis (95/5; CH₂Cl₂/MeOH) displays a low Rf UV active spot corresponding to the diamine. Therefore, the mixture is hot filtered through Celite and the pad rinsed with absolute ethanol (50 mL). The solvent is removed under reduced pressure and the resulting light brown oil is dried in vacuo (in the dark) for 24 h to yield 15.55 g (89%) of 1,2-diamino-4,5-bis[2-(2-methoxyethoxy)ethoxy]hexylbenzene (Formula F). For Formula F: FAB MS, m/z 432; HRMS, M+: 432.2471 (calcd. for C₂₉H₂₃N₂O₂S, 432.2482). This material is taken to the next step without further purification.


[0390] [20.2.1.1], 18, 11.04,10,12, 14,16,18,20,22,24-undecane, Formula H. In an oven dried 5 L round-bottom flask, 2,5-bis[5-formyl-3-(3-hydroxypropyl)-4-methylpyrrol-2-yl]methyl]-3,4-dicyclopropyl (Formula G) (The synthesis of Formula G is provided in U.S. Pat. No. 5,252,720, incorporated by reference herein.) (30.94 g, 0.0644 mol) and 4,5-diamino-bis[2-(2-methoxyethoxy)ethoxy]hexylbenzene (Formula F) (28.79 g, 0.0644 mol) are combined in absolute methanol (600 mL) under an argon atmosphere. To this well stirred suspension, a mixture of concentrated hydrochloric acid (6.7 m) in absolute methanol 200 mL is added in one portion. The mixture is gradually heated to 50°C, at which time the reaction goes from a cloudy suspension of starting materials to a dark red homogeneous solution as the reaction proceeds. After 3 h the reaction is judged complete by TLC analysis and UV/visible spectroscopy (λmax 369 nm). The reaction mixture is cooled to room temperature, 60 g of activated carbon (DARCO™) is added, and the resulting suspension is stirred for 20 min. The dark suspension is filtered through Celite to remove the carbon, the solvent evaporated to dryness, and the crude Formula H dried in vacuo overnight. Formula H is recrystallized from isopropyl alcohol/n-heptane to afford 50 g (85%) of a scarlet red solid. 

For Formula H: 1H NMR (CD₃OD): δ 1.11 (t, 6H, CH₂CH₃), 1.76 (p, 4H, pyrr-CH₂CH₂CH₂OH), 2.36 (s, 6H, pyrr-CH₂), 2.46 (q, 4H, CH₂OCH₂), 2.64 (t, 4H, pyrr-CH₂CH₂CH₂OCH₂), 2.39 (s, 6H, CH₃CH₂OCH₂), 3.29 (t, 4H, pyrr-CH₂CH₂CH₂OCH₂), 3.34-3.38 (m, 20H, CH₂CH₂OCH₂CH₂OCH₂), 4.0 (s, 4H, pyr-CH₂), 4.22 (s, 4H, PhCH₂CH₂OH), 7.45 (s, 2H, PhH), 8.36 (s, 2H, H/C=NC=); UV/vis: [(MeOH)]λmax: 369; FAB MS, [M+H]: m/z 878.5; HRMS, [M+H]: 878.5272 (calcd. for C₄₉H₃₇N₂O₄S) 878.5279.

[0391] Synthesis of the gadolinium (III) complex of 4,5-diethyl-10,23-dimethyl-9,24-bis(3-hydroxypropyl)-16,17-bis[2-(2-methoxyethoxy)ethoxy]hexyl]-13,20,25,26,27-pentaazapenta-
cyclo [20.2.1.3.6.10.11.14.19]heptacosa-1,3,5,7,9,11(27),12,
14,16,18,20,22(25),23-tridecaene, Formula I. Formula I is
prepared according to the process outlined in FIG. 30. In a
dry 2 L three-necked round-bottom flask, Formula H (33.0
g, 0.036 mol) and gadolinium(II) acetate tetrahydrate (15.4
g, 0.038 mol) are combined in methanol (825 mL). To this
well-stirred solution, gadolinium(III) acetate tetrahydrate
(15.4 g, 0.038 mol) and triethylamine (50 mL) are added and
the reaction is heated to reflux. After 1.5 h, air is bubbled
(i.e., at reflux) for 4 h into the dark green reaction solution
with aid of a gas dispersion tube (flow rate=20 cm³/min). At
this point, the reaction mixture is carefully monitored by
UV/Visible spectroscopy (i.e., a spectrum is taken every
0.5-1 h, ~1 drop diluted in 4.5 mL MeOH). The reaction is
deemed complete by UV/Vis (in MeOH ratio: 342 nm/472
nm=0.22-0.24) after 4 h. The dark green reaction is cooled
to room temperature, filtered through Celite into a 2 L round
bottom flask, and the solvent removed under reduced pres-
sure. The dark green solid is suspended in acetone (1 L) and
the resulting slurry is stirred for 1 h at room temperature.
The suspension is filtered to remove the red/brown impuri-
ties (incomplete oxidation products), the solids rinsed with
acetone (300 mL), and air dried. The crude complex (35 g)
is dissolved in MeOH (750 mL), stirred vigorously for 15
min, filtered through Celite, and transferred to a 2 L Erlen-
meyer flask. An additional 300 mL of MeOH and 90 mL
water are added to the flask, along with acetic acid washed
LZY-54 zeolite (150 g). The suspension is agitated with an
overhead mechanical stirrer for approximately 3-4 h. The
zeolite extraction is deemed complete with the absence of
free Gd(III). To test for free gadolinium, the crude Formula
I is spotted heavily onto a reverse phase TLC plate (What-
man KC8F, 15x10 cm) and the chromatogram developed
using 10% acetic acid in methanol. The green complex
moved up the TLC plate close to the solvent front. Any free
gadolinium metal will remain at the origin under these	
conditions. After developing the chromatogram, the plate is
dried and the lower ³ of the plate stained with an Arsenazo
III solution in methanol (4 mg Arsenazo III in 10 mL
methanol). A very faint blue spot (indicative of free metal)
is observed at the origin against a pink background indicat-
ing very little free gadolinium metal. The zeolite is removed
through a Whatman #3 filter paper and the collected solids
rinsed with MeOH (200 mL). The dark green filtrate is
loaded onto a column of Amberlite IRA-904 anion exchange
resin (30 cm lengthx2.5 cm diameter) and eluted through the
resin (ca. 10 mL/min flow rate) into a 2 L round bottom flask
with 300 mL 1-butanol. The resin is rinsed with an addi-
tional 100 mL of MeOH and the combined eluent evaporated
to dryness under reduced pressure. The green shiny solid
Formula I is dried in vacuo for several hours at 40°C, to a
well stirred ethanolic solution (250 mL of Formula I at
55°-60°C, C, n-heptanes (ca. 600 mL) is added dropwise
(flow rate=4 mL/min) from a 1 L pressure-equalizing dropping
dummy. During the course of 1.5 h (300 mL addition) the
green Formula I began to crystallize out of the dark mixture.
After complete addition, the green suspension is cooled and
stirred for 1 h at room temperature. The suspension is
filtered, the solids rinsed with acetone (250 mL), and dried
in vacuo for 24 h to afford 26 g (63%), UV/Vis: [Me(OH)]
λmax nm: 316, 350, 415, 473, 739; FAB MS (M-OAc)−: m/e
1050; HRRMS, (M-20 Ac)−: m/e 1072.4036 (calcd. for
[Cs₂H₂GdN₁₂O₁₄]·0.5H₂O: C, 53.96; H, 6.36; N, 6.05, Gd, 13.59. Found: C, 53.73; H, 6.26; N, 5.82; Gd, 13.92.

[0392] Synthesis of the Lutetium(III) Complex of Formula H: The macroyclic ligand Formula H is oxidatively meta-
lated using lutetium(III) acetate hydrate (9.75 g, 0.0230 mol)
and triethylamine (22 mL) in air-saturated methanol (1500
mL) at reflux. After completion of the reaction (as judged
by the optical spectrum of the reaction mixture), the deep green
solution is cooled to room temperature, filtered through a
pad of celite, and the solvent removed under reduced pres-
sure. The dark green solid is suspended in acetone (600 mL,
stirred for 30 min at room temperature, and then filtered to
wash away the red/brown impurities (incomplete oxidation
products and excess triethylamine). The crude complex is
dissolved into MeOH (300 mL, stirred for ~30 min, and then
filtered through celite into a 1 L Erlenmeyer flask. An
additional 50 mL of MeOH and 50 mL of water are added to
the flask along with acetic acid washed LZY-54 zeolite
(40 g). The resulting mixture is agitated or shaken for 3 h,
then filtered to remove the zeolite. The zeolite cake is rinsed
with MeOH (100 mL) and the rinse solution added to the
filtrate. The filtrate is first concentrated to 150 mL and then
loaded onto a column (30 cm lengthx2.5 cm diameter) of
treated Amberlite IRA-904 anion exchange resin (resin in
the acetate form). The eluant containing the bis- acetate
lutetium(III) texaphyrin complex is collected, concentrated
to dryness under reduced pressure, and recrystallized from
anhydrous methanol/t-butylmethyl ether to afford 11.7 g
(63%) of a shiny green solid. For the complex: UV/vis: [
(MeOH)]λmax nm (log ε): 354, 414, 474 (5.10), 672, 732;
FAB MS, [M-OAc]−: m/e 1106.4; HRRMS, (M-OAc)−:
m/e 1106.4350 (calcd. for [Cs₂H₂GdN₁₂O₁₄]·3H₂O): 1106.4351.
Anal. calcd. for [Cs₂H₂GdN₁₂O₁₄]·3H₂O: C, 72.52; H, 6.30; N, 5.91. Found: C, 72.54; H, 6.18; N, 5.84.

EXAMPLE 2
Synthesis of a 12B1 TXP Metal
Complex-Oligonucleotide Conjugate

[0393] FIGS. 3IA-II illustrate the synthesis of a light
activated drug conjugate. The light activated drug is a
texaphyrin coupled with an oligonucleotide which is
complementary to a DNA site. As a result, the light activated
drug conjugate can bind the complementary DNA site and
will cleave the site upon activation by ultrasound. (Note:
References to “Formula A,”“Formula B,” etc. relate to
FIGS. 3IA, 3IB, etc.)

[0394] Synthesis of 4-Amino-l-[1(ethyloxy)acetyl-2-ox-
yl]-3-nitrobenzene (Formula B of FIG. 19), n=1. Potas-
sium carbonate (14.0 g, 101 mmol) and 4-amino-3-nitro-
phenol (Formula A) (10.0 g, 64.9 mmol) are suspended
in 150 mL dry acetonitrile. Ethyl-2-iodoacetate (10 mL, 84.5
mmol) (or ethyl iodobutyrate maybe used, in that case n=3)
is added via syringe, and the suspension is stirred at ambient
temperature for ca. 21 h. Chloroform (ca. 375 mL) is added
and is used to transfer the suspension to a separatory funnel,
whereupon it is washed with water (2xca. 100 mL). The
water washes are in turn washed with CHCl₃ (ca. 100 mL)
and the combined CHCl₃ extracts are washed with water (ca.
100 mL). Solvents are removed on a rotary evaporator, and
the residue is redissolved in CHCl₃ (ca. 500 mL) and
precipitated into hexanes (1.5 L). After standing two days,
the precipitate is filtered using a coarse fritted funnel and dried in vacuo to provide 14.67 g (Formula B), n=1 (94.1%).

**[0395]** Synthesis of 4-Amino-1-[1-(hydroxyacetyl-2-oxo)]-3-nitrobenzene (Formula C), n=1. 4-Amino-1-[1-(ethylhydroxyacetate-2-oxo)]-3-nitrobenzene (Formula B), n=1, (10.00 g, 37.3 mmol) is dissolved in tetrahydrofuran (100 mL), aqueous sodium hydroxide (1M solution, 50 mL) is added and the solution is stirred at ambient temperature for ca. 21 h. Tetrahydrofuran is removed on a rotary evaporator, and water (100 mL) is added. The solution is washed with 

CHCl₃ (ca. 200 mL), the neutralized by addition of hydrochloric acid (1M solution, 50 mL). The precipitate which formed is filtered after standing a few minutes, washed with water, and dried in vacuo to provide 8.913 g compound (Formula C), n=1 (99.5%). TLC: Rₜ=0.65, 10% methanol/CHCl₃.

**[0396]** Synthesis of 16-[1-Hydroxyacetyl-2-oxo]-9,24-bis(3-hydroxypropyl)-4,5-diethyl-10,23-dimethyl-13,20,25, 26,27-pentaazapentacyclo [20.2.1.1.0⁶,1.0⁸,11] octa-3,5,7,9,11(27),12,14(19),15,17, 20,22(25),23-tridecaene (Formula F), n=1: The metal complex of 16-[1-hydroxyacetyl-2-oxo]-9,24-bis(3-hydroxypropyl)-4,5-diethyl-10,23-dimethyl-13, 20,25,26,27-pentaazapentacyclo [20.2.1.1.0⁶,1.0⁸,11] octa-3,5,7,9,11(27),12,14(19),15,17,20,22(25),23-tridecaene (Formula F), n=1, (about 30 mol) and N-hydroxy succinimide (43 mmol) are dried together overnight in vacuo. The compounds are dissolved in dimethylformamide (anhydrous, 500 mL) and dichloroethylcarbodiimide (10 mg, 48 mmol) is added. The resulting solution is stirred under argon with protection from light for 8 h, whereupon a 110 mL aliquot is added to a solution of oligodeoxynucleotide (Formula G) (87 mmol) in a volume of 350 mL of 0.4M sodium bicarbonate buffer in a 1.6 mL Eppendorf tube. After vortexing briefly, the solution is allowed to stand for 23 h with light protection. The suspension is filtered through 0.45 mm nylon microfiltering tube, and the Eppendorf tube is washed with 250 mL sterile water. The combined filtrates are divided into two Eppendorf tubes, and glycogen (20 mg/mL, 2 μL) and sodium acetate (3M, pH 5.4) are added to each tube. After vortexing, ethanol (absolute, 1 mL) is added to each tube to precipitate the DNA. Ethanol is decanted following centrifugation, and the DNA is washed with an additional 1 mL aliquot of ethanol and allowed to air dry. The pellet is dissolved in 50% formamide gel loading buffer (20 mL), denatured at 90° C, for 1-2 days, and loaded on a 20% denaturing polyacrylamide gel. The band corresponding to conjugate (Formula H), n=1, is cut from the gel, crushed and soaked in 1xTBE buffer (ca. 7 mL) for 1-2 days. The suspension is filtered through nylon filters (0.45 μm) and desalted using a Sep-pak™ reverse phase cartridge. The conjugate is eluted from the cartridge using 40% acetonitrile, lyophilized overnight, and dissolved in 1 mM HEPES buffer, pH 7.0 (500 μL). The solution concentration is determined using UV/vis spectroscopy.

**EXAMPLE 3**

Synthesis of Texaphyrin Metal Complexes with Amine-, Thiol- or Hydroxy-Linked Oligonucleotides

**[0399]** Amides, ethers, and thioethers are representative of linkages which may be used for coupling site-directing molecules such as oligonucleotides to light activated drugs
such as texaphyrin metal complexes as illustrated in FIGS. 32A-F. (Note: References to “Formula A,” “Formula B,” etc. relate to FIGS. 32A, 32B, etc.). Oligonucleotides or other site-directing molecules functionalized with amines at the 5'-end, the 3'-end, or internally at sugar or base residues are modified post-synthetically with an activated carboxylic ester derivative of the texaphyrin complex. In the presence of a Lewis acid such as FeBr₃, a bromide derivatized texaphyrin (for example, Formula C of FIG. 32) will react with an hydroxyl group of an oligonucleotide to form an ether linkage between the texaphyrin linker and the oligonucleotide. Alternatively, oligonucleotide analogues containing one or more thiophosphate or thiol groups are selectively alkylated at the sulfur atom(s) with an alkyl halide derivative of the texaphyrin complex. Oligodeoxy-nucelotide-complex conjugates are designed so as to provide optimal catalytic interaction between the targeted DNA phosphodiester backbone and the texaphyrin.

[0400] Oligonucleotides are used to bind selectively with compounds which include the complementary nucleotide or oligo- or polynucleotides containing substantially complementary sequences. As used herein, a substantially complementary sequence is one in which the nucleotides generally base pair with the complementary nucleotide and in which there are very few base pair mismatches. The oligonucleotide may be large enough to bind probably at least 9 nucleotides of complementary nucleic acid.


[0402] In general, there are three commonly used solid phase-based approaches to the synthesis of oligonucleotides containing conventional 5'-3' linkages. These are the phosphoramidite method, the phosphonate method, and the triester method.

[0403] A brief description of a current method used commercially to synthesize oligomeric DNA is as follows: Oligomers up to ca. 100 residues in length are prepared on a commercial synthesizer, e.g., Applied Biosystems Inc. (ABI) model 392, that uses phosphoramidite chemistry. DNA is synthesized from the 3' to the 5' direction through the sequential addition of highly reactive phosphorus(V) reagents called phosphoramidites. The initial 3' residue is covalently attached to a controlled porosity silica solid support, which greatly facilitates manipulation of the polymer. After each residue is coupled to the growing polymer chain, the phosphorus(V) is oxidized to the more stable phosphorus(III) state by a short treatment with iodine solution. Unreacted residues are capped with acetic anhydride, the 5'-protective group is removed with weak acid, and the cycle may be repeated to add a further residue until the desired DNA polymer is synthesized. The full length polymer is released from the solid support, with concomitant removal of remaining protective groups, by exposure to base. A common protocol uses saturated ethanolic ammonia.

[0404] The phosphonate based synthesis is conducted by the reaction of a suitably protected nucleotide containing a phosphonate moiety at a position to be coupled with a solid phase-derivatized nucleotide chain having a free hydroxylphosphonate ester linkage, which is stable to acid. Thus, the oxidation to the phosphate or thiophosphate can be conducted at any point during synthesis of the oligonucleotide or after synthesis of the oligonucleotide is complete. The phosphonates can also be converted to phosphoramidite derivatives by reaction with a primary or secondary amine in the presence of carbon tetrachloride. [0405] In the triester synthesis, a protected phosphodiester nucleotide is condensed with the free hydroxyl of a growing nucleotide chain derivatized to a solid support in the presence of coupling agent. The reaction yields a protected phosphate linkage which may be treated with an oximate solution to form unprotected oligonucleotide.

[0406] To indicate the three approaches generically, the incoming nucleotide is regarded as having an “activated” phosphate/phosphate group. In addition to employing commonly used solid phase synthesis techniques, oligonucleotides may also be synthesized using solution phase methods such as diester synthesis. The methods are workable, but in general, less efficient for oligonucleotides of any substantial length.

[0407] Preferred oligonucleotides resistant to in vivo hydrolysis may contain a phosphorothioate substitution at each base (J. Org. Chem. 55:4693-4696, (1990) and Agrawal, (1990)). Oligodeoxyxynucleotides or their phosphorothioate analogues may be synthesized using an Applied Biosystem 380B DNA synthesizer (Applied Biosystems, Inc., Foster City, Calif.).

EXAMPLE 4

Synthesis of Diformyl Mononucleate Tripyrrane (FIG. 33. Formulas H) and Oligonucleotide Conjugate (FIG. 33. Formula J)

[0408] The present example provides for the synthesis of a light activated drug conjugate. The light activated drug conjugate includes a oligonucleotide acting as a site directing molecule coupled with the tripyrrane portion of a texaphyrin as illustrated in FIG. 33A-J. (Note: References to “Formula A,” “Formula B,” etc. relate to FIGS. 33A, 33B, etc.).

[0409] Synthesis of Dimethylster Dibenzyester Dipyrromethane (Formula B): A three-neck 1000 mL round-bottom flask set with a magnetic stirring bar, a thermometer, a heating mantle, and a reflux condenser attached to an argon line is charged with methylester acetoxyypprole (Formula A) (100.00 g; 267.8 mmol), 200 proof ethyl alcohol (580 mL), and deionized water (30 mL.) The reaction mixture is heated up and when the resulting solution begins to reflex, 12Naq. hydrochloric acid (22 mL.) is added all at once. The flask contents are stirred under reflux for two hours. The heating element is replaced by a 65°C bath and the resulting thick mixture is stirred for two hours prior to placing it in the freezer overnight.

[0410] The mixture is filtered over medium fritted glass funnel, pressed with a rubber dam, and washed with hexanes until the filtrate comes out colorless. The collected solids are set for overnight high vacuum drying at 30°C to afford slightly yellowish solids (65.85 g, 214.3 mmol, 80.0% yield.)
The solvent of the reaction suspension is removed under reduced pressure. The resulting solids are dried under high vacuum overnight.

The dry solids are suspended in a mixture of saturated aqueous sodium bicarbonate (1500 mL) and ethyl alcohol (200 mL), and stirred at its boiling point for five minutes. The hot suspension is filtered over celite. The filtrate is cooled down to room temperature and acidified to pH 6 with 12N aqueous hydrochloric acid. The resulting mixture is filtered over medium fritted glass. The collected solids are dried under high vacuum to constant weight (21.63 g, 49.78 mmol, 92.5% yield).

Synthesis of Dimethylester Dicarboxylic Acid Dipyrromethane, Formula C: All the glassware is oven dried. A three-neck 2000 mL round-bottom flask set with a magnetic stirring bar, a hydrogen line, and a vacuum line is charged with dimethylester dipyrromethane (Formula B) (33.07 g, 53.80 mmol), anhydrous tetrahydrofuran (1500 mL), and 10% palladium on charcoal (3.15 g). The flask is filled with dry hydrogen gas after each of several purges of the flask atmosphere prior to stirring the reaction suspension under a hydrogen atmosphere for 24 hours.

The solvent of the reaction suspension is removed under reduced pressure. The resulting solids are dried under high vacuum overnight.

The dry solids are suspended in a mixture of saturated aqueous sodium bicarbonate (1500 mL) and ethyl alcohol (200 mL), and stirred at its boiling point for five minutes. The hot suspension is filtered over celite. The filtrate is cooled down to room temperature and acidified to pH 6 with 12N aqueous hydrochloric acid. The resulting mixture is filtered over medium fritted glass. The collected solids are dried under high vacuum to constant weight (21.63 g, 49.78 mmol, 92.5% yield).

Synthesis of Methylester Dibenzyldiptyrranine, Formula E: A three-neck 2000 mL round-bottom flask set with a heating mantle, a magnetic stirring bar, a thermometer, and a reflux condenser attached to an argon line is charged with dimethylester dibenzylpyrramine (Formula C) (21.00 g, 48.33 mmol), ethyl acetate pyrrole (Formula D) (30.50 g), p-toluenesulfonic acid monohydrate (1.94 g), trifluoroacetic acid (39 mL), and methyl alcohol (1530 mL). The flask contents are heated and stirred under reflux for two hours. The heating element is replaced with a 0°C bath and the stirring is continued for half an hour prior to placing the resulting mixture in a freezer overnight.

The cold mixture is filtered over medium fritted glass. The collected solids are washed with hexanes and dried under high vacuum overnight (13.05 g, 19.25 mmol, 39.85% yield).

Synthesis of Methylester Dicarboxylic Acid Triptyrramine, Formula F: All the glassware is oven dried. A three-neck 500 mL round-bottom flask set with a magnetic stirring bar, a hydrogen line, and a vacuum line is charged with methyl ester dibenzyltripyrane (Formula E) (12.37 g, 19.13 mmol), anhydrous tetrahydrofuran (365 mL), and 10% palladium on charcoal (1.13 g). The flask is filled with dry hydrogen gas after each of several purges of the flask atmosphere prior to stirring the reaction suspension for 24 hours under a hydrogen atmosphere at room temperature.

The reaction suspension is filtered over celite. The solvent of the filtrate is removed under reduced pressure to obtain a foam which is dried under high vacuum overnight (10.49 g, 21.99 mmol, 87.0% pure).

Synthesis of Monoacid Triptyrramine, Formula H: All the glassware is oven dried. A three-neck 500 mL round-bottom flask set with a mechanical stirrer, a thermometer, a 0°C bath, and an additional funnel set with an argon line is charged with methyl ester dibenzyltripyrane (Formula F) (10.20 g, 17.83 mmol). Trifluoroacetic acid (32.5 mL) is dripped into the reaction flask from the addition funnel over a 45 minute period keeping the flask contents below 5°C. The resulting reaction solution is stirred at 0°C for 15 minutes, and then at 20°C for three hours. Triethylorthoformate (32.5 mL) is dripped into the flask from the addition funnel over a 20 minute period keeping the flask contents below 25°C by means of a dry ice/ethanol glycol bath. The reaction solution is stirred for one hour at ~25°C. Then a 0°C bath is set up. Deionized water (32.5 mL) is dripped into the reaction flask from the addition funnel keeping the flask contents below 10°C. The resulting two phase mixture is stirred at room temperature for 75 minutes and then added 1-butanol (200 mL). The solvents are removed under reduced pressure. The resulting dark oil is dried under high vacuum overnight to obtain black solids (11.64 g).

A three-neck 2000 mL round-bottom flask set with a thermometer, a heating mantle, a magnetic stirring bar, and a reflux condenser attached to an argon line, is charged with the crude methyl ester diformyl tripyrrane (Formula G) (11.64 g), methyl alcohol (900 mL), deionized water (60 mL), and lithium hydroxide monohydrate (4.7 g). The flask contents are heated, stirred under reflux for two hours, cooled down to room temperature, added deionized water (250 mL), acidified with 12N aqueous HCl to pH 5, and then stirred at 0°C for one hour. The resulting mixture is filtered over medium fritted glass funnel. The collected solids are dried under high vacuum to constant weight prior to their purification by column chromatography (silica gel, MeOH in CH₂Cl₂, 0-10%; 3.64 g, 8.06 mmol, 45.2% yield).

The monoacid tripyrrane (Formula H) is condensed with a derivatized ortho-phenylene diamine to form a non-oligomeric precursor which is then oxidized to an aromatic metal complex, for example, Formula I. An oligonucleotide amine may be reacted with the carboxylic acid derivatized tetrachloroformazine I to form the conjugate Formula J having the site-directing molecule on the T (trypyrane) portion of the molecule rather than the B (benzene) portion.

The following example describes the synthesis of an emulsion including tin ethyl etiopurpurin (SnEt₂) which is illustrated in FIG. 34.

Several emulsions are prepared as described above. In 5 mL glass tubes, medium chain length oil known as MCT oil (Miglyol 801, HVIs America, Piscatway, N.J.) is combined with 10 mg/gm SnEt₂ plus excipients as described above. Current emulsions also included additional excipients in the following concentrations: ethanol at mg/gm oil; egg phospholipids at 75 mg/gm oil; and sodium cholate at 10 mg/gm oil. After incubating for 30 minutes at 55°C, the tubes stand overnight at room temperature (19°-22°C). The tubes are centrifuged to remove bulk precipitates, and supernatants are filtered through 0.45 µm nylon membrane to remove any undissolved drug. Aliquots of filtrate are then diluted in chloroform/isopropyl alcohol (1:1) for spectrophotometric determination of drug concentration (absorbance at 662 nm). Reference standards are prepared with known concentrations of SnEt₂ in the same solvent.

The concentration of SnEt₂ in each of the emulsions is illustrated in Table 2. As illustrated, the concentration of SnEt₂ in the emulsion can be more than ten times the concentration in MCT oil alone.
TABLE 2

<table>
<thead>
<tr>
<th>Excipient Combination Added to MCT Oil</th>
<th>SnE12 Concentration mg/gm oil</th>
<th>SnE12 Concentration Normalized</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCT oil alone</td>
<td>0.38</td>
<td>1.00</td>
</tr>
<tr>
<td>+ ethanol</td>
<td>0.28</td>
<td>0.74</td>
</tr>
<tr>
<td>+ egg phospholipids (EYP)</td>
<td>0.89</td>
<td>2.34</td>
</tr>
<tr>
<td>+ Na cholate</td>
<td>1.17</td>
<td>3.08</td>
</tr>
<tr>
<td>+ ethanol + EYP</td>
<td>1.37</td>
<td>3.61</td>
</tr>
<tr>
<td>+ EYP + Na cholate</td>
<td>1.77</td>
<td>4.66</td>
</tr>
<tr>
<td>+ ethanol + Na cholate</td>
<td>2.20</td>
<td>5.79</td>
</tr>
<tr>
<td>+ ethanol + EYP + Na cholate</td>
<td>4.92</td>
<td>12.95</td>
</tr>
</tbody>
</table>

EXAMPLE 6

[0424] This example illustrates relative efficiencies of several bile salts. Mixtures of MCT oil, egg phospholipids, ethanol, and SnE12 are incubated with different bile salts, all at 4.6 nM, under the same conditions described above. As shown in Table 3, sodium cholate is the most efficient solubilizer. Cholic acid lacks solubilizing action in the oil.

[0428] The filterability of the composition in g/cm² is typically greater than about 10. Moreover, the yield is about 100% by HPLC analysis, with light activated drug potency typically being maintained even after sterile filtration. Average particle sizes vary from about 150 to about 300 nm (±50 nm).

TABLE 3

<table>
<thead>
<tr>
<th>Bile compound</th>
<th>SnE12 Concentration mg/gm oil</th>
<th>SnE12 Concentration Normalized</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>1.26</td>
<td>1.00</td>
</tr>
<tr>
<td>Na Taurocholate</td>
<td>1.13</td>
<td>0.90</td>
</tr>
<tr>
<td>Cholic acid</td>
<td>1.33</td>
<td>1.06</td>
</tr>
<tr>
<td>Na glycocholate</td>
<td>2.22</td>
<td>1.76</td>
</tr>
<tr>
<td>Na deoxycholate</td>
<td>2.31</td>
<td>1.83</td>
</tr>
<tr>
<td>Na cholate</td>
<td>3.70</td>
<td>2.94</td>
</tr>
</tbody>
</table>

EXAMPLE 7

[0425] The following Example describes the delivery of a light activated drug to a tumor. An emulsion is prepared having about 0.6 g SnE12/ml of emulsion and about 20 g of MCT oil based hydrophobic phase/ml of emulsion. The catheter illustrated in FIG. 7C is positioned in a vessel of the cardiovascular system using over the guidewire techniques. The catheter is positioned such that the media delivery port is adjacent to the aneroma using radiopaque markers on the catheter and the balloon is expanded into contact with the vessel wall. The emulsion is delivered via the third utility lumen 16B of the catheter 10. After the delivery of the emulsion, the ultrasound energy is delivered at about 0.3 W/cm² at a frequency of approximately 1.3 MHz for about ten minutes. After the delivery of ultrasound energy has concluded, the catheter is withdrawn from the vascularature of the tumor.

TABLE 4

<table>
<thead>
<tr>
<th>Light activated drug</th>
<th>0.21 g</th>
</tr>
</thead>
<tbody>
<tr>
<td>EPG</td>
<td>0.68 g</td>
</tr>
<tr>
<td>DMPC</td>
<td>1.38 g</td>
</tr>
<tr>
<td>BHT</td>
<td>0.002 g</td>
</tr>
<tr>
<td>Ascorbic acid 6-palmitate</td>
<td>0.002 g</td>
</tr>
<tr>
<td>Lactose NF crystalline injectable</td>
<td>10 g</td>
</tr>
<tr>
<td>Water for injection</td>
<td>100 ml</td>
</tr>
</tbody>
</table>

[0426] Using the above formulation, the total lipid concentration (% w/v) is about 2.06. The resulting solution is filtered through a 0.22 μm filter and then dried under vacuum using a rotary evaporator. Drying is continued until the amount of methylene chloride in the solid residue is no longer detectable by gas chromatography.

[0427] A 10% lactose/water-for-injection solution is then prepared and filtered through a 0.22 μm filter. Instead of being warmed to a temperature of about 35°C, the lactose/water solution is allowed to remain at room temperature (about 25°C) for addition to the flask containing the solid residue of the light activated drug/phospholipid. The solid residue is dispersed in the 10% lactose/water solution at room temperature, stirred for about one hour, and passed through a Microfluidizer™ homogenizer three to four times with the outlet temperature controlled to about 200°-250°C. The solution is then filtered through a 0.22 μm Durapore, hydrophilic filter.

EXAMPLE 8

[0429] The following Example describes the delivery of a light activated drug to a tumor. An emulsion is prepared having about 0.6 g SnE12/ml of emulsion and about 20 g of MCT oil based hydrophobic phase/ml of emulsion. The catheter illustrated in FIG. 7C is positioned in a vessel of the cardiovascular system using over the guidewire techniques. The catheter is positioned such that the media delivery port is adjacent to the aneroma using radiopaque markers on the catheter and the balloon is expanded into contact with the vessel wall. The emulsion is delivered via the third utility lumen 16B of the catheter 10. After the delivery of the emulsion, the ultrasound energy is delivered at about 0.3 W/cm² at a frequency of approximately 1.3 MHz for about ten minutes. After the delivery of ultrasound energy has concluded, the catheter is withdrawn from the vascularature of the tumor.

EXAMPLE 9

[0430] The following Example describes the delivery of a light activated drug to a tumor. An emulsion is prepared having approximately 0.8 g SnE12/ml of emulsion and approximately 30 g of MCT oil based hydrophobic phase/ml of emulsion. The catheter 10 illustrated in FIG. 3A is positioned in the vasculature of a tumor using over the guidewire techniques. The catheter is positioned such that the media delivery port is within the tumor using radiopaque markers included on the catheter. The prepared emulsion is delivered into the vasculature of the tumor via the utility lumen 16A. After the delivery of the emulsion, the ultrasound energy is delivered at about 0.3 W/cm² at a frequency of approximately 1.3 MHz for about fifteen minutes. After the delivery of ultrasound energy has concluded, the catheter is withdrawn from the vascular system of the patient.

EXAMPLE 10

[0431] The following Example describes the delivery of a light activated drug to a potential restenosis site. An emulsion is prepared having approximately 0.6 g SnE12/ml of
emulsion and approximately 30 g of MCT oil based hydrophobic phase/ml of emulsion. The catheter illustrated in FIG. 7C is positioned in the vasculature of a patient using over the guidewire techniques. The catheter is positioned such that the media delivery port is adjacent to a portion of the vessel wall which was previously treated with balloon angioplasty and the balloon is expanded into contact with the vessel wall. The prepared emulsion is delivered into the vasculature of the patient via the utility lumen 16B. Ultrasound energy is delivered from the ultrasound assembly to the potential restenosis site at about 0.3 W/cm² at a frequency of approximately 1.3 MHz for about ten minutes. After the delivery of ultrasound energy has concluded, the catheter is withdrawn from the vascular system of the patient.

EXAMPLE 11

[0432] The following Example describes the delivery of a light activated drug to an atheroma. Liposomes are prepared including BPD-MA (See FIG. 17) as the light activated drug and DMPC and EPG as the phospholipids. The molar ratio of BPD-MA:EPG:DMPC is about 1:3:7. The catheter illustrated in FIG. 7C is positioned in a vessel of the cardiovascular system using over the guidewire techniques. The catheter is positioned such that the media delivery port is adjacent to the atheroma using radiopaque markers included on the catheter and the balloon is expanded into contact with the vessel. Ultrasound energy is delivered at about 0.3 W/cm² at a frequency of approximately 1.3 MHz for about 20 minutes in order to rupture the liposomes and cause tissue death within the atheroma. After the delivery of ultrasound energy has concluded, the catheter is withdrawn from the vascular system of the patient.

EXAMPLE 12

[0433] The following Example describes the delivery of a light activated drug to a tumor. Liposomes are prepared including BPD-MA (See FIG. 17) as the light activated drug and DMPC and EPG as the phospholipids. The molar ratio of BPD-MA:EPG:DMPC is about 1:3:7. The catheter illustrated in FIG. 8 is positioned in the vasculature of a tumor using over the guidewire techniques. The catheter is positioned such that the media delivery port is within the tumor using radiopaque markers included on the catheter. Ultrasound energy is delivered at about 0.3 W/cm² at a frequency of approximately 1.3 MHz for about 20 minutes in order to rupture the liposomes and cause tissue death within the atheroma. After the delivery of ultrasound energy has concluded, the catheter is withdrawn from the vascular system of the tumor.

EXAMPLE 13

[0434] The following Example describes the delivery of a light activated drug to a potential restenosis site. Liposomes are prepared including BPD-MA (See FIG. 17) as the light activated drug and DMPC and EPG as the phospholipids. The molar ratio of BPD-MA:EPG:DMPC is approximately 1:3:7. The catheter illustrated in FIG. 7C is positioned in the vasculature of a patient using over the guidewire techniques. The catheter is positioned such that the media delivery port is adjacent to a portion of the vasculature which was previously treated with balloon angioplasty and the balloon is inflated into contact with the vessel wall. Ultrasound energy is delivered at about 0.3 W/cm² at a frequency of approximately 1.3 MHz for about 15 minutes in order to rupture the liposomes and cause tissue death within the atheroma. After the delivery of ultrasound energy is concluded, the catheter is withdrawn from the vascular system of the patient.

EXAMPLE 14

[0435] The following Example describes the delivery of a light activated drug to an atheroma. Liposomes are prepared including BPD-MA (See FIG. 17) as the light activated drug and DMPC and EPG as the phospholipids. The molar ratio of BPD-MA:EPG:DMPC is about 1:3:7. The phospholipids are systemically delivered. The catheter illustrated in FIG. 7C is positioned in the vasculature of a patient using over the guidewire techniques. The catheter is positioned such that the media delivery port is adjacent to the atheroma and the balloon is inflated into contact with the vessel wall. Ultrasound energy is delivered at about 0.3 W/cm² at a frequency of approximately 1.3 MHz for about 15 minutes. After the delivery of ultrasound energy is concluded, the catheter is withdrawn from the vascular system of the patient.

EXAMPLE 14

[0436] The following Example describes the delivery of a light activated drug to a tumor. Microbubbles are prepared including cisplatin and photofrin according to the methods disclosed in U.S. Pat. No. 5,770,222. The microbubbles are systemically administered. The catheter illustrated in FIG. 1A is positioned within the vasculature of a tumor. Ultrasound energy is delivered at about 0.3 W/cm² at a frequency of approximately 1.3 MHz for about 15 minutes. After the delivery of ultrasound energy is concluded, the catheter is withdrawn from the vascular system of the patient.

EXAMPLE 14

[0437] The following Example describes the delivery of a light activated drug to a tumor. Microbubbles are prepared including cisplatin and photofrin according to the methods disclosed in U.S. Pat. No. 5,770,222. The catheter illustrated in FIG. 3A is positioned within the vasculature of a tumor. The microbubbles are delivered to the tumor via the second utility lumen 16A of the catheter. Ultrasound energy is delivered at about 0.3 W/cm² at a frequency of approximately 1.3 MHz for about 15 minutes. After the delivery of ultrasound energy is concluded, the catheter is withdrawn from the vascular system of the patient.

EXAMPLE 16

[0438] The following Example describes the delivery of a light activated drug to a thrombosis. Microbubbles are prepared including heparin, photofrin and an albumin substrate. The microbubbles are systemically administered. The catheter illustrated in FIG. 1A is positioned adjacent to the thrombosis. Ultrasound energy is delivered at about 0.2 W/cm² at a frequency of approximately 1.3 MHz for about 20 minutes. After the delivery of ultrasound energy is concluded, the catheter is withdrawn from the vascular system of the patient.

EXAMPLE 17

[0439] Small hollow spheres covered with albumin were added to 5% human serum albumin in beakers in the ratio of
approximately 100,000,000 spheres per 1 ml, and these were divided into one group that was treated with the ultrasound-sensitive substance Rose Bengal and one group that was untreated. The beakers containing the respective small hollow spheres were irradiated with 1 MHz ultrasound at 0.5 W/cm² for 30 second and the number of small hollow spheres remaining after irradiation was counted. While nearly all of the small hollow spheres coated with Rose Bengal were ruptured, 70% of the untreated spheres kept their shape. In this manner, the presence of Rose Bengal was found to induce rupturing effects not obtained with the mechanical energy of ultrasound. Note that similar results were obtained using Eosin and other dyes instead of Rose Bengal.

What is claimed is:

1. A method of delivering a therapeutic composition to a target site comprising:
   delivering the therapeutic composition comprising genetic material through a catheter to the target site; and
   delivering ultrasound energy to the target site,
   wherein the catheter has an elongated catheter body with at least one axial lumen for delivery of genetic material therethrough, the catheter comprising at least one ultrasound transducer coupled to an energy source, wherein the at least one ultrasound transducer generates a sufficient level of ultrasound energy.
2. The method of claim 1, wherein the therapeutic composition further comprises a light activated compound.
3. The method of claim 1, wherein the genetic material is selected from the group consisting of DNA, RNA and analogs thereof.
4. The method of claim 1, wherein the genetic material is synthetic.
5. The method of claim 1, wherein the genetic material is recombinant.
6. The method of claim 1, wherein the therapeutic composition further comprises a microbubble.
7. The method of claim 2, wherein the therapeutic composition further comprises a microbubble.
8. The method of claim 1, wherein the genetic material comprises an oligonucleotide.
9. The method of claim 8, wherein the oligonucleotide has an affinity for a DNA in the target site.
10. The method of claim 9, wherein the DNA is a viral DNA.
11. The method of claim 9, wherein the DNA is an oncogene DNA.
12. The method of claim 9, wherein the oligonucleotide is an antisense oligonucleotide.
13. The method of claim 2, wherein the light activated drug is covalently bound to the genetic material.
15. The method of claim 1, wherein the at least one ultrasound assembly is positioned about a circumference of the elongated catheter body, the at least one support member supporting the at least one ultrasound transducer so as to define a chamber between the at least one transducer and the outer circumference of the elongated catheter body.
16. The method of claim 15, wherein the chamber is filled with a media that absorbs ultrasound energy such that a transmission of ultrasound energy from the ultrasound transducer to the elongated catheter body is reduced.
17. The method of claim 16, wherein the media is a gas selected from the group consisting of helium, argon, air and nitrogen.
18. The method of claim 16, wherein the media is a solid medium selected from the group consisting of silicon and rubber.
19. The method of claim 15, wherein the chamber is evacuated using a negative pressure.
20. The method of claim 1, wherein the catheter further comprises:
   a balloon positioned about the circumference of the elongated catheter body;
   at least one media delivery port in fluid communication with the at least one axial lumen for delivery of an expansion media to expand the balloon; and
   at least one media delivery port in fluid communication with the at least one axial lumen for delivery of a medicament.
21. The method of claim 20, wherein the balloon is positioned about the ultrasound assembly.
22. The method of claim 20, wherein the balloon is positioned about the circumference of the elongated catheter body adjacent to the ultrasound assembly.
23. The method of claim 1, wherein the ultrasound transducer is configured to deliver ultrasound energy of approximately 0.3 W/cm² at a frequency of approximately 1.3 MHz.
24. The method of claim 20, wherein pressure is used to drive the media across the balloon.
25. The method of claim 6, wherein the microbubble comprises a lipid substrate.
26. The method of claim 25, wherein the lipid substrate comprises a liposome.
27. The method of claim 6, wherein the interior of the microbubble includes a gas.

* * * * *