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CA 2552735 C 2012/09/11

(11)(21) **2 552 735**

(12) **BREVET CANADIEN
CANADIAN PATENT**

(13) **C**

(86) Date de dépôt PCT/PCT Filing Date: 2005/01/06

(87) Date publication PCT/PCT Publication Date: 2005/07/28

(45) Date de délivrance/Issue Date: 2012/09/11

(85) Entrée phase nationale/National Entry: 2006/07/06

(86) N° demande PCT/PCT Application No.: US 2005/000173

(87) N° publication PCT/PCT Publication No.: 2005/067986

(30) Priorités/Priorities: 2004/01/07 (US60/534,395);
2004/06/01 (US60/575,421); 2004/07/23 (US60/564,589)

(51) Cl.Int./Int.Cl. *A61L 9/00* (2006.01)

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(54) Titre : SYSTEME ET DISPOSITIF DE STERILISATION

(54) Title: STERILIZATION SYSTEM AND DEVICE

(57) Abrégé/Abstract:

A system, device and method for sterilizing or decontaminating an object that includes a sealable sterilizing chamber (12) and a sterilant gas-generating composition (24) that preferably generates NO or a mixture of NO and NO₂. The preferred sterilant gas-generating composition (24) includes a carbon-based diazeniumdiolate compound and a powdered acid.

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

(19) World Intellectual Property
Organization
International Bureau



(43) International Publication Date
28 July 2005 (28.07.2005)

PCT

(10) International Publication Number
WO 2005/067986 A1

- (51) International Patent Classification⁷: **A61L 9/00**
- (74) Agents: **GOTTS, Lawrence, J.** et al.; Shaw Pittman LLP, 1650 Tysons Boulevard, McLean, VA 22102-4859 (US).
- (21) International Application Number: PCT/US2005/000173
- (81) Designated States (*unless otherwise indicated, for every kind of national protection available*): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.
- (22) International Filing Date: 6 January 2005 (06.01.2005)
- (25) Filing Language: English
- (26) Publication Language: English
- (30) Priority Data:
60/534,395 7 January 2004 (07.01.2004) US
60/575,421 1 June 2004 (01.06.2004) US
60/564,589 23 July 2004 (23.07.2004) US
- (84) Designated States (*unless otherwise indicated, for every kind of regional protection available*): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).
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- Published:**
— with international search report
- For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.*

(54) Title: STERILIZATION SYSTEM AND DEVICE

(57) Abstract: A system, device and method for sterilizing or decontaminating an object that includes a sealable sterilizing chamber (12) and a sterilant gas-generating composition (24) that preferably generates NO or a mixture of NO and NO₂. The preferred sterilant gas-generating composition (24) includes a carbon-based diazeniumdiolate compound and a powdered acid.

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STERILIZATION SYSTEM AND DEVICE**[0001]****FIELD OF THE INVENTION**

[0002] This invention relates to sterilization devices and methodologies using gas as the sterilant. Specifically, the invention relates to a device that uses a liquid to activate a composition to rapidly generate nitric oxide, as well as additional oxides of nitrogen, as the sterilant gases.

BACKGROUND OF THE INVENTION

[0003] Steam autoclaving is the hospital standard for sterilizing most medical instruments. This method exposes materials to steam at 121°C at 15-20 lbs of pressure for 15-30 minutes. Killing is mediated by heat denaturation of proteins, DNA, and subsequent interruption of metabolic functions. The method requires cumbersome equipment, a power supply and plumbing, although benchtop models have fillable water tables. Aside from these logistical problems, autoclaving is not suitable for many plastics and other heat labile materials.

[0004] Sterilant gases can kill or control the growth of microbial contaminations. Some of these sterilant gases include chlorine dioxide, sulfur dioxide, hydrogen peroxide, nitric oxide, nitrogen dioxide, carbon dioxide, hydrogen sulfide, ozone and ethylene oxide. One problem with many of the sterilant gases is that they are explosive in high concentrations (e.g. ethylene oxide, hydrogen peroxide, chlorine dioxide, ozone). Thus, storing, containing and using these gases in high concentrations represent a hazard to the user. For safety reasons, this limits the usable concentration of gas and creates an additional disadvantage. That is because the concentration of the sterilant gas must be decreased due to safety concerns, the exposure time must be increased to achieve effective sterilization. In general, therefore, the more hazardous the chemical in

terms of its potential as a chemical hazard, the longer the sterilization cycle becomes.

[0005] Certain sterilant gases, such as chlorine dioxide, ozone and hydrogen peroxide are difficult and expensive to transport. Many of these sterilant gases are powerful oxidizers. Oxidizing gases are expensive and paperwork intensive to ship in bulk tanks, further complicating their use. Gases, such as ozone and chlorine dioxide, must be generated at or near the point of use. On-site plants for generating one such sterilant gas, chlorine dioxide, are costly and require significant space to implement.

[0006] Hamilton U. S. Patent No. 6,607,696 describes device for delivering chlorine dioxide to disinfect or sterilize a liquid or an item contained in the liquid. The device uses a permeable sachet containing gas generating reactants, such as sodium chlorite and citric acid, where the sachet is a receptacle permeable to liquid and gas. Liquid can diffuse into the receptacle to reach the gas generating reactants that then generate a gas, such as chlorine dioxide. The gas that diffuses out of the permeable sachet is not sealed from the environment / atmosphere. . Multi-compartmental devices that employ gas-generating ingredients contained in closed compartments that are permeable and permit the diffusion of liquids and gases through the compartments to produce chlorine dioxide, such as the sachet and envelope compartments used in U.S. Patents 6,602,466 and 6,607,696. Not only are these systems expensive and difficult to manufacture, but they do not contain the generated gases in a manner that prevents their unintended escape to the environment/atmosphere nor do they allow the user to predictably and controllably release the gas into a sealable container that is sealed when the contents are sterilized.

[0007] Thus, there is a need for methods and devices that generate sterilant gases at the point of use in a safe, and efficient manner. There is a further need for processes capable of producing significant concentrations of sterilant gas without the danger of explosion or oxidative fire. There is a need to produce greater concentrations of NO in a short time period to allow a shorter exposure and make the sterilization process more efficient. There is also a need for a system and method to generate small amounts of sterilant gas in an economical manner. The ability to economically generate small amounts of sterilant gases allows for easy

transportation of the sterilizing system, imparting a portability to the system not commonly found with traditional sterilization devices and methods.

[0008] Thus, given the problems with traditional gaseous sterilants and disinfectants, there is a need for a sterilant gas generating system and method where the risk of explosion and oxidative fire is minimized, that produces the sterilant gas rapidly, safely, economically, and in a scaleable manner. In addition, there is a need for a gas that can be safely used at high enough concentrations to act rapidly to minimize the time required for sterilization or disinfecting.

SUMMARY OF THE INVENTION

[0009] The present invention provides a method to generate nitric oxide, nitrogen dioxide, and additional oxides of nitrogen for the purpose of sterilization and disinfecting. By using compounds that generate nitric oxide on acidification, and combining the nitric oxide with ambient air within an exemplary device, the method generates both water soluble and lipid soluble oxides of nitrogen each of which possess anti-microbial properties on their own. In addition, the mixture of gases generated in the present invention has lower oxidation potential than other sterilant gases, making them safer to handle. Furthermore, the mixture of gases lacks the potential for explosive hazard possessed by many currently favored sterilant gases.

BRIEF DESCRIPTION OF THE DRAWINGS

[0010] **FIG. 1** shows a sterilization device having a sterilization chamber (SC) 12, a gas generation chamber (GGC) 14, and a connecting tube 16 having a safety valve 18. The SC 12 has a closure 20, a connecting port 15, and an exhaust port 22 that attaches to exhaust tubing 29. An exhaust valve 23 is attached to the exhaust tubing 29. The GGC 14 contains the composition capable of generating a sterilant gas (sterilant gas-generating composition) 24. The GGC 14 has a fitting 17 to which connecting tubing 16 is attached, and a filling port 21 for adding liquids.

[0011] **FIG. 2** is a drawing of another embodiment of the Sterilization Chamber 12 that has a flap closure 30 for opening or sealing the SC 12.

[0012] **FIG. 3** is a schematic drawing of a sterilization device 100 of the present invention that is comprised of a hard casing with internal gas pumping and scrubbing functions.

[0013] The foregoing features of the invention will be more readily understood by reference to the following detailed description.

DETAILED DESCRIPTION OF THE INVENTION

[0014] The present invention provides systems and devices that generate nitric oxide, along with additional oxides or nitrogen, to sterilize or disinfect instruments, devices, tools and equipment that must be sterile, typically for medical applications. The use of nitric oxide alone, or in combination with oxides of nitric oxide that form in combination with air, as a disinfectant and sterilant gas mixture has several advantages over other gases. Neither nitric oxide or its oxides are explosive at high concentrations. In addition, because nitric oxide and its oxides have a weaker oxidizing potential than peroxides and ozone, they allow for a broader list of materials that can be sterilized. Another advantage to using nitric oxide and or its oxides is that their density is closer to that of air, and thus do not settle to the bottom of a closed compartment when mixed with air, as would chlorine dioxide, which is greater than twice the density of air.

[0015] Generating a mixture of the oxides of nitrogen can have additional advantages over pure nitric oxide and other single entity sterilization gases. Nitric oxide is very lipid soluble and has the ability to disrupt the lipid membranes of microorganisms. Furthermore nitric oxide may inactivate thioproteins thereby disrupting the functional proteins of microbes. Finally, nitric oxide and nitrogen dioxide are extremely effective disruptors of DNA, causing strand breaks and other damage leading to an inability for the cell to function.

Definitions

[0016] As used herein, the term “nitric oxide” or “NO” means the NO free radical or NO•. As used herein, the term “NO-generating” compound or composition means a compound or composition capable of producing or releasing NO, NO₂, and NO_x. As used herein, the term “sterilant gas-generating” compound or composition means a compound or composition capable of producing or releasing a sterilant gas. Examples of preferred sterilant gases include, but are not limited to NO, NO₂, NO_x, and mixtures thereof. An NO-generating compound is one type of sterilant gas-generating compound. The preferred sterilant gas-generating compounds used in the systems, devices and methods of the present

invention are carbon-based diazeniumdiolate compounds that generates at least 1 mole of NO per mole of compound.

[0017] As used herein, the term “sterilization chamber” means any gas tight chamber of any size, whether comprised of hard or soft materials, where items to be sterilized or decontaminated can be contained. One skilled in the art will know that the addition of NO to air will result in the formation of NO₂ because NO reacts with the oxygen in air.

[0018] As used herein, the term “gas generation chamber” means any container, of any size or composition, which may be used to contain a gas. As used herein, the term “microbe” means any bacteria, virus, fungi, parasite, mycobacteria or the like. As used herein, the term “scrubbing” means the removal or conversion of toxic oxides of nitrogen from the exhaust stream of the sterilization device.

[0019] As used herein, the term “impermeable” means a substance, material or object that prohibits over 95% of any liquid or gas from passing or diffusing through it, for at least one hour. As used herein, the term “permeable” means a substance, material or object that allows the passage of gases and/or liquid through it.

[0020] The sterilization system and method of the present invention employs compounds that release a sterilant gas, preferably nitric oxide, upon acidification. The system and method of the present invention generates nitric oxide that is used, typically as a mixture of water soluble and lipid soluble nitrogen oxide gases, to sterilize a wide variety of devices, instruments, materials, human and animal tissues, drugs, biologicals, and a variety of medically relevant materials.

[0021] A preferred embodiment of the system and method of the present invention generates the gases can be generated at the point-of-use. Such point-of-use methods, systems and devices eliminate the need for heavy tanks of potentially hazardous gases or expensive on-site gas generation plants. The point-of-use gas generation employed in the system and method of the present invention can be functional without requiring electricity, which allows the method to be adapted for portable embodiments for sterilization, disinfecting, and decontamination in austere environments such as combat areas, refugee camps, etc. In one aspect, the present invention describes a method to generate a mixture of nitrogen oxides for

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sterilization and disinfecting purposes. The method requires an apparatus that integrates the gas generation and delivery method. The apparatus used in the process may have many potential embodiments.

[0022] One currently preferred embodiment includes a gas generation chamber in fluid connectivity with a sterilization chamber. Another preferred embodiment has the gas generation chamber contained within the sterilization chamber.

[0023] Also preferred, are embodiments of the system and method of the present invention that produces a mixture of nitrogen oxides having less oxidative potential than commonly used sterilant gases, including ozone and hydrogen peroxide. An additional advantage is that the mixture of nitrogen oxides produced has much less explosive potential than the commonly used sterilant gases, including ethylene oxide, hydrogen peroxide, and chlorine dioxide. This allows the use of greater concentrations of the gaseous mixture the system and method of the present invention thereby allowing less exposure time in the sterilization cycle as known to those skilled in the art.

[0024] Yet another advantage is that the method of the present invention is the generation of multiple chemical species with different chemical properties for the purpose of sterilization and disinfecting. Those skilled in the art understand that multiple mechanisms of cell killing or deactivation are often preferred over single mechanisms of action. Antimicrobial agents with different mechanisms of action are often synergistic, producing a greater effect than would be expected by simply adding the effects from each agent together. The same principle is applied to microbial resistance, where multiple, distinctly acting agents are used for treatment.

[0025] A most preferred embodiment of the system and method of the present invention, the NO-releasing compound employed is a carbon-based diazeniumdiolate compound. Carbon-based diazeniumdiolate molecules release a greater amount of nitric oxide and do not form nitrosamines. Preferably, the carbon-based diazeniumdiolate compound produces greater quantities of NO per mole. Preferably, a C-based diazeniumdiolate compound that is capable of producing at least one mole of NO per mole of diazeniumdiolate is used as the sterilant gas generating compound is used in the system and method of the present invention. Such a carbon-based diazeniumdiolate is described in U.S. 2005/0203069.

- [0026] The system and method of the present invention preferably uses a C-based diazeniumdiolate compound that does not result in the formation of carcinogenic nitrosamines when acidified. Another advantage of using a C-based diazeniumdiolate compound as the preferred NO-releasing compound is that it releases a greater quantity of NO per mole of NO-releasing compound. For example, nitrogen-based diazeniumdiolates and nitrosothiols produce lower quantities of NO per mole compound when compared to the carbon-based diazeniumdiolate compounds. Also, the use of a C-based diazeniumdiolate compound as the preferred NO-releasing compound allows the use of an acid to release NO rather than the copper solution required for release of NO from nitrosothiols. Yet another advantage of the method and system of the present invention is that it has a reduced environmental impact as compared to a method requiring a solution containing copper ions.
- [0027] The nitric oxide generating compounds utilized in the system and method of the present invention provide several advantageous elements to the present invention. One advantage is that nitric oxide has a high degree of lipid solubility, making it toxic to almost all microbes, which have lipid membranes (the exception is non-enveloped viruses).
- [0028] Nitrogen dioxide, and other oxides of nitrogen such as dinitrogen tetroxide, are more water soluble than nitric oxide. These, and especially nitrogen dioxide, are highly damaging to DNA, resulting in nitrosation and deamination of DNA bases and single and double strand breaks. Damage to DNA is a powerful killing mechanism. Combined, the mixture of gases in the present invention provide a multi-pronged attack of microbes through a variety of possible mechanisms of action. The antimicrobial benefits of a method that uses multiple mechanisms of action was discussed above.
- [0029] Yet another advantage to the system and method of the present invention is that it can permit the formation of small amounts of nitrous acid in the water that attaches to solids in humid environments, which can enhance the antimicrobial properties of the present invention.

[0030] Methods to generate sterilant gases. A preferred embodiment of the method and system of the present invention generates NO using the class of nitric oxide donors known as diazeniumdiolates. These compounds spontaneously release NO in solution, with rates that are proportional to the acidity of the solution. Highly acidic conditions can be used to generate NO in the method of the present invention, generate NO gas rapidly (complete theoretic release of NO in < 30 sec). A preferred embodiment of the method and system of the present invention uses carbon-based diazeniumdiolates rather than nitrogen-based compounds, due to the ability of the latter to form the highly carcinogenic nitrosamine species. See Parzuchowski et al., J Am Chem. Soc 124: 12182-91 (2002). Also preferred are carbon-based diazeniumdiolates that release large amounts of NO such as but not limited to those described in U.S. 2005/0203069, which produce greater amounts of nitric oxide per mole of compound, as compared to the compounds disclosed in U.S. Pat. No. 6,232,336.

[0031] One NO-generating compound that may be used in the method and system of the present invention, though with caution, is sodium nitroprusside because of its concurrent formation of cyanide in the gas generation chamber. The formation of cyanide represents a human health hazard and creates a disposal safety issue for the gas generation chamber. Nitrosothiols may also be used to generate NO in the current invention, however nitrosothiols have the tendency to reform after they have released NO, thus creating a chemical sink for NO and making the release of NO unpredictable.

[0032] Another embodiment of the system and method of the present invention uses a gas generating chamber that is a pressurized or non-pressurized cylinder containing NO may be used. Though this embodiment sacrifices portability, it is useful in large scale decontaminations, such as military or other very large equipment. The NO may be stored at high concentrations within the cylinder. Although this embodiment is less desirable due to the hazard and added costs and paperwork involved with shipping of concentrated pressurized gases. A more preferred method would be to dilute the NO concentration within the cylinder to a desired concentration by mixing with nitrogen or other inert gas including, but not

limited to argon, helium, and neon. The gas or gas mixture can be delivered to the sterilization chamber through a metered regulator in fluid connectivity with the sterilization chamber, or other gas delivery method known to one skilled in the art. Another embodiment includes computer or microprocessor means to control the delivery of sterilant gas from the gas cylinder.

[0033] In embodiments of the present invention where the NO-releasing entity is activated by acid, any acid can be used to generate NO. In one embodiment of the present invention the NO donors are activated by the addition of an aqueous acid as described in Example 1. Due to the inconvenience of handling and transporting aqueous acids, powdered acids that are activated by water are preferred. While any powdered acid would be acceptable, powdered acids with low pKa are preferred because the preferred method is to rapidly generate the NO, and low pKa acids are more effective as shown in Example 14. These low pKa acids include but are not limited to oxalic and maleic acids. Generally, up to ten-fold molar excess of powdered acid is used, however lower molar ratios may also be acceptable.

[0034] A preferred embodiment of the system and method of the present invention includes a gas generation chamber containing both a carbon-based diazeniumdiolate and a powdered acid, whereby the gas generation chamber includes a rapidly sealing opening that allows the addition of a liquid, preferably water, and is in fluid connectivity with the sterilization chamber so that gas generated upon activation of the carbon-based diazeniumdiolate is transported into the sterilization chamber. Additional connections and/or ports may be included for such purposes as to apply a vacuum, if necessary, to release NO gas from the chamber. Preferably, the NO gas is released into a reusable NOx scrubbing system).

[0035] A desiccant may be included in the gas generation chamber to reduce moisture during manufacture, shipping, and storage of the gas generation chamber. Examples of desiccants may include but not be limited to molecular sieves, silica gels, and other methods known to one skilled in the art. Care should be taken that the amount of desiccant will not prevent the generation of NO on the addition of water.

[0036] One skilled in the art can apply The Ideal Gas Law, the moles of NO released from the various NO-releasing compounds, the molecular weight of the compound in question and derive the weight of the compound necessary in the gas generation chamber to achieve a desired percent of NO added to any specified volume that comprises the sterilization chamber. For example, 1.956 grams of an NO-releasing compound that generates 2 moles of NO per mole of compound having a molecular weight of 163 gms/mole is used to produce 0.0225 moles of NO and provide a concentration of 50% NO in a one liter volume. This allows the user to control the amount of NO added for various sterilization applications. For example, medical practitioners may desire a more rapid sterilization cycle, requiring higher concentrations of added NO. Those users who are more concerned with portability may be less sensitive to speed and cost of the process. Longer sterilization cycles may require less of the NO-releasing compound, i.e. less NO added. Thus, the device 100 and process offer the flexibility to provide potential end users with options regarding cost, speed, portability, and other utilization parameters.

[0037] In one embodiment of the present invention, a lightweight, portable device employing chemically generated NO as a rapid, effective sterilant which requires no electrical power so that it can be used in austere environments.

DETAILED DESCRIPTION OF THE INVENTION

[0038] FIG. 1 shows the device 10 in its most simplistic form. The device 10 is comprised of the subcomponents that include the sterilization chamber (SC) 12, the gas generation chamber (GGC) 14, a connecting tube 16 that allows gas to flow from the GGC 14 to the SC 12, and a safety valve 18, along the length of the connecting tube 16 that separates the GGC 14 from the SC 12. The SC 12 has a closure 20, a connecting port 15, and an exhaust port 22 that attaches to exhaust tubing 29. An exhaust valve 23 is attached to the exhaust tubing 29. The GGC 14 contains the powdered sterilant gas-generation composition or compound 24, which is described in detail below. The GGC 14 is further comprised of a female lure fitting 17 to which the connecting tubing 16 is attached, and a filling port 21 for addition of liquids. Each subcomponent comprising the device 10 is described in greater detail below.

[0039] Sterilization chamber 12 (SC). FIG. 1 also details the SC 12. The SC 12 includes a physical container 13 comprised of a plastic, a closure 20, that is gas impermeable and allows re-opening and re-sealing of the SC 12 for loading and unloading of the materials to be sterilized, a connecting port 15 that allows a gas-tight seal with connecting tube 16, and an exhaust port 22 that allows the removal of the gaseous sterilant from the SC 12 prior to removal of the sterilized materials. The SC 12 can be comprised of any plastic material that can contain a low molecular weight gas for up to 45 minutes. Due to the short duration of the period during which it is necessary to contain the gas, semi-gas permeable materials may be used to construct the SC 12, allowing for optimization of weight, toughness, and cost parameters for each individual application. Plastics used for the physical container 13 of the SC 12 may include highly chemical resistant polymers, such as but not limited to C-FLEX, Chemfluor 367, EPDM, ETFE, Kynar, MFA, PEEK, PFA, FEP, polyimide, PVC. The closure 20 may be located at a variety of locations in the SC, preferably, at a point on the SC 12 opposite from the connecting port 15 and the exhaust port 22, or towards either side of the SC 12. The closure 20 is preferably constructed from polyethylene. One preferred closure is one having interlocking linear fasteners that is resistant to pressure breaches, such as the U-MAXIGRIP (Illinois Tool Works, Inc. Glenview, IL). While many interlocking linear fasteners are available, this model is particularly desirable due to its resistance to pressure breaches.

[0040] An alternate embodiment for the SC 12 is shown in FIG. 2, in which the SC 12 uses a flap closure 30 that is a C-shaped track of interlocking plastic hooks and ribs, sealed or opened by using a hinged zipper-like tab 31 and guide slot 32 that separates or compresses the interlocking components, resulting in the opening or sealing of the SC 12. The flap closure 30 is positioned 1 to 2 cm from the perimeter of the SC 12 over three contiguous sides of the SC 12, allowing for a flap of plastic from the SC 12 to be pulled back for easy loading and unloading of the SC 12.

[0041] The connection port 15 allows for a gas-tight connection between the SC12 and the connecting tube 16. A preferred embodiment includes a female lure fitting 25 at the top of the connection port 15 whereby the end of the connecting tube 16 can be comprised of a male lure fitting 27 or a tapered shaft designed to

snugly fit the female lure fitting 25. Alterations in the configuration are well within the skill of the art, such as having the lure fitting at the top of the connection port 15 be a male lure fitting and having the end of the connecting tube 16 be a female lure fitting.

[0042] In one embodiment, the exhaust port 22 is a plastic flange, that is a contiguous unit with the exhaust tube that flanges out from the SC 12 and tapers into a length of exhaust tubing 29. Exhaust tubing 29 is fitted with a valve 23 that, when closed, seals the SC 12 from the ambient air. In a preferred embodiment, the valve 23 is a roller-activated compression valve are many possible embodiments for means to seal the exhaust tubing 29, known to those of skill in the art.

[0043] The connecting tube may be made of any flexible plastic that is relatively resistant to chemicals. Preferred plastic materials include, but are not limited to, C-FLEX, Chemfluor 367, EPDM, ETFE, Kynar, MFA, PEEK, PFA, FEP, polyimide, PVC. The length of the connecting tube should be sufficient to allow the user to freely manipulate each chamber without disturbing the other chamber. Typically, a length of 20 to 30 inches of connecting tube 16 is preferred, however lengths outside the range of 20 to 30 inches are also be functional. At either end of the connecting tube 16 there is a male lure fitting. Alternatively, there is a tapered hard plastic tip that can be inserted in the female lure fitting to insure a fluid-tight seal.

[0044] A broad range of safety valves 18 may be used to separate the GGC 14 from the SC 12 including, but not limited to, crimp valves, roller-activated compression valves, and the like. Any valve that can seal the flow of fluid from the GGC 14 to the SC 12, A preferred embodiment of the present invention employs an air-venting/vacuum breaking valve because it is self-activated.

[0045] Gas Generation Chamber (GGC) and gas generation compound

The GGC 14 includes a container 19 that can be comprised of a variety of plastics that are chemically resistant. These may include but are not be limited to C-FLEX, Chemfluor 367, EPDM, ETFE, Kynar, MFA, PEEK, PFA, FEP, polyimide, PVC. In a preferred embodiment, the container is comprised of PFTE and/or polyolefins. The GGC 14 includes a female lure fitting 17 integrated for attachment of the GGC 14 to the connecting tube 16, which allows a contiguous flow of fluid from inside the GGC 14 to the SC 12. Preferably, the filling port 21

of the GGC 14 is a large, capped opening, which has a threaded rim protruding at least 0.5 cm above the wall of the GGC to allow easy grasping and capping.

[0046]

Another embodiment of the present invention is presented schematically in FIG. 3. The sterilization device 100 includes a hard case with internal gas pumping and scrubbing means that is attached to a sealable gas generation chamber 102. The device 100 is in fluid connectivity with a gas generation chamber 102, through a sealable port 103. In a preferred embodiment, the sealable port 103 may be comprised of a double shut off quick disconnect coupling (Colder Products St. Paul, MN) where the tubing 104 from the gas generation chamber 102 has the male of female coupling and the sealable port 103 is comprised of the complimentary coupling. The advantage of the double shut off feature is that disconnection is possible without opening either the gas generation chamber 102 or the sterilization chamber 101 to the local environment. Thus, the sterilant gases are contained within the sterilization chamber 101, so that any residual gas from the gas generation chamber 102 will be contained until the scrubbing step. The device 100 has a compartment that is comprised of an electronic or hand operated pump 105 that can be in fluid connectivity to the sterilization chamber 101, or not, depending on the position of an intake valve 106. The intake valve 106 may be manually operated, or microprocessor 110 controlled. The intake valve 106 will allow the pump 105 to remove gas contained in the sterilization chamber 101 and the gas generation chamber 102 if it is in fluid connectivity with the sterilization chamber 101 at the time of pump activation. The gas is pumped through a scrubbing system 107 that deactivates and removes the gases from the exhaust stream. The compartment that comprises the inner lumens of the pump 105 and the scrubbing system 107 may or may not be fluid connectivity with the sterilization chamber 101 depending on the position of the intake valve 106. The device 100 is designed so that after completion of the sterilization cycle, activation of the valves and pump 105 draws gas from the sterilization chamber 101 through the intake valve 106 into the scrubbing system 107, through an exhaust valve 108 that directs the flow of gas out of the device 100, or back into the sterilization chamber 101 for the purpose of cycling the gas through the scrubbing system 107 for an additional period of time in order to reduce the levels of gas to OSHA or other regulating agency standards or

guidelines. During such a recycling of gas, one skilled in the art will realize that the gas that re-enters the sterilization chamber will need to pass through a sterile air filter 109 to insure that any potential microbial contaminants picked up by the gas stream in the pump 105, scrubbing system 107, and the necessary tubing to maintain fluid connectivity between the sterilization chamber 101 and these elements, will not enter the sterilization chamber 101 during the gas recycling process.

[0047] One skilled in the art will realize that there are certain risks inherent with gaseous NO that require specialized delivery methods and handling procedures. Exposure to high concentrations of NO is hazardous. The Occupational Health and Safety Administration (OSHA) has set the current level of NO that poses an Immediate Danger to Life and Health at 100 parts per million (ppm) for a maximum of thirty minutes before the effects of exposure would pose a threat to health or life. OSHA has also set the levels of NO in the workplace at 25 ppm time weighted average for eight hours. Because of the dangers of potentially lethal doses of NO, any device or delivery system must include features to prevent the leakage of NO into the surrounding environment in a manner and at levels that may raise a risk that the leaked NO might be inhaled or otherwise applied to subjects that may be harmed by such exposure. The formation of nitrogen dioxide also represents a severe health hazard. OSHA limits for NO₂ are 1 ppm time weighted average over eight hours.

[0048] The system of the present inventions limits the user's exposure to the gases. The system and methods of the present invention include a system that can remove and/or detoxify the sterilant gases, otherwise known as scrubbing. The method of the present invention preferably includes a scrubbing process that removes and detoxifies these gases, prior to retrieving the sterilized or disinfected materials from the sterilization chamber. The scrubbing process, includes numerous methods for removing and detoxifying the NO, NO₂, and NO_x. Scrubbing systems and processes may employ an adsorbant to trap NO, and an oxidizer to convert NO to NO₂. In appropriate conditions, the sterilant gas may be exhausted to the outside environment, where the concentrations of NO, NO₂, and NO_x will be easily dissipated. The scrubbing process may be achieved using a commercially available scrubbing device, such as the Buchi Analytical B-414

(New Castle, DE). Ideally, the scrubbing device reduces the levels of NO, NO₂, and NO_x in the exhaust gas to levels below the OSHA LTWA. Also, see, Basile R. Dealing with Nitrogen Oxide Emissions. <http://www.finishers-management.com/may2002/nox.htm>. It is also preferred that the method act rapidly.

[0049] The method of the present invention most preferably does not expose the user to concentrations of NO, NO₂, and/or NO_x that are above the OSHA limits. In a preferred embodiment, the gases are removed from the chamber prior to opening the chamber. In some instances such as outdoor use, the chamber may be opened without prior removal of gases. In order to limit the exposure to the sterilant gases, the system and method of the present invention include a system that can remove or detoxify the sterilant gases, otherwise known as scrubbing.

[0050] Examples 2, 3 and 4 describe embodiments of effective scrubbing systems that use Purakol and Purafil Select (Purafil, Doraville, GA). One skilled in the art will realize that many configurations of a scrubbing system for a mixture of oxides of nitrogen can be designed.

[0051] In one embodiment of the present invention, the sterilization system is lightweight, requires no electrical (including battery) power, and can be completely self-contained. The core of the system of a re-usable, sealable sterilization chamber, a disposable gas generation chamber, and connecting tubing. The re-usable sterilization chamber can be loaded with surgical instruments or other materials to be sterilized, sealed, and connected to the gas generation chamber which is pre-filled with nitric oxide (NO) donors and acidic activators. Water is then added to the gas generation chamber, the chamber is sealed, and the generated gas flows into the sterilization chamber. A gas permeable, liquid impermeable valve separates the two chambers to avoid mixing the contents of the separate chambers. Preliminary results indicate a duration of 5 min is sufficient for sterilization, although a safety margin of an additional ten minutes may prove prudent after additional testing.

Example 1 Sterilization with Varying Quantities of Delivered Nitric Oxide

[0052] A blood storage container (Nexell, Irvine, CA; Lifecell PL732 plastic tissue culture flask) is used as the sterilization chamber. Strips of stainless steel are dipped in 10⁶ CFU/ml sporulated *B. subtilis var. niger* (as determined by

ABS₅₉₅ standardization curves). The strips are allowed to air dry, placed in the sterilization chamber, and then the sterilization chamber is heat sealed. The sterilization chamber is evacuated using a syringe and controlling air flow with the sterilization chamber valves. A known quantity of air is added to the vessel using a graduated syringe.

[0053] An NO-generating donor compound is placed in a 7 cc gas generation chamber. The gas generation chamber is attached to the storage container through luer lock connectors. The liquid activator, 3.0N HCl is added to the gas generation chamber and the generated gas is allowed to flow into the sterilization chamber. After a brief generation period, the gas is sealed in the sterilization chamber using a compression valve.

[0054] Varying quantities of NO gas, namely 10%, 5%, 2.5% and 1 % NO, are tested for their efficacy in the sterilization chamber. The quantity(%) of NO gas generated and added to the sterilization chamber is calculated from the number of moles of NO required to be generated to result in the desired percentage of NO. This calculation uses the Ideal Gas Law and formula weight of the NO gas-generating compound, which in this Example is a diazeniumdiolate NO donor, to determine the mass of NO gas generating compound to be used.

[0055] All percentages tested, including 1 %, are effective at killing 10⁶ CFU/ml sporulated *B. subtilis var. niger* in five minutes, as determined by culturing of the contaminated steel strips in LB at 37°C and vigorous shaking for 48 hours, followed by plating onto agar plates. Controls are identically treated with the exception of the addition of percentages of nitrogen in place of NO. Control stainless steel strips exhibited visible growth after 24 hours of incubation under the stated conditions.

Example 2 Scrubbing of NO, NO_x from a portable sterilization chamber

[0056] After the sterilant gas is used in the sterilization chamber, the gas in the chamber is evacuated to another chamber containing scrubbing media. The evacuated gas is allowed to reside over the scrubbing media.

[0057] Two 300 ml PL732 tissue culture bags (Lifecell PL732 plastic tissue culture flask case, Nexell, Irvine, CA) are connected to each other with tubing. A hose clamp is used to close one bag off from the other. An incision is made in one bag, designated to be the 'scrubbing' bag, into which a pre-measured amount of

scrubbing media (6.0 to 60 grams of Purafil Select and Purakol in a 1:1 mixture) is added to the bag. The incision is then heat sealed. Both bags are evacuated with a syringe. Air (180 cc) is injected into the bag designated to be the sterilization chamber. Thereafter, 20 cc of NO gas is injected to reach a final concentration of 10% NO. The mixture of NO and air is allowed to remain in the sterilization chamber for 5 minutes. Thereafter, the hose clamp is removed and the sterilization bag is compressed to push all of the NOx gas into the scrubbing bag containing the Purafil Select and Purakol. The hose clamp is then secured. Immediately thereafter, samples (0.1 to 1.0 cc) of the atmosphere in the scrubbing bag are taken and injected into an NO detector, which measure the concentration of NO in parts per billion (ppb). Thereafter, 1.0 cc samples of atmosphere in the scrubbing bag are taken at timed intervals and injected into the NOx detector. Results of three successive trials are shown in Table 1. The scrubbing materials need not be changed between successive trials.

Table 1. Scrubbing of NOx gas

Time minutes	Trial 1 [NO] ppb	Trial 2 [NO] ppb	Trial 3 [NO] ppb
0	32556	69685	69094
5	686	nd	999
6	nd	1484	nd
10	76	nd	253
12	nd	102	nd

nd=no data

Example 3

[0058]

This example provides a method of scrubbing NOx by flowing the NOx gas through tubing filled with scrubbing media, which is connected to a container. Tubing (30 inches of 3/8 inch ID and 1/2 inch OD silastic tubing) is filled with 13.3 grams of a 1:1 mixture of Purafil Select and Purakol. Some of the media is crushed in this process. Glass wool plugs are inserted in the ends of the tubing. Each end of the tube is connected to separate plastic tissue culture bag (Lifecell PL732 plastic

tissue culture flask, Nexell, Irvine, CA). One bag includes an inline valve. The bags are evacuated of atmosphere and the valve is closed. One bag is designated the sterilization chamber, and injected with 180 cc of air and 20 cc of NO gas. The gas is allowed to remain in the sterilization back for five minutes. The valve is then opened and the gas pushed through the tubing into the receiving bag. A 0.5 cc sample of the atmosphere in the receiving bag is injected into the NOx detector. The results show that the contents of the receiving bag was 30 ppb NOx, a concentration well below the OSHA guidelines.

Example 4 Scrubbing of NO, NOx from sterilization chamber.

[0059] A sealable case (Pelican Products, Inc. , Torrance, CA) is modified with additional ports comprised of EFC 12 Series Quick Disconnect Couplings (Colder Products Company, St. Paul, MN) and a plastic shelf with a self-sealing gasket edge which divided the case into upper and lower sections of approximately equal volume. The upper section is the sterilization chamber, which has a volume of 20.3 liters (4.5 in by 19 in by 14.5 in). One port into the sterilization chamber is used to introduce a known amount of NO gas into the sterilization chamber, and optionally, allowing for a recirculating flow. An exhaust port on the opposite end of the case is in the disconnected (sealed) state for the steps involving the addition of NO gas and during the 5 minute time period to approximate sterilization cycle time.

[0060] The lower chamber stores the pump, microprocessor and electric components if any, the valves, the scrubbing system, the sterile air filters and, optionally, additional components. The scrubbing system is connected to the exhaust port and includes tubing having a male end of the EFC 12 Series Quick Disconnect Couplings. Distal to the exhaust port, the tubing is connected to a pump (Gast, Benton Harbor, MI; Model DOA-P104-AA; 35 lit/min flow rate), followed by columns that comprise the scrubbing system. One column is filled with Purafil Select (Doraville, GA); the other is filled with Purakol (approximately 200 to 300 grams of material for each column). NO is injected into the upper sterilization chamber and held for a 5 minute period. After 5 minutes, the scrubbing system is engaged by attaching the male end of the EFC 12 Series Quick Disconnect Couplings to the female end of the exhaust port, thus opening the port, and activating the pump. Prior to pump activation, the pump exhaust is reconnected

to the sterilization chamber via the same port that is used to add NO to the sterilization chamber, using tubing ending with a male end of the EFC 12 Series Quick Disconnect Couplings, and also comprised of a sterile air filter (ACRO 50, Pall Corporation, Port Washington, NY). The gas from the sterilization chamber is sampled using a syringe from an in-line sampling vessel fitted with a rubber septum after 1 minute of pumping. The sampled gas is then injected into and quantified by the ThermoEnvironmental (Waltham, MA) 42C chemilluminescence NOx detector. In addition, NO from the NO storage vessel is injected on the machine as a positive control. The system can recirculate, for example, by adding the gas, disconnecting the gas generation chamber, add tubing from the exhaust port, back to the "intake" port where the NO was added originally, and, when the pump is turned on, the gas will recycle through the system.

[0061] One set of experiments is performed in quadruplicate on the device using 1% added NO. After 1 minute of recirculating the gas from the exhaust port back through the intake port (using sterile air filters to eliminate contaminating the sterilization chamber), as described above, sampling the gas content of the sterilization chamber and measuring showed that virtually all of the NO and NOx components are removed. Each of the four samples barely raised the baseline of the NO detector, resulting in a reading estimated to be approximately 2 ppb, far below the OSHA guidelines of 25 ppm for NO and 1 ppm for NO₂.

[0062] Experiments are performed using 5% added NO. One liter of air (5%) is removed from the sealed case prior to addition of 5% NO, so that the experiment is performed at atmospheric pressure. One liter of NO is then added to the sealed sterilization chamber and allowed to remain for 5 minutes. The scrubbing system is then activated as described above. After one minute of gas recirculation, samples showed approximately 4 ppb for NO and Nox, in each experiment, again far below OSHA guidelines. The Purafil Select and Purakol columns were not changed over the course of these 6 experiments.

Example 5

[0063] A glass pressure vessel is connected to a scrubbed NO gas tank source. The pressure vessel is purged five times with Argon gas to eliminate atmospheric oxygen (preventing formation of NO₂•) and an additional three purges of NO are used to ensure a pure NO atmosphere and consistent results of bactericidal activity.

To test the sterilization method, *Bacillus subtilis* var *niger* 9372 is used (after obtaining >80% endospore formation; a standard for ethylene oxide and autoclave test sterilization), as well as organisms commonly found on the epidermis: *Staphylococcus aureus* (str 21769) and *Staphylococcus epidermides* (str 21977), and the enteric organisms: *Klebsiella pneumoniae* (str 21991) and *Serratia marcesens* (str 21140). This particular *Serratia* strain has been found in previous studies to be one of the most resistant bacteria to the bactericidal effects of NO in culture (Rauli et al., 2002).

[0064] The organisms are cultured overnight in BHI. The cultures contained at least 10^8 CFU/ml based on standardized ABS₅₉₅ curves for each organism. Stainless steel coupons, 3 x 1 cm, are dipped in the cultures, and either dried in ambient air first or placed in the pressure vessel still wet from the culture dip. The coupons are exposed to NO gas at atmospheric pressure for decreasing time periods starting with 45 min and working back to 5 min. Control samples are handled identically, except the pressure vessel was gassed with nitrogen.

Table 2. Results from Pressure Vessel Experiments at Five Minute Sterilization Cycle

	<i>Bacillus subtilis</i>	<i>Serratia marcesens</i>	<i>Staphylococcus epidermides</i>	<i>Klebsiella pneumoniae</i>	<i>Staphylococcus aureus</i>
Stainless steel	3/3 Killed	3/3 Killed	3/3 Killed	3/3 Killed	3/3 Killed

[0065] The sealed vessel is carefully opened in a laminar flow hood after all the NO had been purged with Argon. The samples are removed aseptically with sterilized tongs and placed in culture tubes containing sterile BHI media. The samples are incubated in a vigorously shaking waterbath at 35°C. The samples are observed (digitally photographed) 24 hrs later, placed back in the waterbath and are measured for absorbance 72 hrs later. The controls had > 10^8 CFU/ml after 24 hrs. The results shown in Table 2 are from three separate experiments and the results (3/3) indicate that 3 out of 3 trials showed no bacterial growth.

Example 6

[0066] Similar to Example 1, a portable system is devised using blood storage containers and other laboratory items. In this construct, a blood storage container (Nexell, Irvine, CA; Lifecell PL732 plastic tissue culture flask) serves as the sterilization chamber. It has multiple ports, is easily attached to tubing or other chambers, and is easily cut and heat sealed for insertion and removal of contaminated/sterilized samples. The heat seal is strong and holds well at pressure, even though 1 ATM of pressure is used experimentally. Two 60 ml syringes connected to each other and a line of tubing by a three-way stopcock are employed to mix acidic buffer in one syringe with NO-releasing diazeniumdiolate in the other syringe. The tubing is connected to the blood container/sterilization chamber.

[0067] The stopcock is turned so that the acidic buffer can be added to the syringe containing the diazeniumdiolate. The valve is immediately closed to the buffer syringe and opened to the sterilization chamber. The 300 cc sterilization chamber inflates in about 15 seconds. Experiments are performed as described above, except that the devised system is used instead of the pressure vessel.

[0068] The test organisms are cultured overnight in BHI. The cultures contained at least 10^8 CFU/ml (100-fold greater than FDA testing guidelines) based on standardized ABS₅₉₅ curves for each organism. Stainless steel coupons, 3 x 1 cm, are dipped in the cultures, and either dried in ambient air first or placed in the pressure vessel still wet from the culture dip. The dried samples are dipped in sterile BHI media before being placed in the sterilization chamber. This prototype is shown to exhibit bactericidal activity against wet dipped stainless steel coupons contaminated with *B. subtilis* (endospore), *B. subtilis* (vegetative), *S. marcesens*, and *S. epidermides* in 15 minutes and it may be possible to achieve sterilization in less time.

[0069] **Table 3. Results from Fifteen Minute Sterilization Cycle**

	<i>Bacillus subtilis</i> (spore)	<i>Bacillus subtilis</i>	<i>Staphylococcus epidermides</i>	<i>Serratia marcesens</i>

Stainless steel coupon	2/2 Killed	2/2 Killed	3/3 Killed	3/3 Killed
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Example 7**[0070]**

An additional study is performed to test medically relevant materials such as needles and plastic tubing. Teflon (1/8' ID), polyethylene (1.77 mm ID), vinyl (0.5 mm ID) tubing and a 30 ga. disposable needle are dipped in a bacterial cocktail containing *B. subtilis*, *S. marcesens*, and *S. epidermides* at about 10^8 total CFU/ml. The samples are placed in the sterilization chamber and sealed. In each case, the lumen of the tubing or contained at least some visually confirmed inoculum. Table 4 shows the results from the study. The controls for each material reached at least 10^6 total CFU/ml in 24 hrs as determined by ABS₅₉₅ standardization curves.

Table 4. Sterilization of Medical Materials

	Vinyl Tubing 0.5 mm ID	Polyethylene Tubing 1.77 mm ID	Teflon Tubing 1/8' ID	Disposable Needle 30 ga.
15 min Sterilization Cycle	2/2 Killed	2/2 Killed	2/2 Killed	2/2 Killed

Example 8 Humidity Effects**[0071]**

Several humidifying paradigms and the ability to sterilize through a gas sterilization seal pouch is tested. A cocktail of bacteria is used, all grown to about 10^8 CFU/ml and mixed in equal volume. Stainless steel coupons are dipped, allowed to dry, and subject to one of three methods. Method A samples are wrapped in a moist Kimwipe, Method B samples are left dry, and Method C samples are sealed, dry, in a V. Mueller™ Dual Peel Seal Pouch, designed for gas and autoclave sterilizations. Samples from methods B and C are placed in the

sterilization chamber with a moist Kimwipe, which is placed in the chamber so as to insure maximal separation of the Kimwipe and the sample to be sterilized. The chamber is re-sealed carefully so as not to disturb the positioning of the samples relative to the Kimwipe. The samples are subject to a 15 min sterilization cycle at 1 ATM, removed under sterile conditions, and the samples are placed in BHI media as described above. Control samples are handled identically except for the chamber being gassed with NO. The results are shown in Table 5. All controls reached greater than 10^6 total CFU/ml in 24 hrs as determined by ABS₅₉₅ standardization curves.

Table 5. Effect of Moisture on Sealed and Unsealed Dry Samples

	Method A Moist Kimwipe Wrap	Method B Dry sample	Method C Dry, Sealed Sample
15 min Sterilization cycle	Killed 2/2	Killed 2/2	Killed 2/2

[0072] This experiment suggests two highly significant findings. One is that the samples do not necessarily need to be wet, as a small amount of moisture from a Kimwipe was able to achieve a sterilization from dry samples within the chamber. The second key finding is that the sterilization can occur in individual wrappers that can preserve the sterility of the instrument after the chamber is opened. Our results with this sterilization method to date show a promising, lightweight method that requires no electrical power and is highly transportable.

Example 9 Testing of powdered acids

[0073] A preferred sterilant gas-generating composition is comprised of a nitrogen-based diazeniumdiolate and oxalic acid. Addition of the oxalic acid in a 10:1 molar ratio with the diazeniumdiolate provides produces the sterilant gas, NO, from the diazeniumdiolate, filling a blood storage container in about 20 sec. This capability eliminates the necessity to add 3N HCl to the diazeniumdiolate to generate NO, instead allowing the addition of water to activate the release of NO

gas. The elimination of the need for acid makes the device significantly more convenient to ship, store, and use.

[0074] A disposable, plastic gas generation chamber that can be pre-filled with a carbon-based diazeniumdiolate (nitrogen-based diazeniumdiolates can possibly decompose to form carcinogenic nitrosamines) plus an activating powdered acid, have a large, capped opening to ease the addition of water, and have appropriate attachment lines to transport the gas into the sterilization chamber. Other utility lines or ports may be subsequently added to pull a vacuum if necessary, and to release NO gas from the chamber (through a reusable NOx scrubbing system).

[0075] A polyolefin material chosen for its flexibility, puncture resistance, light weight, and ease of manufacture. The size is approximately a flat 10 inch square. The sterilization chamber's bottom edge will have a tabbed "Ziploc" like re-sealable opening, allowing quick and easy loading of instruments and re-sealing. After the user places the instruments in the pouch, the top portion of the pouch is sealed with a simple quick motion of the tab, resulting in a complete gas tight seal.

[0076] One edge of the pouch sterilization chamber will have an embedded tubing port and approximately 10 inches of tubing to provide a connection to the gas generation chamber. The end of the delivery tube will have "quick disconnect" fittings to facilitate easy connection to the gas generation chamber, and each piece of tubing will have compressing roller valves to seal the tube.

[0077] The chamber will also be made of a polyolefin material, be 3.5 inches square, and have a large hard plastic screw cap protruding from the top side of the container for easy filling of powders and water. The chamber will have a lure lock port to allow easy connection to the sterilization chamber.

Example 10 Synthesis of a soluble, carbon-based diazeniumdiolate

[0078] While a variety of nitrogen-based diazeniumdiolates are commercially available and would work in this application, the ability of nitrogen-based diazeniumdiolates to form highly carcinogenic nitrosamines limits their use in medical applications (Parzuchowski et al., 2002). The carbon-based diazeniumdiolates cannot form nitrosamines and can produce up to three times more NO on a molar basis than nitrogen-based NO donors. By using a carbon-based NO donor, the margin of safety for the product is increased while decreasing the total weight.

[0079] A carbon-based diazeniumdiolates can be produced through the use of a benzylic intermediate. Benzyl methyl ether, $\text{PhCH}_2\text{OCH}_3$ (commercially available from Sigma-Aldrich, St. Louis, MO) is one starting material. In a Parr pressure vessel, 3 ml (0.024 moles) of benzyl methyl ether is added to 30 ml of methanol. To this solution, 11 ml (0.048 moles) of 25% sodium methoxide is added with stirring. Oxygen is removed from the flask by alternating cycles (10) of inert gas pressurization and venting. The solution will then be exposed to between 40 and 80 psi NO gas at room temperature for 1 to 5 days. When the reaction is complete, i.e. when no more NO gas is being consumed, then the head space is purged of NO gas. Diethyl ether will then be added to precipitate out all of the anionic diazeniumdiolated salts, which may then be filtered and dried. The product, $\text{PhC}(\text{N}_2\text{O}_2\text{Na})_2\text{OCH}_3$ is tested for its ability to release NO using a chemilluminescent method described below as well as subject to structure verification by spectrophotometry, elemental analysis, and NMR confirmation.

[0080] An alternate synthetic scheme can be derived from the commercially available benzyl thiocyanate (PhCH_2SCN , Sigma-Aldrich, St. Louis, MO). In a Parr pressure vessel, 3 g (0.020 moles) of benzyl thiocyanate is added to 30 ml of tetrahydrofuran. To this solution, 40 ml (0.040 moles) of 1.0 M sodium silanolate is added with stirring. Oxygen is removed from the flask by alternating cycles (10) of inert gas pressurization and venting. The solution is then exposed to between 40 and 80 psi NO gas at room temperature for 1 to 5 days. When the reaction is complete, i.e. when no more NO gas is being consumed, the head space is purged of NO gas. Diethyl ether is then be added to precipitate out all of the anionic diazeniumdiolated salts, which may then be filtered and dried. The product, $\text{PhC}(\text{N}_2\text{O}_2\text{Na})_2\text{SCN}$ is tested for it ability to release NO using a chemilluminescent method described below as well as subject to structure verification by spectrophotometry, elemental analysis, and NMR confirmation.

[0081] Preferred sterilant gas-generating compounds are these carbon-based diazeniumdiolate compounds for generating NO because their NO release rate under acidic conditions rapid and close to identical. The likely criteria for choice of NO donor is yield and cost.

Example 11 Determination of NO release from diazeniumdiolates

[0082] The method will follow Smith et al. (1996). Weights of the samples is recorded and placed in 0.1 M phosphate buffer (pH 7.4) and the mixture is allowed to stand open to the air at 25°C in a water bath. The buffer is then purged with argon gas via a fritted glass tube at the bottom of the vessel, such that the gaseous effluent gas is passed through a chemilluminiscent NO_x detector calibrated to measure NO content. Bubbling is continued until a steady and horizontal trace is achieved, whereupon the signal is integrated over a span of several minutes. The number of integral units are converted to a value for moles of NO by comparisons with integrals obtained for certified gaseous standards of NO in helium (MG Industries, Morrisville, PA). The rate of NO release over that time increment, calculated by dividing the integrated signal by the number of minutes the integration is conducted, are plotted versus the total elapsed time since the sample was first placed in the buffer.

Example 12 Environmental containment of NO

[0083] All experiments involving NO are performed in a certified fume hood. NO is an environmental pollutant and may be harmful to humans at concentrations higher than 100 ppm. NO contained in synthesis vessels or in sterilization chambers are bled into a container that holds 10-fold the volume of ambient air for a duration of 5 min. This step serves to turn all of the NO into NO₂. The NO₂ from the chamber will then be passed over a column of NaOH which effectively scrubs out the NO₂. This is a well characterized method commonly used in industrial processing (Basile, 2002).

Example 13 Optimization of Sterilization Conditions

[0084] The following parameters are optimized: cycle duration, ambient temperature, percent air allowable, humidity, internal pressure (amount of NO), and instrument loading parameters (surface area of instruments, types of instruments [i.e. narrow lumens, dead end lumens], use of pre-packaged materials in sterilization pouches, salt-crusting instruments, protein crusted instruments). The biological indicator organism chosen for testing is sporulated *Bacillus subtilis var niger*, a standard organisms for Et₂O process validation and commonly used for other sterilization process validation (Hoxey et al., 1985).

[0085] *B. subtilis var niger* 9372 is cultured overnight in Luria Broth (LB) media at 37°C in a shaking waterbath. This usually results in a culture of greater than 10⁸ CFU/ml. The ABS₅₉₅ is measured for each overnight culture and compared to a standard curve to determine the approximate CFU/ml. The density of the cultures is adjusted to 10⁶ CFU/ml by dilution with sterile LB. The bacillus will then be sporulated according to the following method. The cultures are centrifuged at 2500 RPM (1000 x g, Sorvall GLC-1) for 5 min and resuspended in a low nutrient salt media as described by Sterlini and Mendelstam (1969). The bacteria are washed twice more in this sporulation media, and the final pellet suspended in an appropriate amount of sporulation media to retain a density of 10⁶ CFU/ml. This method typically results in greater than 80% endospore formation.

[0086] Paired stainless steel coupons, Teflon® tubing sections 1" long, 1/8" I.D., and (polyethylene)terephthalate (PET) coupons are used for general studies of the Sterilization Cycle Parameters studies ("Materials Panel"). Items from the Materials Panel are dipped in 10⁶ CFU/ml of the endosporulated bacillus suspension. Paired material samples are handled identically, with the control materials being placed in a sterilization chamber and exposed to nitrogen under the same conditions as the group undergoing sterilization with NO gas. After processing, the materials are placed in LB media for incubation at 37°C in a shaking waterbath for 24 hrs. The culture broth for control and processed groups are observed visually and photographed at 24 hrs. An aliquot is removed, serially diluted with sterile LB, and plated onto LB agar plates to determine the CFU/ml. The culture is incubated for an additional 24 hrs to total 48 hrs, and if needed the ABS₅₉₅ is measured at 48 hrs (along with additional confirmatory photographs). The inoculated LB agar plates are incubated for 37°C and assessed for colony growth 24 and 48 hrs after plating.

[0087] Any plate colonies that arise from materials that have been through the sterilization process is tested to confirm their identity as *B. subtilis var niger* through morphological, Gram stain and/or other necessary means. The same confirmatory step applies to any cultures in LB that turn positive. Any tested parameter that results in material(s) that exhibit a *B. subtilis var niger* positive culture after being exposed to the sterilization process is considered a parameter beyond the usable range.

Use of dried spores as a biological indicator organism

[0088] The method of Doyle and Ernst (1967) is used to obtain purified spores from the bacillus. Briefly, heat shocked spores (65°C 30 min) is added to LB and cultured overnight with vigorous shaking at 32°C. This culture is used to inoculate large flasks of LB. These cultures are incubated for 4 days at 32°C. The sporulated culture is kept at 45°C overnight to allow for autolysis of the vegetative cells. The spores are harvested by centrifugation, washed 8 times in distilled water, and checked by phase microscopy to assess purity. Spores are stored as a lyophilized powder for long term storage. Each batch is cultured to insure the viability of the batch before use in experiments.

[0089] Once the limits for each parameter have been found using sporulated *B. subtilis* inocula, the same limits, along with the condition of most likely use, is challenged by inoculating with a suspension of purified spores (10^6 CFU/ml). While no differences are expected, in the interest of thoroughness, the dried spores are tested at the found limits and likely use point. Failure to sterilize spores at the established limits for the sporulated bacillus will result in a re-adjustment of the set limits and a re-evaluation of sporulated *B. subtilis* as the biological indicator, with dried *B. subtilis* spores as the replacement.

[0090] Determination of experimental outcomes using dried spores are similar to those described above, and cultures of the inoculated materials are performed as described in the Loading Parameters section below.

Example 14 Assessment of Sterilization Cycle Duration on Sterilization Efficacy

[0091] Sterilization of the materials are tested at 5, 10, 20, 40, 80, and 120 min at room temperature on the Materials Panel. Each processed group will have a control group treated identically with the exception of using nitrogen gas instead of NO. The experiments are repeated three times, and the criteria for successful sterilization at any particular time point is 0 CFU/ml in all three trials. One failure (positive *B. subtilis var niger* culture) in three trials is considered a failure at that measurement therefore setting the limit at the measured parameter.

Effect of Ambient Temperature on Sterilization Efficacy.

[0092] Items from the Materials Panel are dipped in 10^6 CFU/ml *B. subtilis* in LB. An appropriate time point is selected using data from the previous experiment, using the penultimate minimal successful time point (i.e. if 5 min is successful,

them 10 minutes are used. Experiments are performed from -10°C to 50°C in ten degree increments. Should either of the extreme temperatures fail to produce a successful outcome, the temperature is increased or decreased by 10°C and the trial repeated until a successful outcome is reached. Cold experiments are performed in a calibrated refrigeration unit capable of temperatures from -20° to 20° C. Beyond 20°C, the experiments are performed in a standard incubator. The sterilization device components are equilibrated to the test temperature for 20 min prior to the sterilization process trial. Each processed group will have a control group treated identically with the exception of using nitrogen gas instead of NO. A successful outcome at any temperature point is 0 CFU/ml in all three trials. One failure (positive *B. subtilis var niger* culture) in three trials is considered a failure at that measurement therefore setting the limit at the measured parameter.

[0093] One possible interdependency would be the relationship between higher ambient temperatures and NO gas pressure used in the process. It may be reasonably predicted that higher temperatures may result in an equal or greater degree of efficacy with less NO gas pressure. This would not necessarily be a problem. A problem that may surface is the ability to humidify the sterilization chamber under freezing temperature conditions. In this case, the inability to humidify the chamber may impose a limit on utilization of the process in freezing temperatures.

Effect of Humidity on Sterilization Efficacy.

[0094] While many gas sterilization methods require some degree of humidity, preliminary data (performed at ambient levels of humidity) indicate high levels of humidity may not be required for this sterilization system.

Determination of reproducible humidity conditions in the sterilization chamber.

[0095] A manufactured sterilization chamber prototype is altered to allow the insertion of a hygrometer probe. The probe is sealed inside the chamber using a non-hardening silicon sealant. A NIST traceable hygrometer (Fisher Scientific) with a range of 2 to 98% relative humidity (RH) is used to measure humidity levels. The calibration of the meter is checked once a week using dedicated nitrogen-gassed chambers containing salt baths mixed to generate RH environments of 11, 43, and 75%.

[0096] Once a method to produce reproducible RH levels in the sterilization chamber have been established, items from the Material Panel is contaminated with *B. subtilis*, allowed to dry in ambient air, placed in the sterilization chamber along with an appropriate weight of water (absorbed on swatches) and the sterilization process is tested along the linear range of RH achievable in increments of 10 to 15% RH. Experiments performed at non-room temperatures is allowed to equilibrate at the test temperature for 20 min prior to beginning the sterilization process. Again, the penultimate minimal effective time point is used. A successful outcome at any RH level is 0 CFU/ml in all three trials. One failure (positive *B. subtilis var niger* culture) in three trials is considered a failure at that measurement therefore setting the limit at the measured parameter.

[0097] Should the sterilization fail at a point in the RH spectrum i.e. fails at 0 % RH but is effective at 15%, additional experiments testing RH between 0 and 15 % is added to attempt to establish a more precise range of effective conditions to determine if a mechanism for humidification or dessication is needed in the sterilization chamber

[0098] Interrelationships include the previously discussed potential relationship between humidity and freezing temperatures. Potential also exists for high humidity and low NO gas pressure to produce an inadequate (diluted) level of nitrous acid, resulting in a reduced sterilization efficacy, if in fact nitrous acid formation is important in the sterilization process. This interdependency is testable by testing the sterilization outcome at the lowest possible NO gas pressure and varying the humidity from low to high.

[0099] A requirement for inclusion of a small amount of ambient air may drastically increase the variation in RH levels in normal use conditions.

Effect of NO gas pressure on sterilization efficacy.

[00100] A low pressure gauge is fitted to the sterilization chamber tubing. A three-way stopcock (lure lock) is adapted to the gauge directly or via a short length of tubing. From there a vacuum can be pulled with a 60 cc syringe or pump if required. The chamber can be sealed with the stopcock, thus maintaining the vacuum. The NO gas pressure used for the sterilization trial is regulated by altering the mass of diazeniumdiolate in the gas generation chamber from the normal level of 6.8 gm per 1000 cc of volume. Sterilizations are attempted using 1.7, 3.4, 6.8

gms (control) of diazeniumdiolate in the gas generation chamber, keeping the 10:1 ratio of oxalic acid throughout the experiment. Dead space, once measured in the new prototype, will also be accounted for. A successful outcome is 0 CFU/ml in three trials. One failure (positive *B. subtilis var niger* culture) in three trials is considered a failure at that measurement therefore setting the limit at the measured parameter.

The effect of ambient air on sterilization efficacy.

[00101] The inclusion or exclusion of ambient air is a critical parameter, as the ultimate mechanism of NO sterilization in this method can include the formation of nitrous acid (HNO_2) on surface condensates. A small percentage of ambient air may be advantageously included in the process. The small amount of O_2 dissolved in a humid condensate can also suffice to produce enough nitrous acid under conditions used in the method of the present invention.

[00102] A low pressure gauge is fitted to the sterilization chamber tubing. A three-way stopcock (lure lock) is adapted to the gauge directly or via a short length of tubing. From there a vacuum can be pulled with a 60 cc syringe or pump if required. The chamber can be sealed with the stopcock, thus maintaining the vacuum. Graduated syringes filled with ambient air can be attached to the stopcock and a known quantity of air can be added to the sterilization chamber. The amount of diazeniumdiolate in the gas generation chamber is adjusted based on 2 mol of NO per mol of diazeniumdiolate and using the Ideal Gas Law to reach what would be 1 ATM in 1 L, or 6.8 gm per liter of volume. Dead space, when it is determined from the produced prototype, will also be accounted for with diazeniumdiolate mass. Volumes of ambient air representing 1, 2.5, 5, 10, 15, and 20% in the sterilization chamber is tested. These trials are performed at 25°C, and 2 relevant extremes of temperature determined from experiments outlined above. Zero growth from *B. subtilis* contaminated items from the Material Panel in three trials is considered a successful outcome. One failure (positive *B. subtilis var niger* culture) in three trials is considered a failure at that measurement therefore setting the limit at the measured parameter. A failure at the low level(s) of ambient air may be an indication that oxygen is required, thus providing further evidence that the mechanism of action for NO in this process may be related to the formation of nitrous acid.

[00103] The potential for an interdependency between ambient air and humidity has been previously discussed.

Effect of insoluble crystal occlusion of sterilization efficacy

[00104] Numerous studies have demonstrated the difficulty, especially with gaseous sterilizing agents, of killing spores occluded within water insoluble crystals (Abbott et al., 1956; Doyle and Ernst, 1967). The method of Doyle and Ernst is used for the production and isolation of spores, occlusion of spores in crystalline calcium carbonate, and recovery of occluded spores for determination of sterilization efficacy.

[00105] Ten ml solution of 1.11 % CaCl_2 is prepared containing 10^6 spores per ml. To this, ten ml of 1.06% Na_2CO_3 is rapidly added and the mixture vigorously shaken. Crystals of Ca_2CO_3 will immediately form, occluding large quantities of spores per crystal. The crystals are washed using distilled water in three 20,000 x g centrifugation steps. The crystals are brought back to 10ml in distilled water and 0.2 % methylcellulose for easy handling. Ten μl of the crystal suspension is blotted onto strips of filter paper, dried at room temperature, and further dried at 90°C for 16 hrs.

[00106] After exposure to the sterilization agent, the strips are placed in 25 ml of sterile 3.0 % NH_4Cl for 3 days at 0°C to dissolve the crystals. The strip and solution will then be placed in a blender, followed by sonication for 5 min. The samples will then be diluted and plated on tryptone glucose yeast extract agar for counting. Zero growth in three separate experiments would be considered a successful outcome.

Sterilization Efficacy in devices with long, narrow and dead-end lumens.

[00107] Many studies have documented the difficulties in reliably sterilizing long, narrow, and dead-end lumens (Alfa, 1997; Rutala and Weber, 1998). To test the ability of this sterilization process to effectively sterilize these types of devices, non-porous Teflon tubing ($\leq 3\text{mm}$ I.D.) are cut into 125 cm lengths and a culture of *B. subtilis var niger* (10^6 CFU/ml) is pushed through the tubing using a 60 ml syringe. The tubing is drained and allowed to air dry. Some tubing is plugged at one end with a tight fitting plug. The gas tightness of the plug will have been previously tested by applying a small amount of air pressure using a 60 cc syringe. Alternate methods of sealing one end of the tubing will include heat sealing,

solvent welding, and clamping. Open or sealed end tubing is coiled with care to not crimp the tubing and placed in the sterilization chamber for processing. After the sterilization process is complete, the tubing is cut into 4 inch sections and placed into sterile culture tubes containing enough LB to completely submerge the tubing sections. Sterilization efficacy is evaluated as described above.

Sterilization Efficacy in instruments with occluded joints.

- [00108] Surgical scissors and forceps are contaminated to beyond the swivel joint by dipping in contaminated broth at 10^6 CFU/ml . The swivel joint is actuated while the instrument is submerged in bacterial broth to allow bacteria to get between the arms of the instrument. The instrument is allowed to air dry, and subject to the sterilization process. Zero growth in three separate trials is considered a successful outcome.

Sterilization of instruments in individual sterilization pouches.

- [00109] Surgical scissors and forceps are contaminated to beyond the swivel joint by dipping in contaminated broth at 10^6 CFU/ml. The instruments are dried and sealed in a V. Mueller™ Dual Peel Seal Pouch (Fisher Scientific) and inserted into the sterilization chamber of the device for processing. After processing, the contaminated forceps are carefully removed from the pouches using sterile technique and under sterile conditions, and placed in a culture flask containing sterile LB media, and sterilization efficacy is assessed as described above. Zero growth in three separate trials is considered a successful outcome. Other items, such as long narrow lumen tubing may also be added to this protocol for study.

- [00110] In describing representative embodiments of the invention, the specification may have presented the method and/or process of the invention as a particular sequence of steps. However, to the extent that the method or process does not rely on the particular order of steps set forth herein, the method or process should not be limited to the particular sequence of steps described. As one of ordinary skill in the art would appreciate, other sequences of steps may be possible. Therefore, the particular order of the steps set forth in the specification should not be construed as limitations on the claims. In addition, the claims directed to the method and/or process of the invention should not be limited to the performance of their steps in the order written, and one skilled in the art can readily appreciate that

the sequences may be varied and still remain within the spirit and scope of the invention.

[00111] The foregoing disclosure of the embodiments of the invention has been presented for purposes of illustration and description. It is not intended to be exhaustive or to limit the invention to the precise forms disclosed. Many variations and modifications of the embodiments described herein will be apparent to one of ordinary skill in the art in light of the above disclosure.

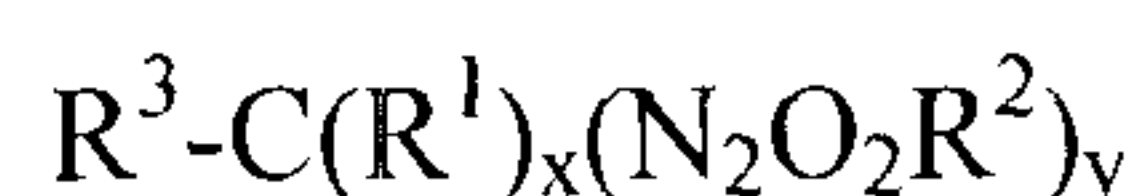
[00112] In describing representative embodiments of the invention, the specification may have presented the method and/or process of the invention as a particular sequence of steps. However, to the extent that the method or process does not rely on the particular order of steps set forth herein, the method or process should not be limited to the particular sequence of steps described. As one of ordinary skill in the art would appreciate, other sequences of steps may be possible. Therefore, the particular order of the steps set forth in the specification should not be construed as limitations on the claims. In addition, the claims directed to the method and/or process of the invention should not be limited to the performance of their steps in the order written, and one skilled in the art can readily appreciate that the sequences may be varied and still remain within the spirit and scope of the invention.

[00113] The foregoing disclosure of the embodiments of the invention has been presented for purposes of illustration and description. It is not intended to be exhaustive or to limit the invention to the precise forms disclosed. Many variations and modifications of the embodiments described herein will be apparent to one of ordinary skill in the art in light of the above disclosure.

WHAT IS CLAIMED IS:

1. A method for sterilizing or decontaminating an object comprising the steps of:
 - placing the object in a sterilizing chamber;
 - generating a sterilant gas comprising a mixture of gaseous NO and gaseous NO₂ in a gas generating chamber comprising sterilant gas generating composition;
 - allowing the sterilant gas to flow from the gas generating chamber to the sterilizing chamber;
 - permitting the sterilant gas to reach a concentration of at least 0.1% in the sterilizing chamber; and
 - exposing the object to the sterilant gas for a time period sufficient to sterilize the desired object.
2. The method as claimed in claim 1, wherein said sterilant gas generating composition comprises a carbon-based diazeniumdiolate compound.
3. The method as claimed in claim 1, wherein said sterilant gas-generating composition comprises a nitrogen-based diazeniumdiolate compound.
4. The method as claimed in claim 1, wherein said sterilant gas-generating composition further comprises an activator, wherein said activator is an acid.
5. The method as claimed in claim 4, wherein said acid is selected from the group consisting of oxalic acid and maleic acid.
6. The method as claimed in claim 1, wherein the gas generating chamber is a pressurized chamber comprising sterilant gas generating composition.
7. The method as claimed in claim 1, wherein the sterilant gas-generating composition requires from about 2 seconds to about 30 seconds to release sufficient quantities of NO to sterilize an object in the sterilization chamber.

8. The method as claimed in claim 1, wherein said carbon-based diazeniumdiolate compound produces quantities of NO that are greater than 1 mole of NO per mole of said carbon-based diazeniumdiolate compound.
9. The method as claimed in claim 2, wherein said carbon-based diazeniumdiolate has a carbon bearing a diazeniumdiolate group, wherein said carbon does not comprise part of an imidate, thioimidate, amidine or enamine.
10. The method as claimed in claim 2, wherein said carbon-based diazeniumdiolate compound is a C-based diazeniumdiolate compound having the formula:



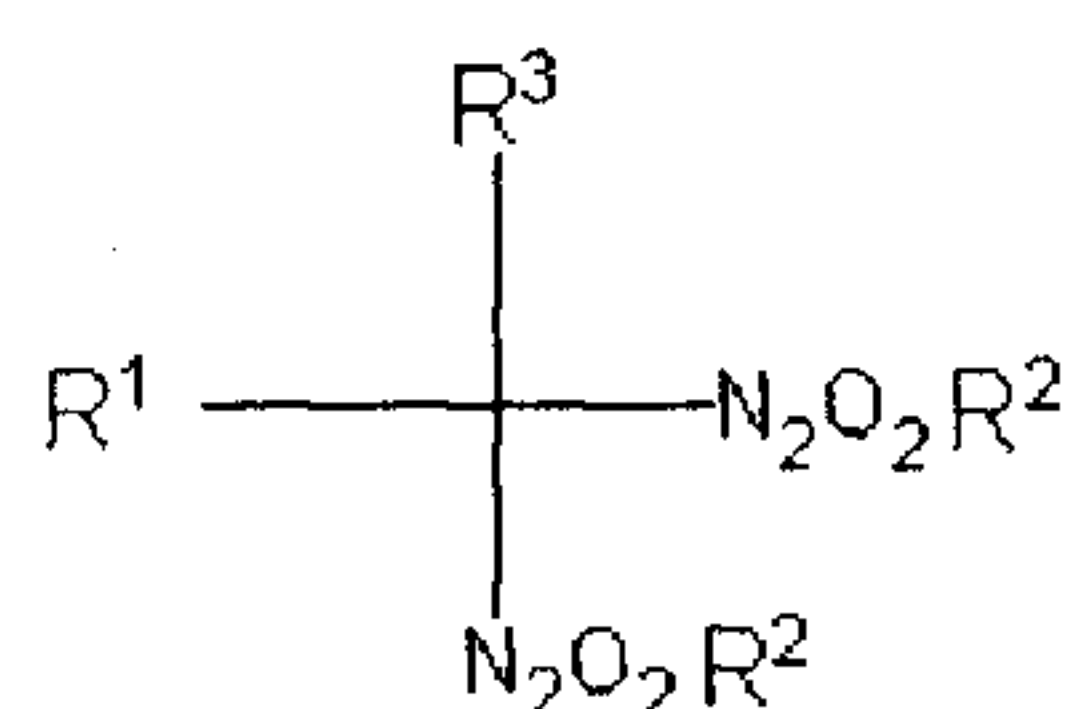
wherein x is an integer from 0 to 2 and y is an integer from 1 to 3 and the sum of x plus y equals 3;

wherein R^1 is not an imidate, thioimidate, amidine or enamine;

wherein R^2 is selected from the group consisting of a counteranion and a protecting group on the terminal oxygen; and

wherein R^3 is a phenyl group.

11. The method as claimed in claim 2, wherein said carbon-based diazeniumdiolate compound has the formula:



wherein R^1 is not an imidate, thioimidate, amidine or enamine;

wherein R^2 is selected from the group consisting of a counteranion and a protecting group on the terminal oxygen; and wherein R^3 is a phenyl.

12. The method as claimed in claim 11, wherein R^1 is selected from the group consisting of an electron withdrawing group, a nitro group, an ether, a thioether, and a non-enamine amine;
 wherein the R^3 substituent is selected from the group consisting of aliphatic, aromatic, and non-aromatic cyclic groups; and
 wherein the R^3 substituent is selected from the group consisting of mono- or di-substituted amino, unsubstituted amino, ammonium, alkoxy, acetoxy, aryloxy, acetamide, aldehyde, benzyl, cyano, nitro, thio, sulfonic, vinyl, carboxyl, nitroso, trihalosilane, trialkylsilane, trialkylsiloxane, trialkoxysilane, diazeniumdiolate, hydroxyl, halogen, trihalomethyl, ketone, benzyl, and alkylthio.
13. The method as claimed in claim 11, wherein the counteraction is selected from the group consisting of ammonium and other quaternary amines;
 and further wherein the protecting group is selected from the group consisting of aryl, sulfonyl, glycosyl, acyl, alkyl and olefinic groups.
14. The method as claimed in claim 1, further comprising activating a gas-generating composition to generate the sterilant gas in the gas generating chamber and wherein the step of activating the sterilant gas-generating composition precedes the step of placing the object in the sterilizing chamber.
15. The method as claimed in claim 1, wherein the sterilant gas-generating composition comprises a sterilant gas-generating compound and an acid compound.
16. The method as claimed in claim 1, wherein said method is conducted at a temperature less than 90°C , at ambient pressure for time period in the range of 2 minute to 30 minute.
17. The method of claim 16, wherein said method is conducted at a temperature in the range of 10°C to 90°C .
18. The method of claim 16, wherein said method is conducted at room temperature.
19. The method as claimed in claim 1 or 16, wherein said time period is from about 4 minutes to about 15 minutes.
20. The method as claimed in claim 1 or 16, wherein said time period is from about 4 minutes to about 10 minutes.
21. The method as claimed in claim 1, wherein said method requires no electricity or other power source.

22. The method as claimed in claim 1, wherein the sterilant gas is generated by reacting the sterilant gas composition with addition of a liquid activator, wherein the liquid activator is selected from the group consisting of: water and acids.
23. A system or device for performing a method of sterilization an object as claimed in claim 1, comprising:
- a. a gas generating chamber;
 - b. a sealable sterilizing chamber, in fluid communication with the gas generating chamber and configured to hold the material to be sterilized;
 - c. a valve, configured and arranged to controllably allow flow between the gas generating chamber and the sterilizing chamber; wherein the gas generating chamber comprising a gas generating composition for producing, a sterilant gas comprising a mixture of gaseous NO and gaseous NO₂.
24. The system as claimed in claim 23, wherein said gas generating chamber comprises a port for introducing a liquid or solid.
25. The system as claimed in claim 23, further comprises a scrubbing system.
26. The system as claimed in claim 25, wherein said scrubbing system comprises an oxidizer to convert NO to NO₂.
27. The system as claimed in claim 23, wherein said scrubbing system comprises an adsorbant to trap NO₂.

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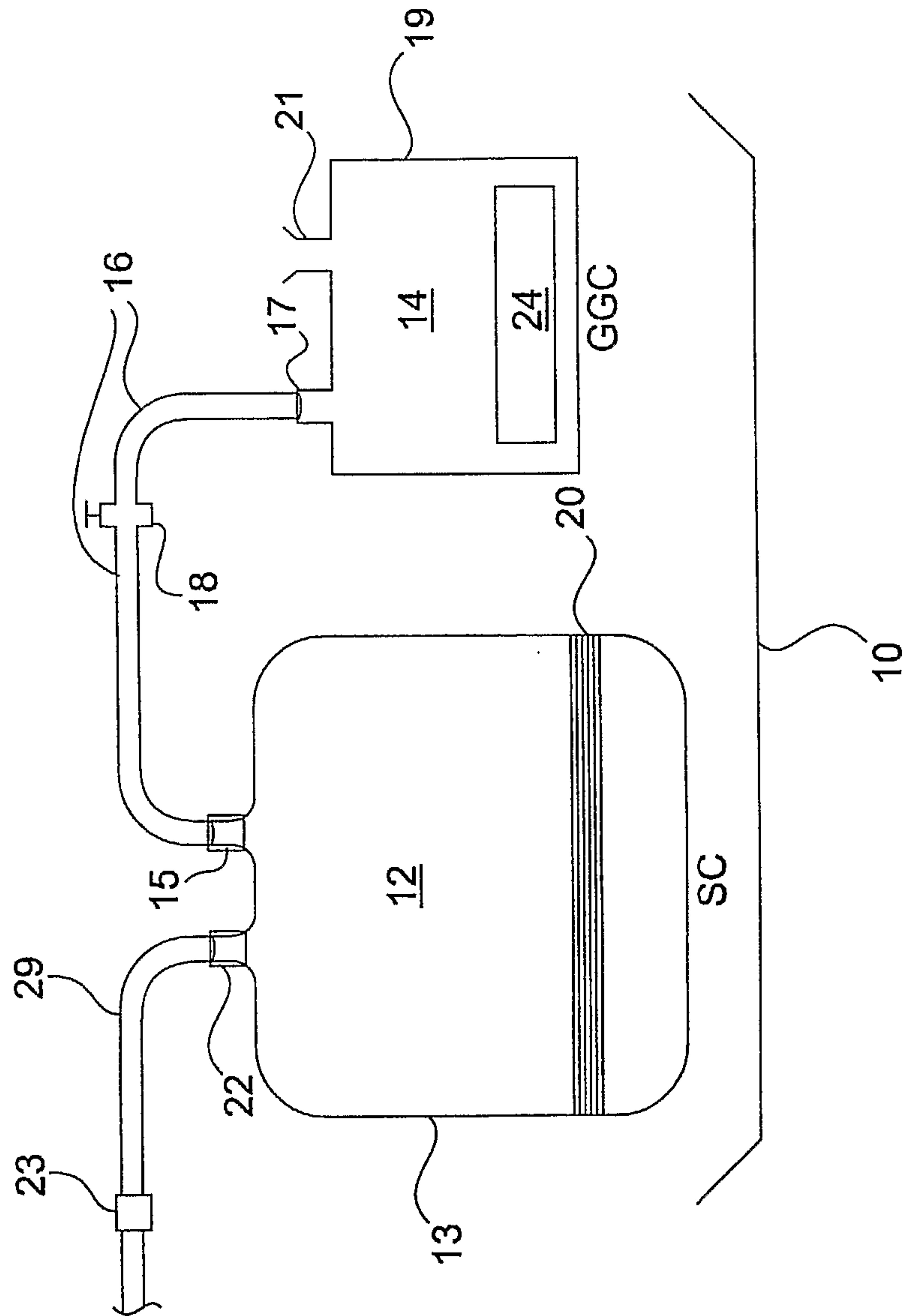
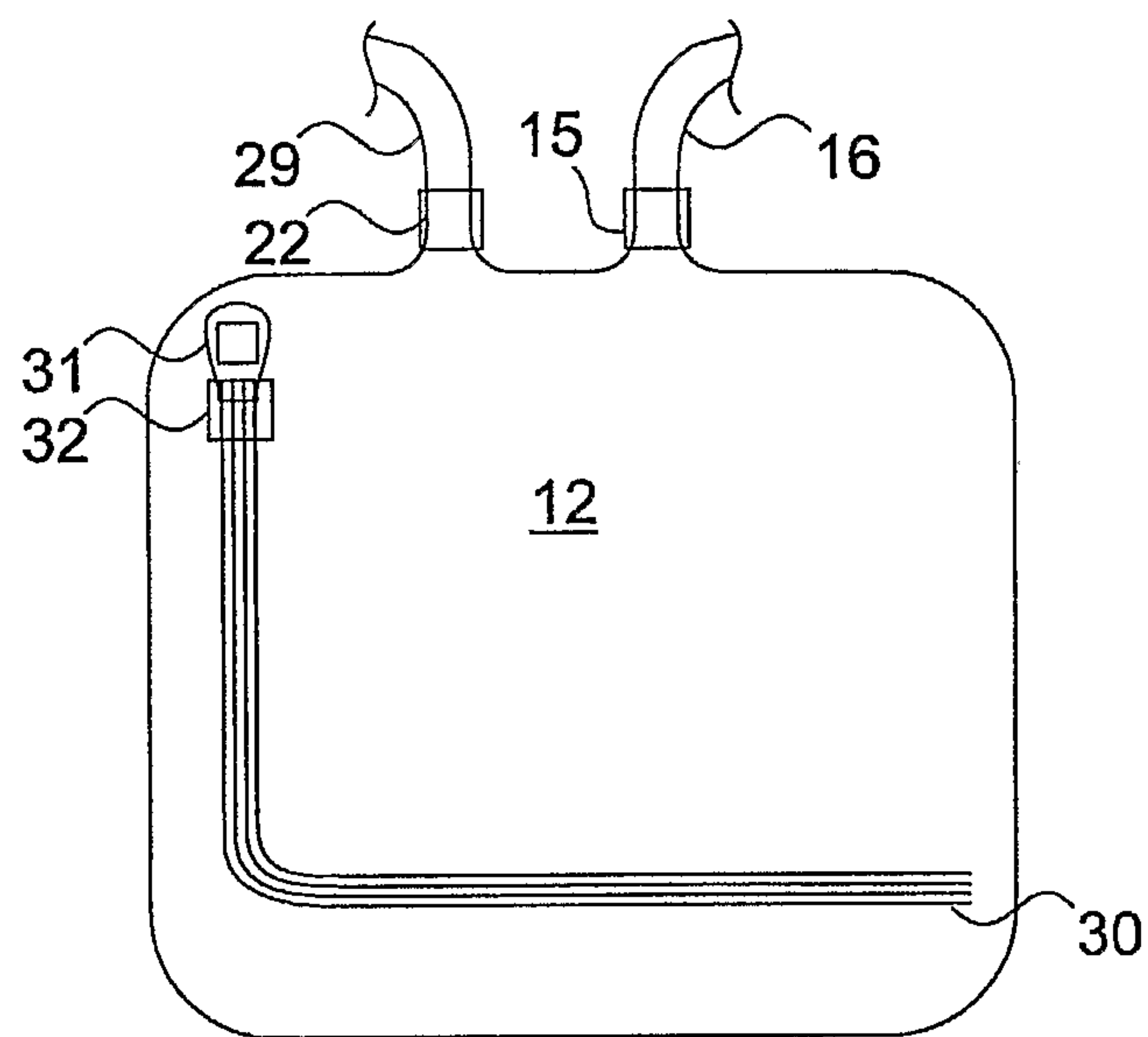


FIG. 1

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**FIG. 2**

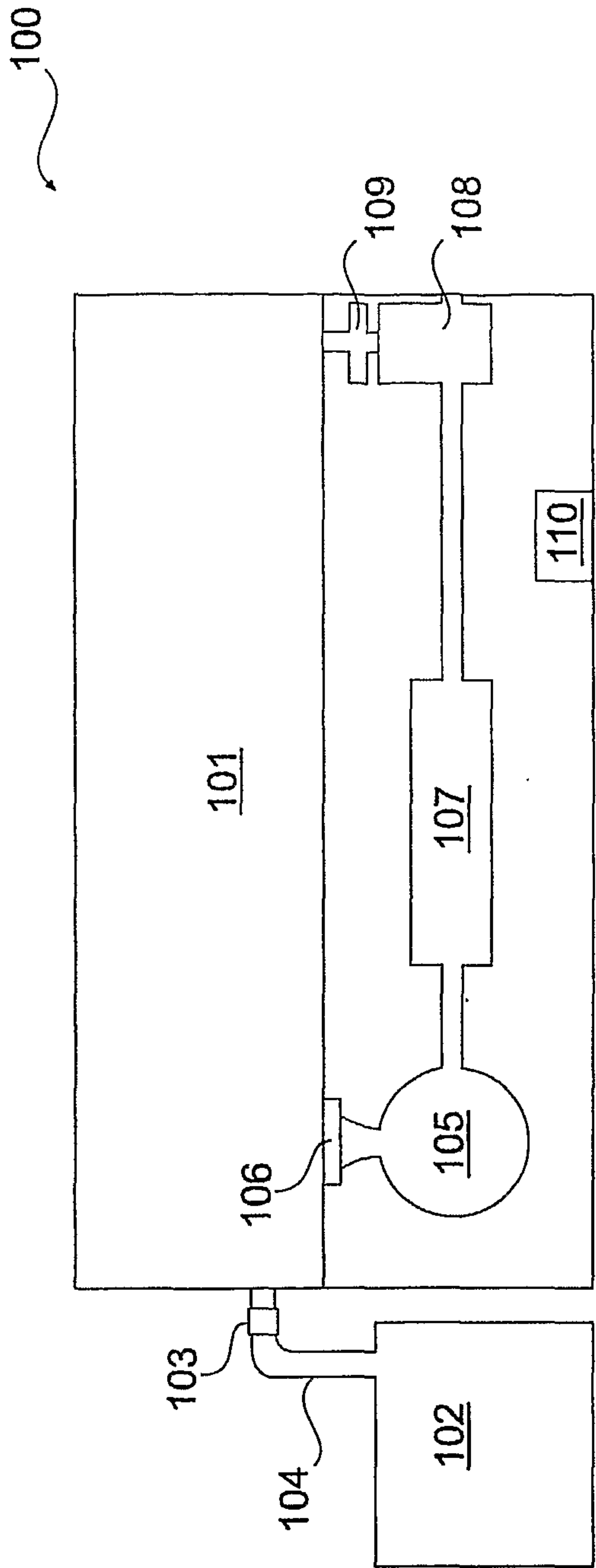


FIG. 3