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Method and apparatus for packaging a drug-device combination product

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(71) Applicant(s)
Codman & Shurtleff, Inc.

(72) Inventor(s)
Destraduer, Alan J.; Gupta, Rainuka

(74) Agent/Attorney
Callinans, 1193 Toorak Road, Camberwell, VIC, 3124

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ABSTRACT

A package for a drug-device combination product includes an outer package including a first gas impermeable sheet and a second gas impermeable sheet hermetically sealed there to on three sides. A gas permeable header is attached to an unsealed side of
5 the first sheet and sealed to the second sheet on two sides. The first and second gas impermeable sheets and the header form an interior and an opening communicating with the interior. A gas permeable inner package is disposed within the outer package. A product is sealed within the inner package. The inner package is placed within the outer package and a top end of the header is sealed to the second sheet. The outer package is
10 then sealed by sealing the first gas impermeable sheet to the second gas impermeable sheet at a seal point below the point where the header attaches to the first sheet.

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Name of Applicant: CODMAN & SHURTLEFF, INC.

Actual Inventors: Rainuka Gupta and Alan J. Destradeur

Address for Service: CALLINAN LAWRIE, 711 High Street,
Kew, Victoria 3101, Australia

Invention Title: METHOD AND APPARATUS FOR PACKAGING A DRUG-
DEVICE COMBINATION PRODUCT

The following statement is a full description of this invention, including the best method of performing it known to us:-

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METHOD AND APPARATUS FOR PACKAGING
A DRUG-DEVICE COMBINATION PRODUCT

BACKGROUND OF THE INVENTION

5 **1. Field of the Invention**

This invention relates to a package for a drug-device combination product having an outer package including a first gas impermeable sheet, a second gas impermeable sheet and a gas permeable header.

10 **2. Discussion of the Related Art**

Packaging a product in an inner package and then packing the inner package in an outer package is common in the packaging arts. Vacuum packaging, packaging with an inert gas and multiple methods of sterilizing medical products are known in the medical device packaging art. However, drug-device combination products offer a new and unique
15 problem in the packaging and sterilizing of the product, while refraining from altering the chemical structure of the drug incorporated in the device.

Numerous inventions relate to the use of radiation to sterilize products for medical use, for example, U.S. Patent No. 5,577,368 to Hamilton et al. ("Hamilton") and U.S. Patent No. 6,174,934 to Sun et al. ("Sun"). Both Hamilton and Sun first remove the
20 oxygen/atmosphere from the packaging prior to radiation sterilizing medical implants made of polymeric material. Hamilton's and Sun's goal is to reduce the wear resistance of the polymeric implant and, radiation typically alters the chemical structures of incorporated drugs.

Other inventions known in the art require numerous complex steps to sterilize and
25 seal a medical device in one or more packages. U.S. Patent No. 4,709,819 to Lattuada et al. discloses first sterilizing an outer package, placing an inner package in the outer package and then evacuating both the inner and outer packages. This process is used because Lattuada et al. are packaging biological samples and any sterilization of the inner package would kill the sample.

30 Further, U.S. Patent No. 4,941,308 to Grabenkort et al. discloses sterilizing the interior of the package before placing the product in the inner package, sterilizing the product in the inner package, and then placing the inner package in the outer package. Additionally, Grabenkort et al. uses ethylene oxide gas (EtO) for the sterilization.

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Both Lattunda et al. and Grabenkort et al. require a separate sterilizing step prior to inserting and sealing the inner package in the outer package. This adds steps and cost to the handling of the already sterilized product/inner package prior to inserting it into the outer package.

- 5 Thus, there is a need in the art for a packaging product and method to sterilize a drug-incorporated device in the minimum number of steps while refraining from altering the chemical structure of the drug incorporated in the device.

SUMMARY OF THE INVENTION

- 10 According to the present invention there is provided a package for a drug-device combination product, comprising: an outer package having a first gas impermeable sheet, a second gas impermeable sheet hermetically sealed on three sides with the first gas impermeable sheet, a gas permeable header disposed at an unsealed side of the first gas impermeable sheet and sealed to the second gas
- 15 impermeable sheet on two sides, wherein the first and the second gas impermeable sheets and the gas permeable header form an interior and an opening communicating with the interior and a gas permeable inner package in the form of a pouch designed to hold said drug-device combination product and be disposed within the outer package.
- 20 The invention also provides a method of packaging a drug-device combination product, including the steps of placing the product inside a gas permeable inner package; sealing the inner package; placing the inner package inside an outer package, wherein the outer package has a first gas impermeable sheet and a second gas impermeable sheet hermetically sealed on three sides and a gas
- 25 permeable header disposed at an unsealed side of the first gas impermeable sheet and sealed to the second gas impermeable sheet on two sides, sealing a top end of the header to the second sheet to seal the inner package in the outer package; sterilizing the product and sealing the first sheet to the second sheet to seal the inner package in a gas impermeable outer package in the form of a pouch. The impermeable sheet
- 30 material should be flexible and can be selected from among many types of high barrier, flexible packaging materials that are commonly used to enclose medical devices. Preferably the impermeable sheet material is a multilayered, heat seal

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- peelable packaging material that includes one or more foil layers, various polymer layers and a heat seal coating. Examples of suitable materials are those that include the following layers: polyester film-low density polyethylen-foil-ionomer-heat seal coating. Packaging materials having the following layers can also be used:
- 5 polyester-low density polyethylene-foil-EAA-linear low density polyethylene-heat seal coating; and polyester-Surlyn-nylon-Surlyn-foil-EAA-linear low density polyethylene-heat seal coating. Additionally, polyvinylidene chloride (PVDC) and ethylene vinyl alcohol copolymer, (EVOH), are generally key components of a high-barrier film. Nylons, acrylonitrile methacrylate copolymer (AN-MA), and other
- 10 specialty polymers such as certain copolyesters may potentially be used.

- The gas permeable material for both the header and/or the inner package can be Tyvek® or any other durable gas permeable material such as polyethylene, polystyrene or polypropylene. The gas permeable inner package can be a blister tray, a pouch, or any other gas permeable container designed to hold a product to be
- 15 sterilized. The product can

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be incorporated with drugs that are antimicrobial agents, antiangiogenesis, antiproliferatives, and anti-inflammatories. Incorporating the drug into the product can include, but is not limited to, impregnating, coating and sandwiching the drug between layers of the device.

5 In an embodiment of the product, the antimicrobial agent can be selected from the group comprising antibiotics, antiseptics, and disinfectants. Further, the antibiotics can be selected from a group comprising tetracyclines (i.e. minocycline), penicillins, (i.e. nafcillin), macrolides (i.e. erythromycin), rifampin, gentamicin, vancomycin, clindamycin, azithromycin, enoxacin, and combinations thereof. A preferred product is incorporated with Rifampicin or Clindamycin.

10 Antiangiogenesis drugs (also called angiogenesis inhibitors) deprive the cancer cells of their blood supply. Antiangiogenesis can be selected from the group including angiostatin, thalidomide (Thalomid™), CC-5013 (Revimid™), bevacizumab (Avastin™), squalamine, endostatin, angiostatin, and angiozyme. Other antiangiogenesis drugs can include drugs derived from chemotherapy drugs, for example, paclitaxel (Taxol™), doxorubicin (Adriamycin™), epirubicin, mitoxantrone, and cyclophosphamide.

Antiproliferative drugs can prevent restenosis (a re-narrowing or blockage of an artery at the same site where treatment, such as an angioplasty or stent procedure, has already taken place) of the implanted device. Examples of antiproliferatives are 20 Sirolimus™ and Paclitaxel™. Examples of anti-inflammatories, including non-steroidal anti-inflammatory drugs (NSAID's), are ibuprofen, ketoprofen, motrin, and naproxen.

Once the product is sealed within the inner package, the inner package is placed within the outer package and a top end of the header is sealed to the second sheet. This forms a sealed outer package that has gas permeable and impermeable sections. Up to this 25 point, only minimal care is required, neither the product, inner package, nor outer package has been sterilized and special handling or handling in a clean environment is not required. However, a preferred embodiment includes handling in a clean room environment to minimize the introduction of contaminants.

The sterilizing compound is then introduced into the interior of the outer package 30 through the header. Since the inner package is also permeable, the sterilizing compound can permeate through the inner package and sterilize the product. As above, the sterilizing compound can be steam, ethylene oxide gas (EtO), gas plasma/radio frequency-peroxide (e.g. Sterrad™), chemical vapor (e.g. alcohol, formaldehyde, etc), and cold sterilization using liquid chemical sterilants/disinfectants that require immersion (e.g. glutaraldehyde

and chlorine dioxide). The invention allows for any sterilization process that utilizes a sterilizing agent that can only pass through a permeable layer.

Once the product is sterilized, the atmosphere can be evacuated from the interior to 'vacuum seal' the product, as well as retard further oxidation. The outer package is then sealed by sealing the first gas impermeable sheet to the second gas impermeable sheet at a seal point below the point where the header attaches to the first sheet. The outer package is now a gas impermeable package. The header can either be removed or folded over to complete the packaging.

A key feature of the sterilizing step includes using any sterilizing process that maintains a chemical structure of the drug as well as prevents oxidation of the drug, and still sterilizes the product for medical use.

BRIEF DESCRIPTION OF THE DRAWING FIGURES

The above and still further objects, features and advantages of the present invention will become apparent upon consideration of the following detailed description of a specific embodiment thereof, especially when taken in conjunction with the accompanying drawings wherein like reference numerals in the various figures are utilized to designate like components, and wherein:

Figure 1 is an exploded perspective view of the outer package of the present invention;

Figure 2 is a perspective view of the outer package of the present invention;

Figure 2A is a cross-sectional view along line 2A-2A of Figure 2;

Figure 3 is a perspective view of the of the assembled outer package prior to the insertion of the inner package;

Figure 3A is a cross-sectional view along line 3A-3A of Figure 3;

Figure 4 is a perspective view of the of the inner package and outer package during sterilization;

Figure 4A is a cross-sectional view along line 4A-4A of Figure 4;

Figure 5 is a perspective view of the inner package and outer package during evacuation;

Figure 5A is a cross-sectional view along line 5A-5A of Figure 5;

Figure 6 is a perspective view of the inner package and outer package after evacuation;

Figure 6A is a cross-sectional view along line 6A-6A of Figure 6;

Figure 7 is a perspective view of the final packaging; and

Figure 8 is a flow chart outlining the method of the present invention.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

5 Referring now to Figures 1 through 3A, a package 100 for a drug-device combination product is illustrated. Package 100 includes an outer package 102 including a first gas impermeable sheet 104 and a second gas impermeable sheet 106 hermetically sealed on three sides 112 (Figure 3). A gas permeable header 108 is attached at an attachment point 109 located on an unsealed side 110 of first gas impermeable sheet 104 and sealed to second gas impermeable sheet 106 on two sides. First and second gas impermeable sheets 104, 106 and gas permeable header 108 form an interior 114 with an opening 116 communicating with interior 114. The impermeable sheet material should be flexible and can be selected from among many types of high barrier, flexible packaging materials that are commonly used to enclose medical devices. Preferably the impermeable sheet material is a multilayered, heat seal peelable packaging material that includes one or more foil layers, various polymer layers and a heat seal coating. Examples of suitable materials are those that include the following layers: polyester film-low density polyethylene-foil-ionomer-heat seal coating. Packaging materials having the following layers can also be used: polyester-low density polyethylene-foil-EAA-linear low density polyethylene-heat seal coating; and polyester-Surlyn-nylon-Surlyn-foil-EAA-linear low density polyethylene-heat seal coating. Additionally, polyvinylidene chloride (PVDC) and ethylene vinyl alcohol copolymer, (EVOH), are generally key components of a high-barrier film. Nylons, acrylonitrile methacrylate copolymer (AN-MA), and other specialty polymers such as certain copolyesters may potentially be used.

25 Referring to Figures 3 through 4A, a gas permeable inner package 118 is disposed within outer package 102. Additionally, gas permeable inner package 118 is preferably disposed only between first and second gas impermeable sheets 104, 106. The gas permeable material for both the header and/or the inner package can be Tyvek® or any other durable gas permeable material such as polyethylene, polystyrene or polypropylene. Gas permeable inner package 118 can be a blister tray, a pouch, or any other gas permeable container designed to hold a product 120 to be sterilized.

Figure 4A illustrates product 120 which, in an embodiment, can be incorporated with a drug (not illustrated) and gas permeable inner package 118 is sized to contain product 120 and be placed only between gas impermeable section of first sheet and second

sheet 104, 106. The drugs that the product can be incorporated with are antimicrobial agents, antiangiogenesis, antiproliferatives, and anti-inflammatory. Incorporating the drug into the product can include, but is not limited to, impregnating, coating and sandwiching the drug between layers of product 120.

5 In an embodiment of product 120, the antimicrobial agent can be selected from the group comprising antibiotics, antiseptics, and disinfectants. Further, the antibiotics can be selected from a group comprising tetracyclines (i.e. Minocline™), penicillins, (i.e. Nafcillin™), macrolides (i.e. Erythromycin™), rifampin, gentamicin, vancomycin/clindamycin, azithromycin, enoxacin, and combinations thereof. A preferred
10 product 120 is incorporated with Rifampicin™ or Clindamycin™.

Antiangiogenesis drugs (also called angiogenesis inhibitors) deprive the cancer cells of their blood supply. Antiangiogenesis can be selected from the group including angiostatin, thalidomide (Thalomid™), CC-5013 (Revimid™), bevacizumab (Avastin™), squalamine, endostatin, angiostatin, and angiozyme. Other antiangiogenesis drugs can
15 include drugs derived from chemotherapy drugs, for example, paclitaxel (Taxol™), doxorubicin (Adriamycin™), epirubicin, mitoxantrone, and cyclophosphamide.

Antiproliferative drugs can prevent restenosis (a re-narrowing or blockage of an artery at the same site where treatment, such as an angioplasty or a stent procedure, has already taken place) of product 120. Examples of antiproliferatives are Sirolimus™ and
20 Paclitaxel™. Examples of anti-inflammatories, including non-steroidal anti-inflammatory drugs (NSAID's), are ibuprofen, ketoprofen, motrin, and naproxen.

Figures 3 through 4A further illustrate that once product 120 is sealed within inner package 118, inner package 118 is placed within outer package 102. A top end 122 of header 108 is sealed to second sheet 106. A sterilizing compound 124 is then introduced
25 into interior 114 through header 108. Sterilizing compound 124 then permeates through inner package 118 and sterilizes product 120. Sterilizing compound 124 can be steam, ethylene oxide gas (EtO), gas plasma/radio frequency-peroxide (e.g. Sterrad™), chemical vapor (e.g. alcohol, formaldehyde, etc), and cold sterilization using liquid chemical sterilants/disinfectants that require immersion (e.g. glutaraldehyde and chlorine dioxide).
30 The invention allows for any sterilization process that utilizes a sterilizing agent that can only pass through a permeable layer and that process is compatible with the incorporated drug. For example, Rifampicin is not compatible with EtO sterilization, but is compatible with steam sterilization.

Referring to Figures 5 and 5A, optionally, atmosphere 126 can be evacuated from interior 114 to 'vacuum seal' product 120. Outer package 102 is then sealed by sealing first gas impermeable sheet 104 to second gas impermeable sheet 106 at a seal point 128 below attachment point 109 of header 108 to first sheet 104. Outer package 102 is now a gas impermeable package. Optionally, header 108 can either be removed or folded over to complete the packaging (Figure 7).

Figure 8 illustrates a method of packaging a drug-device combination product 120 such as, for example, but not limited to a drug incorporated catheter (e.g. Bactiseal™) and a drug-eluting stent (e.g. Cypher™). The steps include placing product 120 inside gas permeable inner package 118 (step 202), sealing inner package 118, and placing inner package 118 inside outer package 102 (step 204). Outer package 102 is formed as described above. Further steps include sealing the top end 122 of header 108 to second sheet 106 to seal inner package 118 in outer package 102 (step 204). Once inner package 118 is sealed, product 120 is sterilized with sterilizing compound 126 (step 206) and the first sheet 104 is sealed to second sheet 106 to seal inner package 118 in a gas impermeable outer package (step 210). An embodiment includes, after the sterilizing step (206), removing atmosphere 126 from inside outer package 102 (step 208). Once the gas impermeable outer package is sealed, another step includes removing the gas permeable header 108 (step 212), or as an alternative header 108 can be folded under outer package 102.

An embodiment includes, replacing atmosphere 126 with an inert gas (not illustrated) prior to removing the atmosphere 128 (step 208). Another embodiment includes, after removing the atmosphere (step 208), filling the outer package 102 with an inert gas (not illustrated).

A key feature of the sterilizing step includes using steam, EtO, gas plasma/radio frequency-peroxide, chemical vapor, cold sterilization or any sterilizing process that maintains the chemical structure of the incorporated drug, prevents oxidation of the incorporated drug and also maintains the desired mechanical properties (rigidity, elasticity, etc.) of the device, while still sterilizing the product for medical use.

While there have been shown, described, and pointed out fundamental novel features of the invention as applied to a preferred embodiment thereof, it will be understood that various omissions, substitutions, and changes in the form and details of the devices illustrated, and in their operation, may be made by those skilled in the art without departing from the spirit and scope of the invention. For example, it is expressly intended

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that all combinations of those elements and/or steps which perform substantially the same function, in substantially the same way, to achieve the same results are within the scope of the invention. Substitutions of elements from one described embodiment to another are also fully intended and contemplated. It is also to be understood that the drawings are not
5 necessarily drawn to scale, but that they are merely conceptual in nature. It is the intention, therefore, to be limited only as indicated by the scope of the claims appended hereto.

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The claims defining the invention are as follows:

1. A package for a drug-device combination product, said package including: an outer package having:
 - 5 a first gas impermeable sheet;
 - a second gas impermeable sheet hermetically sealed on three sides with the first gas impermeable sheet;
 - a gas permeable header disposed at an unsealed side of the first gas impermeable sheet and sealed to the second gas impermeable sheet on two sides;
 - 10 wherein the first and the second gas impermeable sheets and the gas permeable header form an interior and an opening communicating with the interior; and
 - a gas permeable inner package in the form of a pouch designed to hold said drug-device combination product and be disposed within the outer package.
- 15 2. The package of claim 1, wherein the gas permeable inner package is disposed only between the first and the second gas impermeable sheets.
3. The package of claim 1, further including:
 - a drug-device combination product incorporated with a drug;
 - wherein the gas permeable inner package is sized to contain the product.
- 20 4. The package of claim 3, wherein the gas permeable inner package is disposed only between the gas impermeable section of the first sheet and the second sheet.
5. The package of any one of claims 1 to 4, wherein the drug is an antimicrobial agent, antiangiogenesis agent, antiproliferative or anti-inflammatory.
6. The package of claim 5, wherein the antimicrobial agent is an antibiotic,
 - 25 antiseptic or disinfectant.

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7. The package of claim 6, wherein the antibiotic is a tetracycline, penicillin, macrolide, rifampin, gentamicin, vancomycin, clindamycin, azithromycin, enoxacin or a combination thereof.
8. The package of claim 5, wherein the antimicrobial agent is at least one of Rifampicin and Clindamycin.
9. The package of claim 5, wherein the antiangiogenesis agent is angiostatin, thalidomide, CC-5013, bevacizumab, squalamine, endostatin, angiostatin, angiozyme, paclitaxel, doxorubicin, epirubicin, mitoxantrone or cyclophosphamide.
10. The package of claim 5, wherein the antiproliferative is at least one of Sirolimus and Paclitaxel.
11. The package of claim 5, wherein the anti-inflammatory is ibuprofen, ketoprofen, motrin or naproxen.
12. A method of packaging a drug-device combination product, including the steps of:
 - placing the product inside a gas permeable inner package; sealing the inner package;
 - placing the inner package inside an outer package, wherein the outer package has a first gas impermeable sheet and a second gas impermeable sheet hermetically sealed on three sides and a gas permeable header disposed at an unsealed side of the first gas impermeable sheet and sealed to the second gas impermeable sheet on two sides;
 - sealing a top end of the header to the second sheet to seal the inner package in the outer package; sterilizing the product; and
 - sealing the first sheet to the second sheet to seal the inner package in a gas impermeable outer package in the form of a pouch.
13. The method of claim 14, further including, after the sterilizing step, removing the atmosphere inside the outer package.

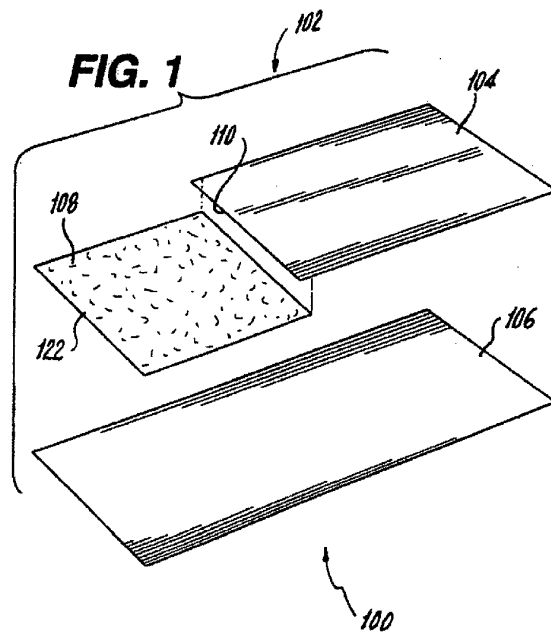
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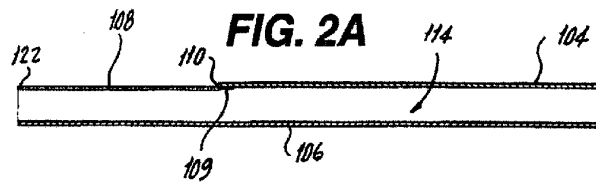
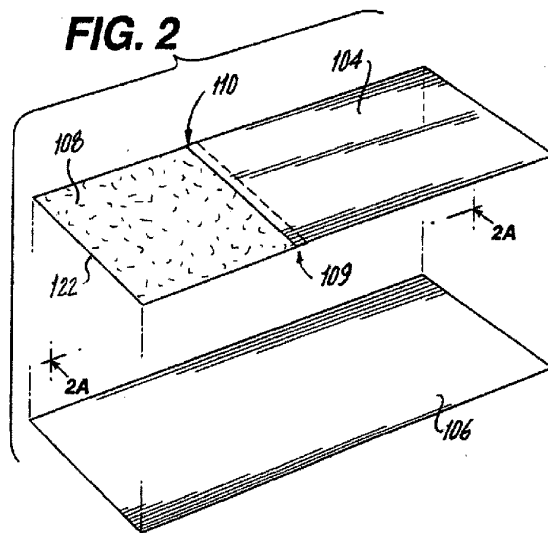
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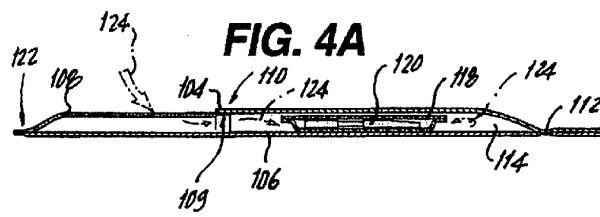
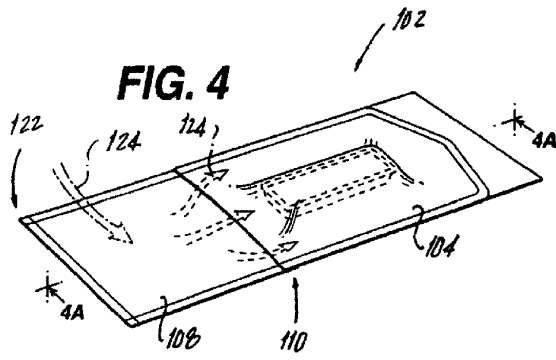
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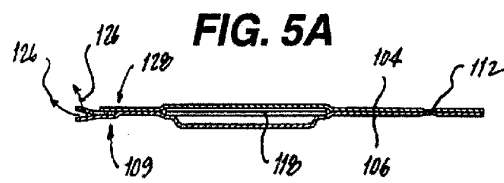
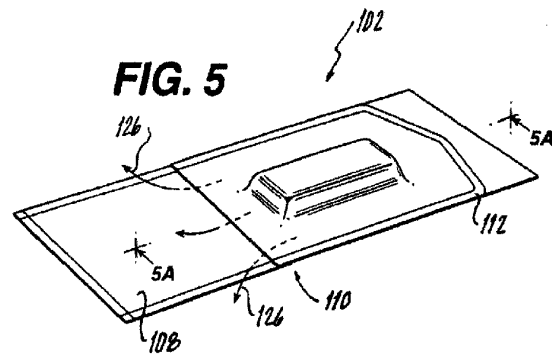
14. The method of claim 13, further including, prior to the removing step, filling the outer package with an inert gas.
15. The method of claim 13 or claim 14, further including, after the removing step, filling the outer package with an inert gas.
- 5 16. The method of any one of claims 12 to 15, further including removing the gas permeable header.
17. The method of any one of claims 12 to 16, wherein the sterilizing step includes at least one of steam sterilization, EtO sterilization, gas plasma/radio frequency-peroxide sterilization, chemical vapor sterilization, and cold sterilization.
- 10 18. The method of any one of claims 12 to 17, wherein the product is incorporated with at least one of an antimicrobial agent, an antiangiogenesis agent, an antiproliferative agent and an anti-inflammatory.
- 15 19. A package, substantially as hereinbefore described with reference to the drawings.
20. A method of packaging a drug-device combination product, substantially as hereinbefore described with reference to the drawings.

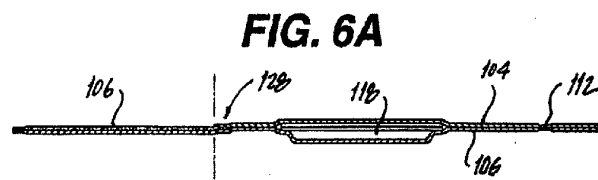
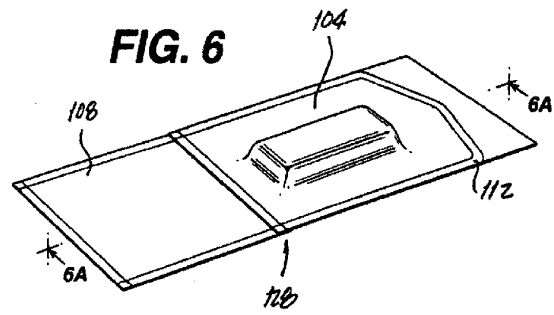
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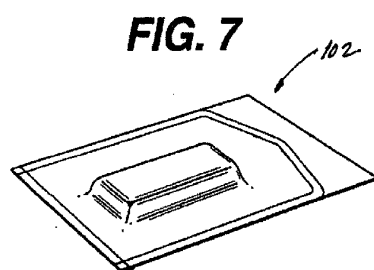


FIG. 8

