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(54) Title: METHODS OF TREATING CANCER

(57) **Abrégé/Abstract:**

The disclosure pertains to drug delivery conjugates for targeted therapy. The disclosure relates to methods of treating PSMA expressing cancers with a combination of compounds of the formulas I-Lu or Ia-Lu, and I-Ac or Ia-Ac. The disclosure also relates to methods of treating PSMA-expressing cancers with a combination of compounds of the formulas I-Lu or Ia-Lu, and I-Ac or Ia-Ac in patients where stable disease results after treatment with a combination of compounds of the formulas I-Lu or Ia-Lu, and I-Ac or Ia-Ac.

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(54) Title: METHODS OF TREATING CANCER

(57) Abstract: The disclosure pertains to drug delivery conjugates for targeted therapy. The disclosure relates to methods of treating PSMA expressing cancers with a combination of compounds of the formulas I-Lu or Ia-Lu, and I-Ac or Ia-Ac. The disclosure also relates to methods of treating PSMA-expressing cancers with a combination of compounds of the formulas I-Lu or Ia-Lu, and I-Ac or Ia-Ac in patients where stable disease results after treatment with a combination of compounds of the formulas I-Lu or Ia-Lu, and I-Ac or Ia-Ac.

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METHODS OF TREATING CANCER

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims priority under 35 U.S.C. § 119(e) to U. S. Provisional Application Serial No. 62/734,649 filed on September 21, 2018, the entire disclosure of which
5 is incorporated herein by reference.

TECHNICAL FIELD

The present disclosure relates to drug delivery conjugates for targeted therapy. The present disclosure relates to methods of treating PSMA expressing cancers with a combination
10 of compounds of the formulas I-Lu or Ia-Lu and I-Ac or Ia-Ac, wherein ^{177}Lu or ^{225}Ac are complexed to compounds I and Ia. The present disclosure also relates to methods of treating PSMA-expressing cancers with a combination of compounds of the formulas I-Lu or Ia-Lu and I-Ac or Ia-Ac.

15 BACKGROUND

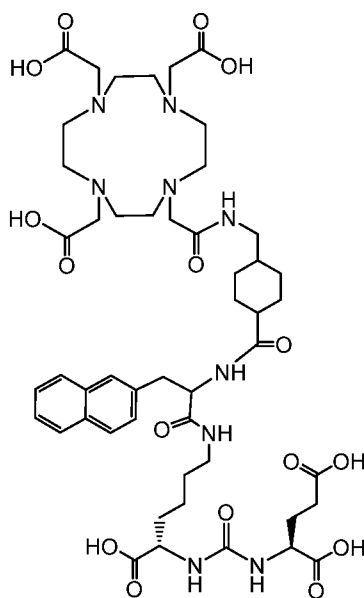
Prostate specific membrane antigen (PSMA) is a type II cell surface membrane-bound glycoprotein with ~110 kD molecular weight, including an intracellular segment (amino acids 1-18), a transmembrane domain (amino acids 19-43), and an extensive extracellular domain (amino acids 44-750). While the functions of the intracellular segment and the transmembrane
20 domains are currently believed to be insignificant, the extracellular domain is involved in several distinct activities. PSMA plays a role in the central nervous system, where it metabolizes N-acetyl-aspartyl glutamate (NAAG) into glutamic and N-acetyl aspartic acid. Accordingly, it is also sometimes referred to as an N-acetyl alpha linked acidic dipeptidase (NAALADase). PSMA is also sometimes referred to as a folate hydrolase I (FOLH I) or
25 glutamate carboxypeptidase (GCP II) due to its role in the proximal small intestine where it removes γ -linked glutamate from poly- γ -glutamated folate and α -linked glutamate from peptides and small molecules.

PSMA is named largely due to its higher level of expression on prostate cancer cells; however, its particular function on prostate cancer cells remains unresolved. PSMA expression
30 is highly restricted in man, present in only salivary gland tissue, renal tissue small numbers of cells in the small and large intestine. PSMA is over-expressed in the malignant prostate tissues when compared to other organs in the human body such as kidney, proximal small intestine, and salivary glands. Higher PSMA expression is associated with high grade, metastatic and

castration resistance disease. Tumor expression in prostate cancer is typically 100 to 1,000-fold higher. Unlike many other membrane-bound proteins, PSMA undergoes rapid internalization into the cell in a similar fashion to cell surface bound receptors like vitamin receptors. PSMA is internalized through clathrin-coated pits and subsequently can either recycle to the cell surface or go to lysosomes. It has been suggested that the dimer and monomer form of PSMA are inter-convertible, though direct evidence of the interconversion is being debated. Even so, only the dimer of PSMA possesses enzymatic activity, and the monomer does not.

PSMA is also expressed on the neovasculature of other tumors, such as thyroid cancer, renal clear cell carcinoma, transitional cell carcinoma of bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, and soft tissue sarcoma, breast carcinoma. These cancers represent a large range of different tumors with different histological subtypes, growth rates and cell cycle times. In some cases, the cancers are imbedded within normal tissues having variable radiation tolerances. In addition, hypoxic areas of larger deposits may also lead to radio resistance. These and other factors are known to result in different intrinsic response to traditional external beam radiation therapy.

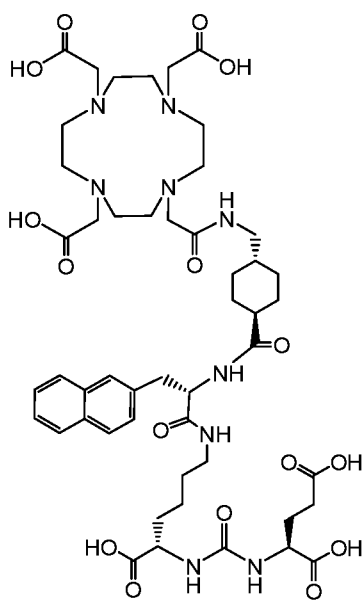
Though the activity of the PSMA on the cell surface of the prostate cells remains under investigation, it has been recognized by the inventors herein that PSMA represents a viable target for the selective and/or specific delivery of biologically active agents or combinations of biologically active agents, including drug compounds to such prostate cells. One such drug compound is the compound of Formula I



I

wherein ^{177}Lu is complexed to the compound to provide I-Lu, or ^{225}Ac is complexed to compound I to provide I-Ac, useful for the treatment of cancer as described in WO2015/055318. Compounds I-Lu and I-Ac can be prepared according to the methods described in WO2015/055318, incorporated by reference for the preparation of Compounds I-Lu and I-Ac, as described in Example 3 and Example 5.

Another such drug compound is Compound Ia



Ia

(a.k.a. (3S,10S,14S)-3-[(naphthalen-2-yl)methyl]-1,4,12-trioxo-1-[(1R,4S)-4-[[2-[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododecan-1-yl]acetamido]methyl]cyclohexyl]-2,5,11,13-tetraazahexadecane-10,14,16-tricarboxylic acid) wherein ^{177}Lu is complexed to compound Ia to provide Ia-Lu, or ^{225}Ac is complexed to the compound to provide Ia-Ac, useful for the treatment of cancer as described in WO2015/055318. Compounds Ia-Lu and Ia-Ac can be prepared according to the methods described in WO2015/055318, incorporated by reference for the preparation of Compounds Ia-Lu and Ia-Ac, as described in Example 3 and Example 5.

Compound I or Ia can be described as a small molecule that specifically binds to PSMA (prostatic specific membrane antigen) which is expressed on the surface of prostate cancer cells. Compound I or Ia can be characterized as composed of a pharmacophore ligand, glutamate-urea-lysine; a chelator, DOTA (able to complex ^{177}Lu and ^{225}Ac); and a linker connecting the ligand and the chelator. Without being bound by theory, it is believed that the urea-based pharmacophore ligand allows the agent to bind to, and be internalized by PSMA at the site of disease. It is further believed that the binding of I-Lu, I-Ac, Ia-Lu, or Ia-Ac can lead to

internalization through endocytosis which can provide a sustained retention of the ligand and its bound radioactive cargo within the cancer cell.

Previous radioligand therapy (RLT) used in the clinic includes ^{131}I in thyroid cancer, and elements emitting alpha radiation, such as $^{223}\text{Radium}$ or $^{89}\text{Strontium}$, for the treatment of
5 bone metastases.

^{177}Lu has a half-life of 6.7 days. It emits a combination of 0.5MeV energy consisting of negatively charged Beta particles (electrons) that travel chaotically through tissues for approximately 20-80 cells or 0.5-2mm and cause predominantly base damage and single strand breaks. At high dose these lesions can interact to convert sublethal damage (SLD) or
10 potentially lethal damage (PLD) to irreparable, lethal damage. ^{177}Lu also emits 113Kv and 208kV radiation which can be used for imaging.

^{225}Ac has a half-life of 9.9 days, and in contrast emits 8.38MV energy alpha particles. Only 0.5% of energy is emitted as 142Kv photon emissions. The majority of radiation particles are therefore positively charged, and about 8,000 times larger than β particles. Furthermore, the
15 energy from these particles is deposited over relatively short distances (2-3 cells). As a result, there is dense and severe tissue damage in the form of double strand breaks with multiply damaged sites that represent irreparable lethal damage. This is called High Linear Energy Transfer (LET) or densely ionizing ionization and it delivers 3-7 x more absorbed dose than β .

The type of cellular damage inflicted by either isotope (^{177}Lu or ^{225}Ac) is expected to be
20 different due to the difference of the characteristics of each warhead. ^{177}Lu is believed to provide a longer path length of radiation and therefore can be effective in delivering radiation to adjacent cells. The preponderance of single strand breaks, especially in the presence of oxygen, provides the opportunity to repair sub lethal damage (SLD) and or potentially lethal damage (PLD) providing the optimal conditions for normal tissue repair. On the contrary,
25 ^{225}Ac delivers extremely powerful, high LET radiation, and the potential for repair of normal tissue is much more limited. The radiological biological effectiveness of alpha radiation is at least 5 times that of beta irradiation and administered doses the relative biological effectiveness (RBE) has to be taken into account. With ^{225}Ac therapy, the type of DNA damage inflicted
30 does not require the presence of oxygen so it will also be more effective in hypoxic tumor regions. A possible disadvantage of ^{225}Ac therapy is that the short path length can lead to large amounts of damaging radiation deposited only within a short distance of 2-4 cells.

Another such compound is the PSMA-imaging conjugate 4

In some embodiments, the present disclosure provides use of Compound I-Lu or Ia-Lu, in the preparation of a medicament useful for the treatment of a cancer in a patient in combination with Compound I-Ac or Ia-Ac. In some aspects, the medicament comprises a therapeutically effective amount of Compound I-Lu or I-Lu.

5 In some embodiments, the present disclosure provides use of Compound I-Ac or Ia-Ac, in the preparation of a medicament useful for the treatment of a cancer in a patient in combination with Compound I-Lu or Ia-Lu. In some aspects, the medicament comprises a therapeutically effective amount of Compound I-Ac or I-Ac.

10 In some embodiments, the present disclosure provides use of Compound I-Lu or Ia-Lu, in the preparation of a first medicament useful for the treatment of a cancer in a patient in combination with a second medicament comprising Compound I-Ac or Ia-Ac. In some aspects, the first medicament comprises a therapeutically effective amount of Compound I-Lu or I-Lu, and the second medicament comprises a therapeutically effective amount of Compound I-Ac or I-Ac.

15 In some aspects of these embodiments, the cancer is a PSMA expressing cancer. In some aspects of these embodiments, the compound is at least about 98 percent pure. In some embodiments, the cancer is selected from the group consisting of a glioma, a carcinoma, a sarcoma, a lymphoma, a melanoma, a mesothelioma, a nasopharyngeal carcinoma, a leukemia, an adenocarcinoma, and a myeloma.

20 In some aspects of these embodiments, the cancer is selected from the group consisting of lung cancer, bone cancer, pancreatic cancer, skin cancer, cancer of the head, cancer of the neck, cutaneous melanoma, intraocular melanoma uterine cancer, ovarian cancer, endometrial cancer, rectal cancer, stomach cancer, colon cancer, breast carcinoma, triple negative breast cancer, metastatic breast cancer, carcinoma of the fallopian tubes, carcinoma of the
25 endometrium, carcinoma of the cervix, carcinoma of the vagina, carcinoma of the vulva, Hodgkin's Disease, cancer of the esophagus, cancer of the small intestine, cancer of the endocrine system, cancer of the thyroid gland, cancer of the parathyroid gland, non-small cell lung carcinoma, cancer of the adrenal gland, soft tissue sarcoma, cancer of the urethra, cancer of the penis, prostate cancer, metastatic castration-resistant prostate cancer (mCRPC), thyroid
30 cancer, transitional cell carcinoma of the bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, chronic leukemia, acute leukemia, lymphocytic lymphomas, pleural mesothelioma, cancer of the bladder, Burkitt's lymphoma, cancer of the ureter, cancer of the kidney, renal cell carcinoma, carcinoma of the renal pelvis, neoplasms of the central nervous system (CNS), primary CNS

lymphoma, spinal axis tumors, glioma, brain stem glioma, pituitary adenoma, and adenocarcinoma of the gastroesophageal junction. In some aspects of these embodiments, the cancer is a primary or secondary brain cancer. In some aspects of these embodiments, the cancer is prostate cancer. In some aspects of these embodiments, the cancer is metastatic prostate cancer.

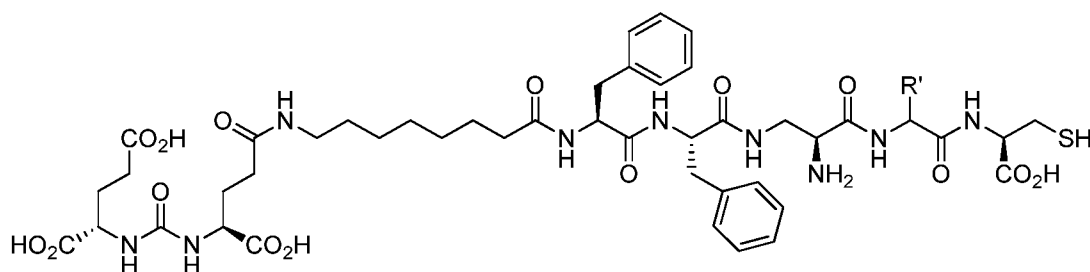
In some aspects of these embodiments, a combination of Compounds I-Lu or Ia-Lu and I-Ac or Ia-Ac is administered in a parenteral dosage form. In some aspects of these embodiments, the parenteral dosage form is selected from the group consisting of intradermal, subcutaneous, intramuscular, intraperitoneal, intravenous, and intrathecal. In some aspects of these embodiments, the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 13 GBq. In some aspects of these embodiments, the therapeutically effective amount of I-Lu or Ia-Lu is from about 4 GBq to about 11 GBq. In some aspects of these embodiments, the therapeutically effective amount of I-Lu or Ia-Lu is from about 5 GBq to about 10 GBq. In some aspects of these embodiments, the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 9 GBq. In some aspects of these embodiments, the therapeutically effective amount of I-Lu or Ia-Lu is from about 6.5 GBq to about 8.5 GBq. In some aspects of these embodiments, the therapeutically effective amount of I-Lu or Ia-Lu is from about 7 GBq to about 8 GBq. In some aspects of these embodiments, the therapeutically effective amount of I-Lu or Ia-Lu is about 7.4 GBq. In some aspects of these embodiments, the total dose of I-Lu or Ia-Lu ranges from about 15 GBq to about 200 GBq. In some aspects of these embodiments, the total dose of I-Lu or Ia-Lu ranges from about 25 GBq to about 185 GBq. In some aspects of these embodiments, the total dose of I-Lu or Ia-Lu ranges from about 35 GBq to about 150 GBq. In some aspects of these embodiments, the total dose of I-Lu or Ia-Lu ranges from about 40 GBq to about 100 GBq. In some aspects of these embodiments, the total dose of I-Lu, or Ia-Lu is about 44 GBq. In some aspects of these embodiments, the maximum duration of treatment of a subject is about 19 to 23 months.

In some aspects of these embodiments, the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 20 MBq. In some aspects of these embodiments, the therapeutically effective amount of I-Ac or Ia-Ac is from about 4 MBq to about 14 MBq. In some aspects of these embodiments, the therapeutically effective amount of I-Ac or Ia-Ac is from about 5 MBq to about 10 MBq. In some aspects of these embodiments, the therapeutically effective amount of I-Ac or Ia-Ac is from about 6 MBq to about 8 MBq. In some aspects of these embodiments, the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 4 MBq. In some aspects of these embodiments, the therapeutically effective amount of

I-Ac or Ia-Ac is from about 2 MBq to about 3 MBq. In some aspects of these embodiments, the therapeutically effective amount of I-Ac or Ia-Ac is about 2.5 MBq.

In other aspects, the methods and uses described herein further comprise imaging PSMA expression by the cancer. In some aspects of these embodiments, the step of imaging occurs before the step of administering. In some aspects of these embodiments, the step of imaging occurs after the step of administering. In some aspects of these embodiments, the imaging is performed by imaging wherein the imaging is selected from the group consisting of SPECT imaging, PET imaging, IHC, and FISH. In some aspects of these embodiments, the imaging is performed by SPECT imaging.

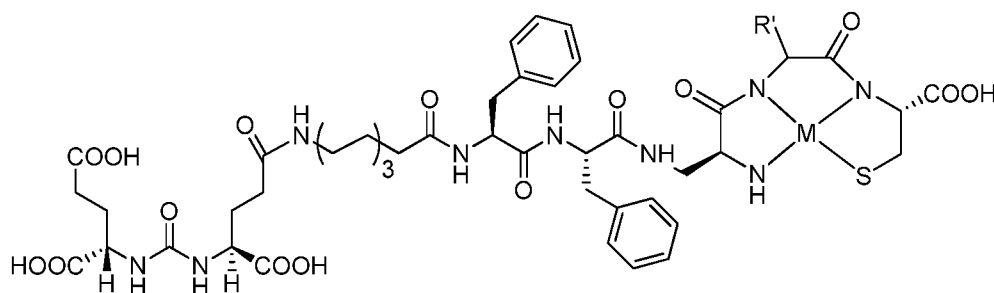
In some aspects of these embodiments, the step of imaging comprises administering to the patient a PSMA ligand-imaging conjugate of the formula 2



2

or a pharmaceutically acceptable salt thereof, wherein R' is hydrogen, or R' is selected from the group consisting of alkyl, aminoalkyl, carboxyalkyl, hydroxyalkyl, heteroalkyl, aryl, arylalkyl and heteroarylalkyl, each of which is optionally substituted, and wherein a radionuclide is bound to the conjugate.

In some aspects of these embodiments, the step of imaging comprises administering a PSMA ligand-imaging conjugate of the formula 3

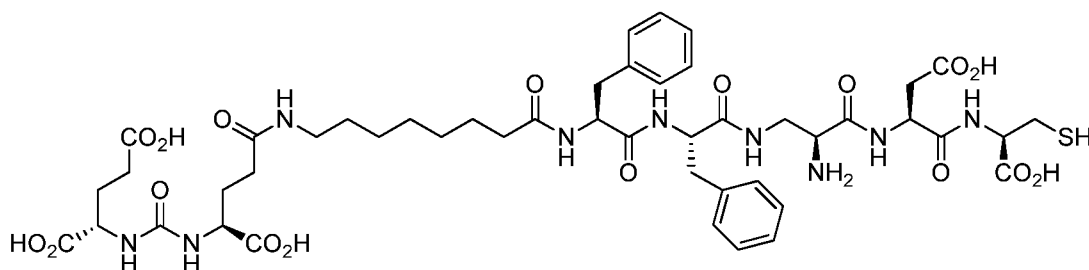


3

or a pharmaceutically acceptable salt thereof, wherein R' is hydrogen, or R' is selected from the group consisting of alkyl, aminoalkyl, carboxyalkyl, hydroxyalkyl, heteroalkyl, aryl, arylalkyl and heteroarylalkyl, each of which is optionally substituted, and wherein M is a cation of a

radionuclide. In some aspects of these embodiments, M in the conjugate, or a pharmaceutically acceptable salt thereof, is selected from the group consisting of an isotope of gallium, an isotope of indium, an isotope of copper, an isotope of technetium, and an isotope of rhenium. In some aspects of these embodiments, M in the conjugate, or a pharmaceutically acceptable salt thereof, is an isotope of technetium.

In some aspects of these embodiments, the PSMA ligand-imaging conjugate is of the formula 2a

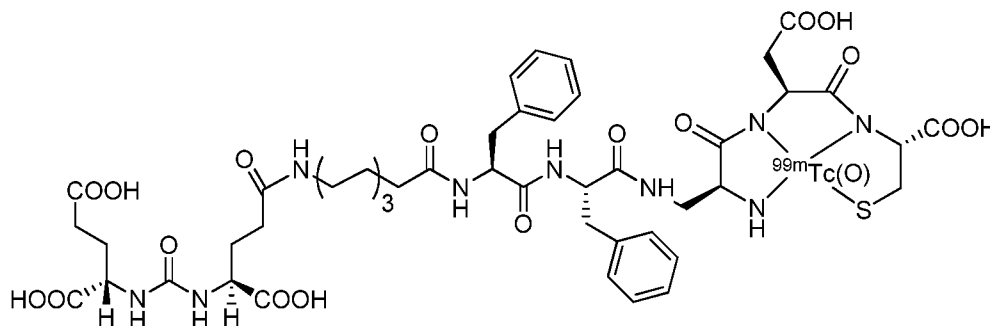


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2a

or a pharmaceutically acceptable salt thereof, wherein a radionuclide is bound to the conjugate.

In some aspects of these embodiments, the PSMA ligand-imaging conjugate is of the formula 3a



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3a

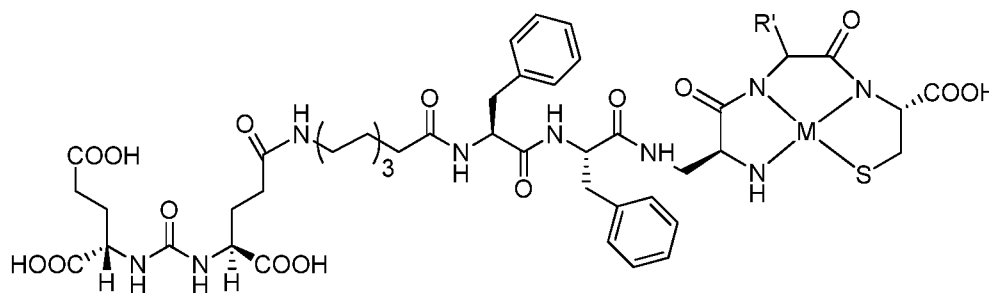
or a pharmaceutically acceptable salt thereof.

In some aspects of these embodiments, the step of imaging comprises administering to the patient a PSMA ligand-imaging conjugate of the formula 4

2

or a pharmaceutically acceptable salt thereof, wherein R' is hydrogen, or R' is selected from the group consisting of alkyl, aminoalkyl, carboxyalkyl, hydroxyalkyl, heteroalkyl, aryl, arylalkyl and heteroarylalkyl, each of which is optionally substituted, and wherein the conjugate is bound to a radionuclide.

In some aspects of these embodiments, the step of determining comprises administering a PSMA ligand-imaging conjugate of the formula 3

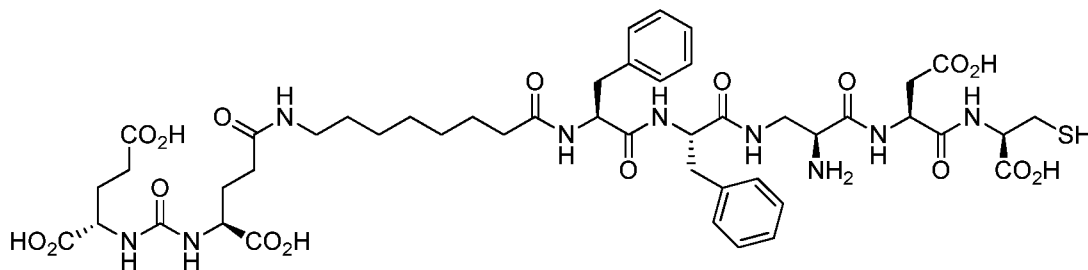


3

or a pharmaceutically acceptable salt thereof, wherein R' is hydrogen, or R' is selected from the group consisting of alkyl, aminoalkyl, carboxyalkyl, hydroxyalkyl, heteroalkyl, aryl, arylalkyl and heteroarylalkyl, each of which is optionally substituted, and wherein M is a cation of a radionuclide.

In some aspects of these embodiments, M in the conjugate, or a pharmaceutically acceptable salt thereof, is selected from the group consisting of an isotope of gallium, an isotope of indium, an isotope of copper, an isotope of technetium, and an isotope of rhenium. In some aspects of these embodiments, M in the conjugate, or a pharmaceutically acceptable salt thereof, is an isotope of technetium. In some aspects of these embodiments, the PSMA ligand-imaging conjugate is of the formula 2a

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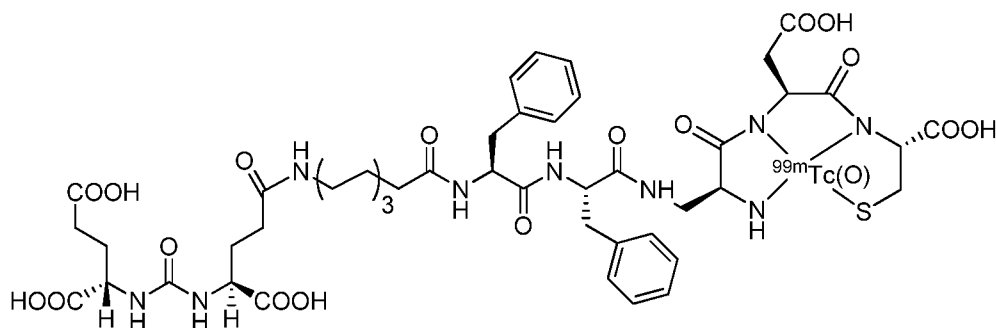
2a

or a pharmaceutically acceptable salt thereof, wherein a radionuclide is bound to the conjugate.

In some aspects of these embodiments, the PSMA ligand-imaging conjugate is of the formula 3a

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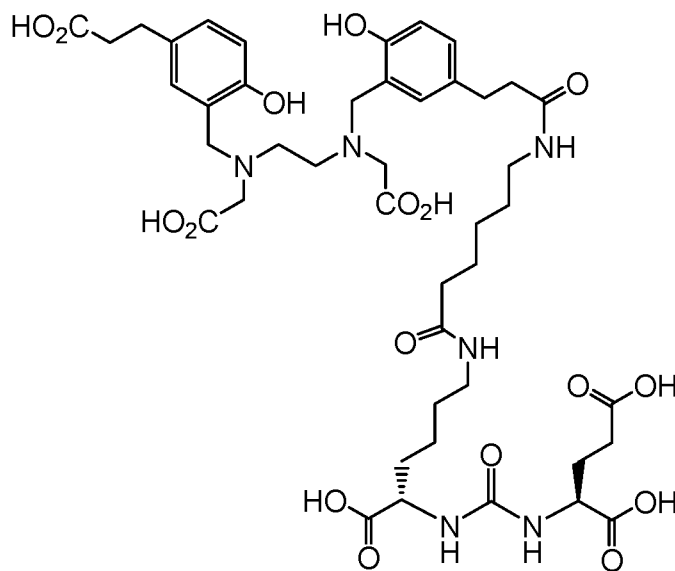
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3a

or a pharmaceutically acceptable salt thereof.

In some aspects of these embodiments, the step of determining comprises administering
5 to the patient a PSMA ligand-imaging conjugate of the formula 4



4

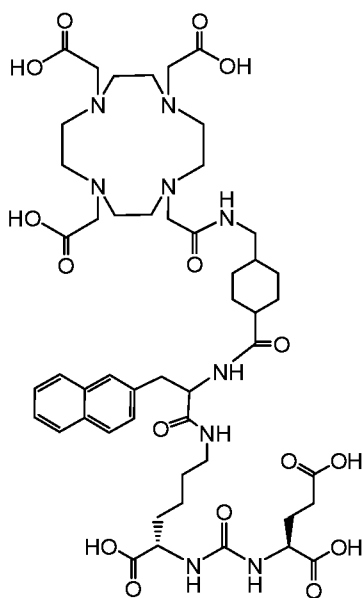
or a pharmaceutically acceptable salt thereof, wherein a radionuclide is bound to the conjugate.

In some aspects of these embodiments, the radionuclide is ^{68}Ga .

10 In some aspects of these embodiments, the step of determining comprises detecting the compound of the formula I-Lu or Ia-Lu administered for the purpose of treating.

In other embodiments, the present disclosure provides a method of treating a cancer in a patient in need of such treatment comprising, administering to the patient a therapeutically effective combination of Compounds I-Lu and I-Ac

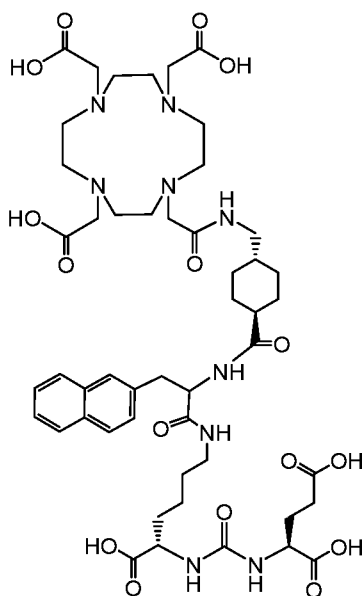
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I

wherein ^{177}Lu is complexed to the Compound in I-Lu, and ^{225}Ac is complexed to the
 Compound in I-Ac, wherein stable disease results after a combination of Compounds I-Lu and
 5 I-Ac.

In other embodiments, the present disclosure provides a method of treating a cancer in a
 patient in need of such treatment comprising, administering to the patient a therapeutically
 effective combination of Compounds Ia-Lu and Ia-Ac



Ia

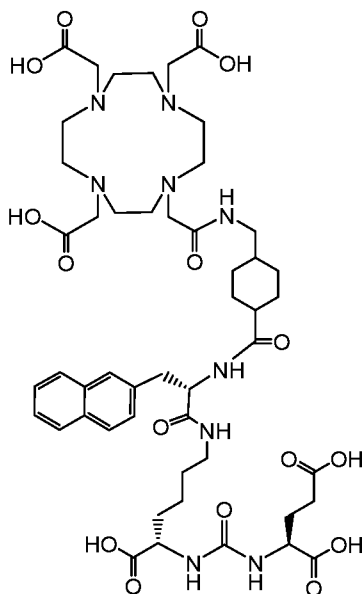
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wherein ^{177}Lu is complexed to the Compound in Ia-Lu, and ^{225}Ac is complexed to Compound I
 in Ia-Ac, wherein stable disease results after a combination of Compounds Ia-Lu and Ia-Ac.

wherein ^{177}Lu is complexed to the Compound in Ia-Lu, and ^{225}Ac is complexed to Compound Ia in Ia-Ac, wherein stable disease results after a combination of Compounds Ia-Lu and Ia-Ac is administered. In some aspects of these embodiments, the use comprises administering to the patient a therapeutically effective combination of Compounds Ia-Lu and

5 Ia-Ac.

In other embodiments, the present disclosure provides use of Compounds I-Lu and I-Ac



I

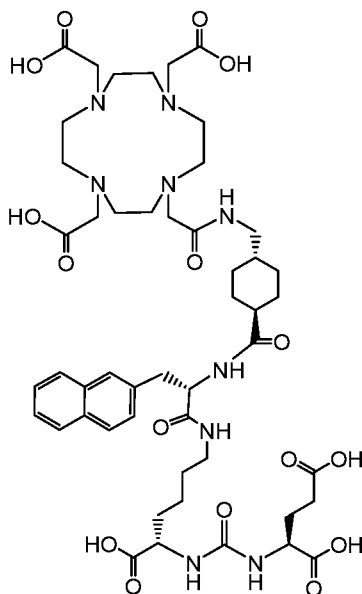
10 wherein ^{177}Lu is complexed to the Compound I in I-Lu, and ^{225}Ac is complexed to the Compound I in I-Ac, in the preparation of a medicament useful for the treatment of a cancer in a patient. In some aspects, the medicament comprises a therapeutically effective combination of the Compounds I-Lu and I-Ac.

In other embodiments, the present disclosure provides use of Compounds I-Lu and I-Ac

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wherein ^{177}Lu is complexed to the Compound Ia in Ia-Lu, and ^{225}Ac is complexed to the Compound Ia in Ia-Ac, in the preparation of a medicament useful for the treatment of a cancer in a patient. In some aspects, the medicament comprises a therapeutically effective combination of the Compounds Ia-Lu and Ia-Ac.

5 In other embodiments, the present disclosure provides use of Compounds Ia-Lu and Ia-Ac



Ia

10 wherein ^{177}Lu is complexed to the Compound Ia in Ia-Lu, and ^{225}Ac is complexed to the Compound Ia in Ia-Ac, in the preparation of a medicament useful for the treatment of a cancer in a patient, wherein stable disease results after a combination of Compounds Ia-Lu and Ia-Ac are administered. In some aspects, the medicament comprises a therapeutically effective combination of the Compounds Ia-Lu and Ia-Ac.

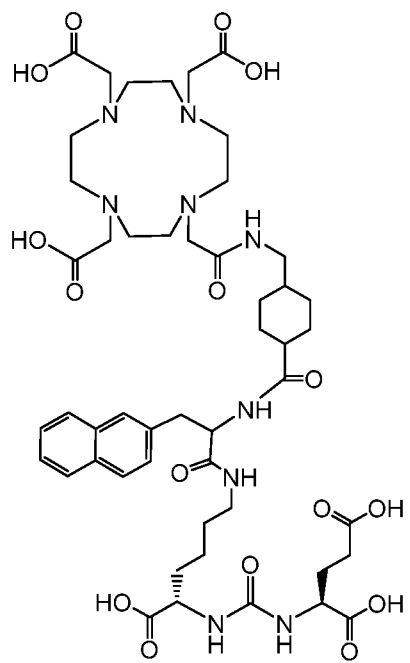
15 In some aspects of these embodiments, the patient has been treated with at least one prior treatment. In some aspects of these embodiments, the at least one prior treatment is selected from the group consisting of an androgen axis systemic treatment, a chemotherapeutic agent, surgery, radiation therapy, immunotherapy, photodynamic therapy, stem cell therapy, and hyperthermia. In some aspects of these embodiments, the at least one prior treatment is a
 20 systemic treatment. In some aspects of these embodiments, the systemic treatment is selected from the group consisting of palifosfamide, 5-fluorouracil, capecitabine, pemetrexed, cisplatin, carboplatin, gemcitabine, paclitaxel, vinorelbine, eribulin, docetaxel, cyclophosphamide, doxorubicin, regorafenib, and combinations thereof. In some aspects of these embodiments, the

cancer is a PSMA expressing cancer. In some aspects of these embodiments, the compound is at least about 98 percent pure.

In some aspects of these embodiments, I-Lu or Ia-Lu is administered prior to I-Ac or Ia-Ac. In some aspects of these embodiments, I-Lu or Ia-Lu is administered prior to I-Ac or Ia-Ac on the same day. In some aspects of these embodiments, I-Lu or Ia-Lu is administered at the same time as I-Ac or Ia-Ac. In some aspects of these embodiments, I-Ac or Ia-Ac is administered prior to I-Lu or Ia-Lu. In some aspects of these embodiments, I-Ac or Ia-Ac is administered prior to I-Lu or Ia-Lu on the same day.

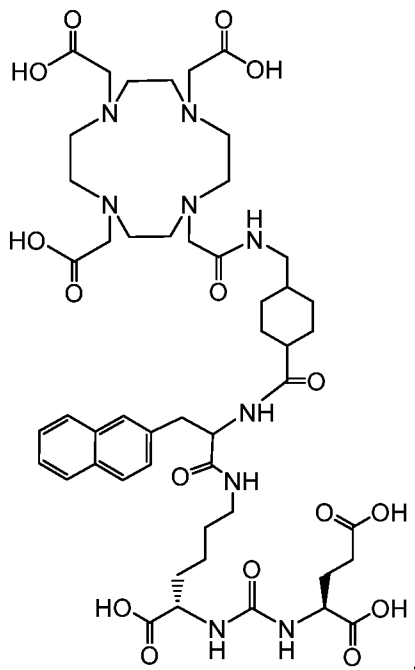
Certain embodiments are further described by the following enumerated clauses:

- 10 1. A method for treating cancer in a host animal, the method comprising the step of administering to the host animal a therapeutically effective amount of a first compound having the Formula I



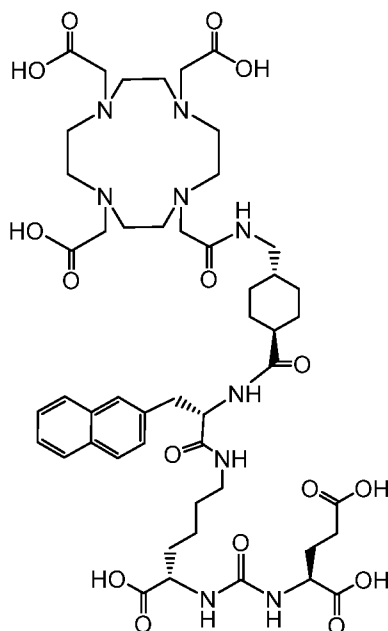
wherein the compound is complexed with ^{177}Lu (I-Lu);

- 15 in combination with a therapeutically effective amount of a second compound having the Formula I



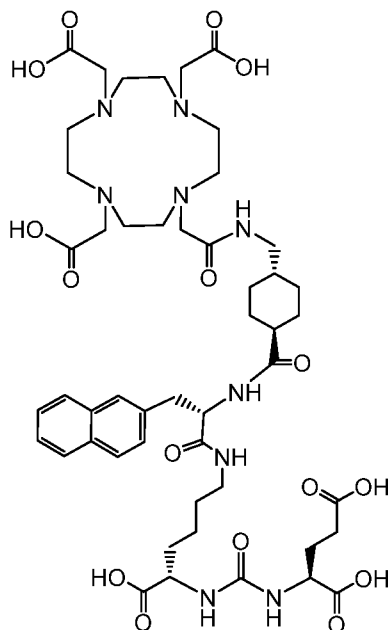
wherein the compound is complexed with ^{225}Ac (I-Ac).

2. The method of clause 1, wherein the first compound is of the Formula Ia



- 5 wherein the compound is complexed with ^{177}Lu .

3. The method of clause 1 or 2, wherein the second compound is of the Formula Ia



wherein the compound is complexed with ^{225}Ac .

4. The method of clauses 1 to 3, wherein the cancer is associated with expression of prostate specific membrane antigen (PSMA).
5. The method of any one of the preceding clauses, wherein the cancer is selected from the group consisting of prostate cancer, metastatic castration-resistant prostate cancer (mCRPC), thyroid cancer, renal cell carcinoma, transitional cell carcinoma of the bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, soft tissue sarcoma, and breast carcinoma.
6. The method of any one of the preceding clauses, wherein the cancer is prostate cancer.
7. The method of any one of the preceding clauses, wherein the cancer is metastatic castration-resistant prostate cancer (mCRPC).
8. The method of any one of the preceding clauses, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq.
9. The method of any one of the preceding clauses, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq.
10. The method of any one of the preceding clauses, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 7.4 GBq.
11. The method of any one of the preceding clauses, wherein the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq; or about 5 MBq to about 10 MBq; or about 5 MBq to about 7MBq.

12. The method of any one of the preceding clauses, wherein the therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq.

13. The method of any of the preceding clauses, wherein the compound of Formula I-Lu or Ia-Lu is administered at the same time as the compound of Formula I-Ac or Ia-Ac.
5

14. The method of any one of clauses 1-12, wherein the compound of Formula I-Lu or Ia-Lu is administered about 1 hour prior to the compound of Formula I-Ac or Ia-Ac; or about 12 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 24 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 48 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 72 hours prior to the compound of Formula I-Ac or Ia-Ac.
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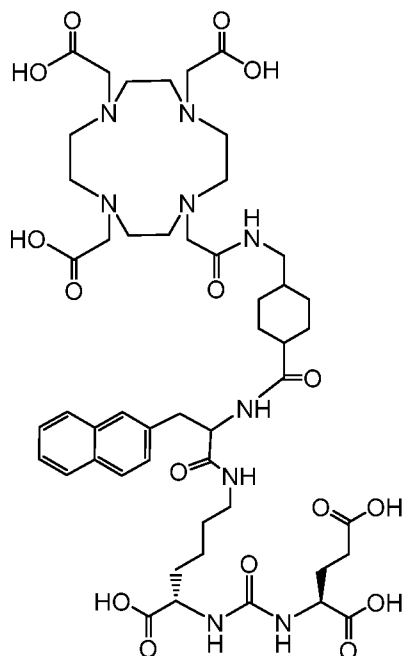
15. The method of any one of clauses 1-12, wherein the compound of Formula I-Ac or Ia-Ac is administered about 1 hour prior to the compound of Formula I-Lu or Ia-Lu; or about 12 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 24 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 48 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 72 hours prior to the compound of Formula I-Lu or Ia-Lu.
15

16. The method of clause 13, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of I-Lu or Ia-Lu and I-Ac or Ia-Ac.
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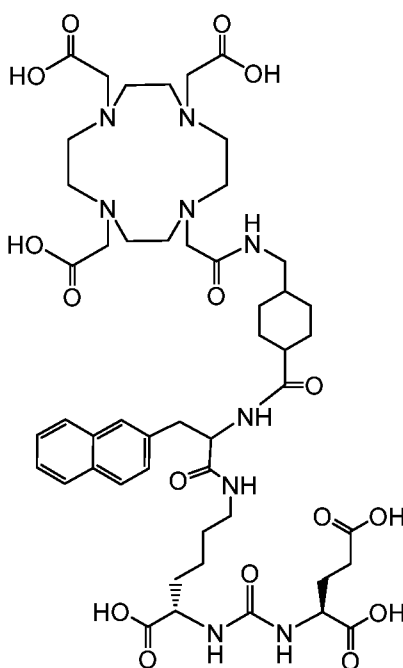
17. The method of clause 14, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

18. The method of clause 15, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.
25

19. A compound of the Formula I-Lu



wherein the compound is complexed with ^{177}Lu , for use in the treatment of cancer in a patient, in combination with a therapeutically effective amount of a compound of the Formula I-Ac



5 wherein the compound is complexed with ^{225}Ac .

20. The compound of clause 19, wherein the compound is of the Formula Ia-Lu

cancer (mCRPC), thyroid cancer, renal cell carcinoma, transitional cell carcinoma of the bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, soft tissue sarcoma, and breast carcinoma.

5 24. The compound of any one of clauses 19 to 23, wherein the cancer is prostate cancer.

 25. The compound of any one of clauses 19 to 24, wherein the cancer is metastatic castration-resistant prostate cancer (mCRPC).

10 26. The compound of any one of clauses 19 to 25, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq.

 27. The compound of any one of clauses 19 to 26, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq.

 28. The compound of any one of clauses 19 to 27, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 7.4 GBq.

15 29. The compound of any one of clauses 19 to 28, wherein the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq; or about 5 MBq to about 10 MBq; or about 5 MBq to about 7MBq.

 30. The compound of any one of clauses 19 to 29, wherein the therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq.

20 31. The compound of any one of clauses 19 to 30, wherein the compound of Formula I-Lu or Ia-Lu is administered at the same time as the compound of Formula I-Ac or Ia-Ac.

 32. The compound of any one of clauses 19 to 30, wherein the compound of Formula I-Lu or Ia-Lu is administered about 1 hour prior to the compound of Formula I-Ac or Ia-Ac; or about 12 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 24 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 48 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 72 hours prior to the compound of Formula I-Ac or Ia-Ac.

 33. The compound of any one of clauses 19 to 30, wherein the compound of Formula I-Ac or Ia-Ac is administered about 1 hour prior to the compound of Formula I-Lu or Ia-Lu; or about 12 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 24 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 48 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 72 hours prior to the compound of Formula I-Lu or Ia-Lu.

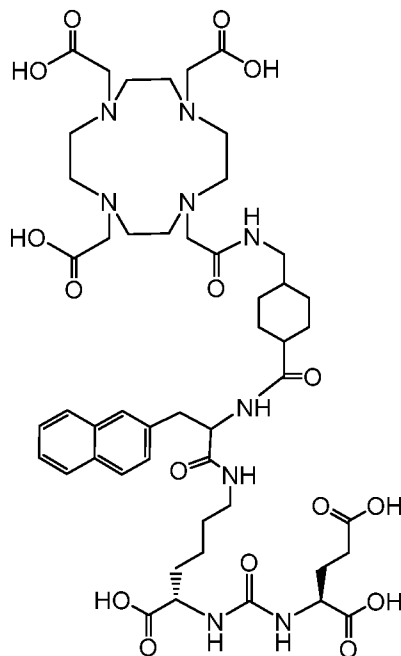
 34. The compound of clause 31, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly

cycle for from about 1 to about 7 cycles following the administration of I-Lu or Ia-Lu and I-Ac or Ia-Ac.

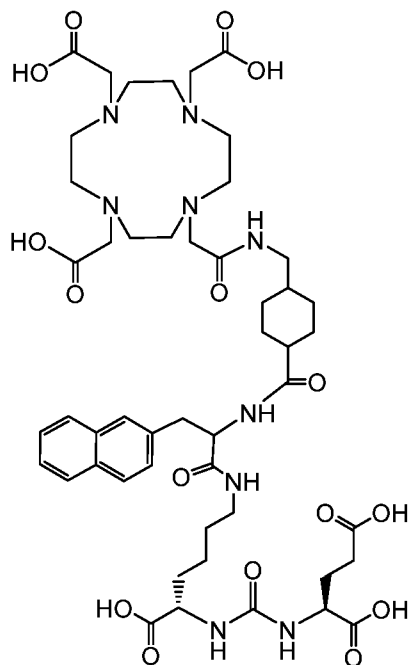
35. The compound of clause 32, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

36. The compound of clause 33, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

37. Use of a compound of the Formula I-Lu,

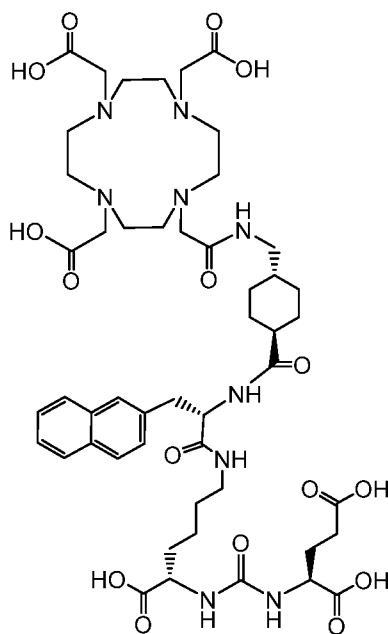


wherein the compound is complexed with ^{177}Lu , in the preparation of a medicament comprising a therapeutically effective amount of the compound of the Formula I-Lu, for treating cancer in a patient in combination with a therapeutically effective amount of a compound of the Formula I-Ac



wherein the compound is complexed with ^{225}Ac .

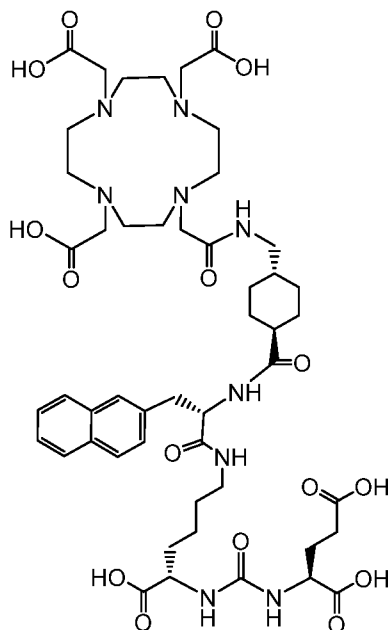
38. The use of clause 37, wherein the compound of the Formula I-Lu is of the Formula Ia-Lu



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wherein the compound is complexed with ^{177}Lu .

39. The use of clause 37, wherein the compound of the Formula I-Ac is of the Formula Ia-Ac



wherein the compound is complexed with ^{225}Ac .

40. The use of any one of clauses 37 to 39, wherein the cancer is associated with expression of prostate specific membrane antigen (PSMA).

5 41. The use of any one of clauses 37 to 40, wherein the cancer is selected from the group consisting of prostate cancer, metastatic castration-resistant prostate cancer (mCRPC), thyroid cancer, renal cell carcinoma, transitional cell carcinoma of the bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, soft tissue sarcoma, and
10 breast carcinoma.

42. The use of any one of clauses 37 to 41, wherein the cancer is prostate cancer.

43. The use of any one of clauses 37 to 42, wherein the cancer is metastatic castration-resistant prostate cancer (mCRPC).

15 44. The use of any one of clauses 37 to 43, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq.

45. The use of any one of clauses 37 to 44, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq.

20 46. The use of any one of clauses 37 to 45, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 7.4 GBq.

47. The use of any one of clauses 37 to 46, wherein the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq; or about 5 MBq to about 10 MBq; or about 5 MBq to about 7MBq.

48. The use of any one of clauses 37 to 47, wherein the therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq.

49. The use of any one of clauses 37 to 48, wherein the compound of Formula I-Lu or Ia-Lu is administered at the same time as the compound of Formula I-Ac or Ia-Ac.

50. The use of any one of clauses 37 to 49, wherein the compound of Formula I-Lu or Ia-Lu is administered about 1 hour prior to the compound of Formula I-Ac or Ia-Ac; or about 12 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 24 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 48 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 72 hours prior to the compound of Formula I-Ac or Ia-Ac.

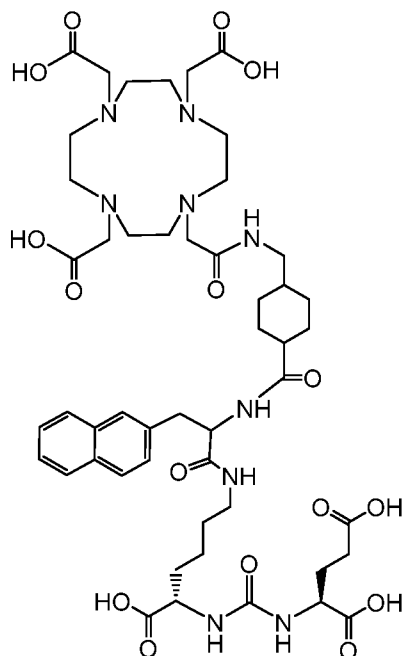
51. The use of any one of clauses 37 to 50, wherein the compound of Formula I-Ac or Ia-Ac is administered about 1 hour prior to the compound of Formula I-Lu or Ia-Lu; or about 12 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 24 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 48 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 72 hours prior to the compound of Formula I-Lu or Ia-Lu.

52. The use of clause 49, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of I-Lu or Ia-Lu and I-Ac or Ia-Ac.

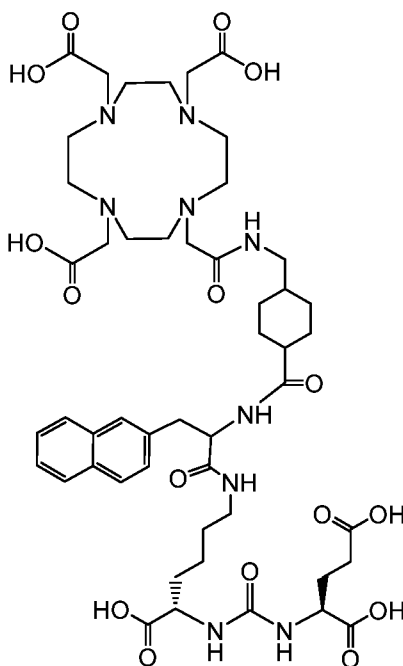
53. The use of clause 50, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

54. The use of clause 51, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

55. A composition comprising a compound of the Formula I-Lu,



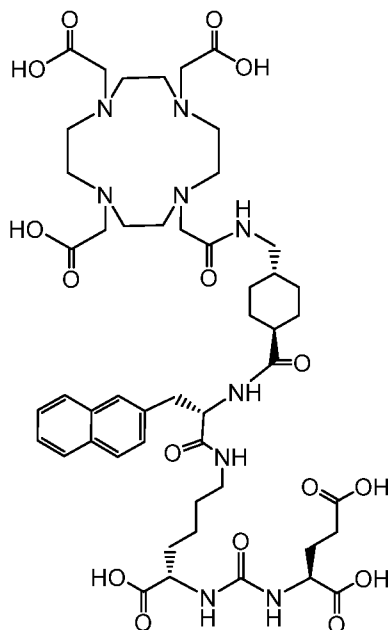
wherein the compound is complexed with ^{177}Lu , in a therapeutically effective amount, for use in the treatment of cancer in a patient, in combination with a therapeutically effective amount of a compound of the Formula I-Ac



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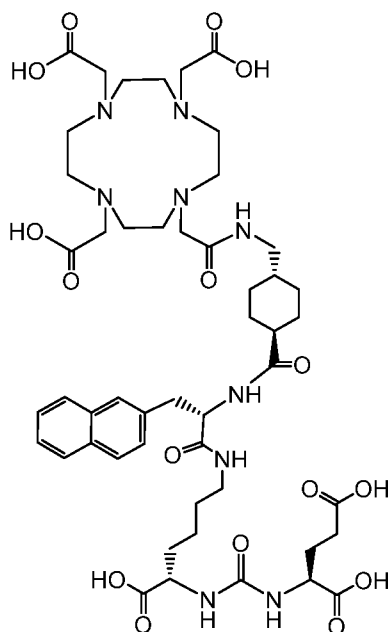
wherein the compound is complexed with ^{225}Ac .

56. The composition of clause 55, wherein the compound of the Formula I-Lu is of the Formula Ia-Lu



wherein the compound is complexed with ^{177}Lu .

57. The composition of clause 55, wherein the compound of the Formula I-Ac is of the Formula Ia-Ac



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wherein the compound is complexed with ^{225}Ac .

58. The composition of any one of clauses 55 to 57, wherein the cancer is associated with expression of prostate specific membrane antigen (PSMA).

59. The composition of any one of clauses 55 to 58, wherein the cancer is selected from the group consisting of prostate cancer, metastatic castration-resistant prostate cancer (mCRPC), thyroid cancer, renal cell carcinoma, transitional cell carcinoma of the

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bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, soft tissue sarcoma, and breast carcinoma.

5 60. The composition of any one of clauses 55 to 59, wherein the cancer is prostate cancer.

61. The composition of any one of clauses 55 to 60, wherein the cancer is metastatic castration-resistant prostate cancer (mCRPC).

62. The composition of any one of clauses 58 to 61, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq.

10 63. The composition of any one of clauses 58 to 62, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq.

64. The composition of any one of clauses 58 to 63, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 7.4 GBq.

15 65. The composition of any one of clauses 58 to 64, wherein the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq; or about 5 MBq to about 10 MBq; or about 5 MBq to about 7MBq.

66. The composition of any one of clauses 58 to 65, wherein the therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq.

20 67. The composition of any one of clauses 58 to 66, wherein the compound of Formula I-Lu or Ia-Lu is administered at the same time as the compound of Formula I-Ac or Ia-Ac.

25 68. The composition of any one of clauses 58 to 67, wherein the compound of Formula I-Lu or Ia-Lu is administered about 1 hour prior to the compound of Formula I-Ac or Ia-Ac; or about 12 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 24 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 48 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 72 hours prior to the compound of Formula I-Ac or Ia-Ac.

30 69. The composition of any one of clauses 58 to 68, wherein the compound of Formula I-Ac or Ia-Ac is administered about 1 hour prior to the compound of Formula I-Lu or Ia-Lu; or about 12 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 24 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 48 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 72 hours prior to the compound of Formula I-Lu or Ia-Lu.

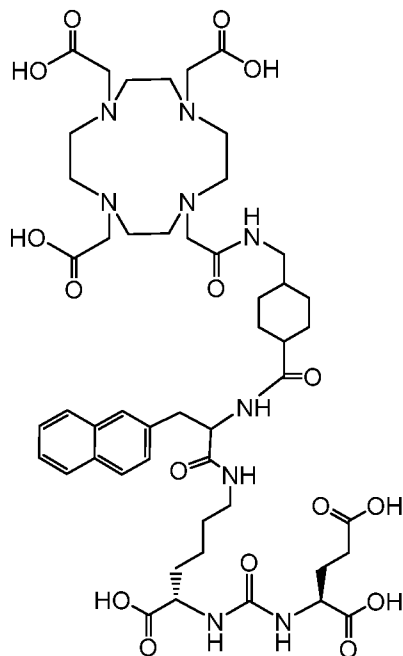
70. The composition of clause 67, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly

cycle for from about 1 to about 7 cycles following the administration of I-Lu or Ia-Lu and I-Ac or Ia-Ac.

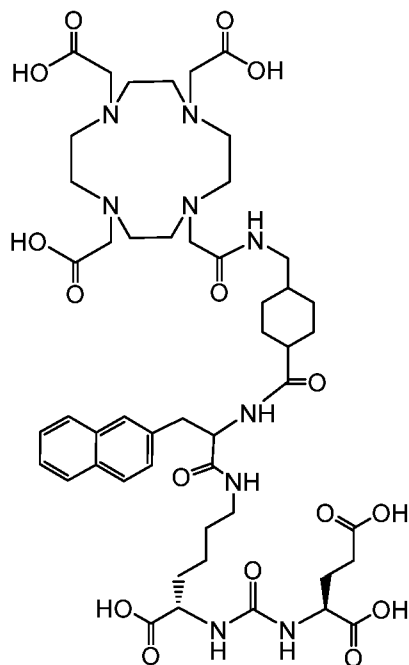
71. The composition of clause 68, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

72. The composition of clause 69, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

73. A medicament comprising a compound of the Formula I-Lu

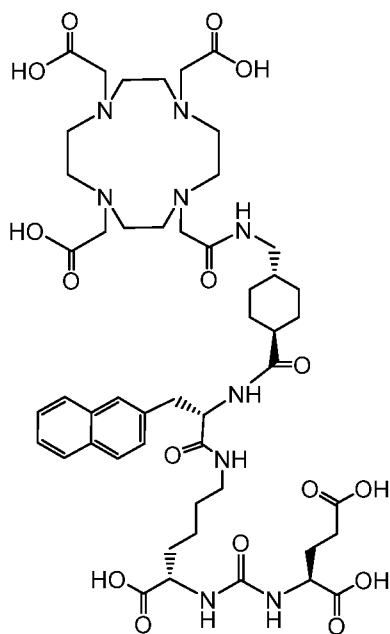


wherein the compound is complexed with ^{177}Lu , in a therapeutically effective amount, combined with a therapeutically effective amount of a compound of the Formula I-Ac



wherein the compound is complexed with ^{225}Ac .

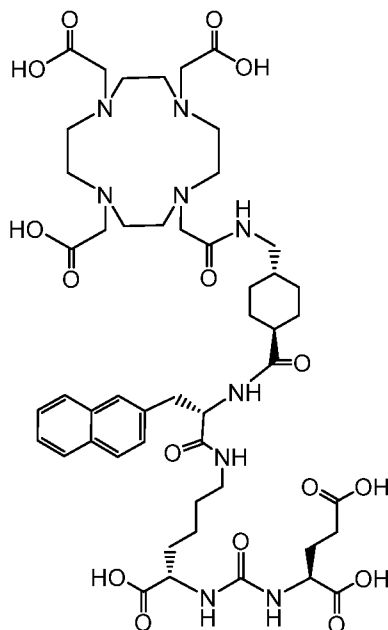
74. The medicament of clause 73, wherein the compound of the Formula I-Lu is of the Formula Ia-Lu



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wherein the compound is complexed with ^{177}Lu .

75. The medicament of clause 73, wherein the compound of the Formula I-Ac is of the Formula Ia-Ac



wherein the compound is complexed with ^{225}Ac .

76. The medicament of any one of clauses 73 to 75, wherein medicament provides a synergistic effect on a cancer associated with expression of prostate specific membrane antigen (PSMA).

77. The medicament of any one of clauses 73 to 76, wherein the cancer is selected from the group consisting of prostate cancer, metastatic castration-resistant prostate cancer (mCRPC), thyroid cancer, renal cell carcinoma, transitional cell carcinoma of the bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, soft tissue sarcoma, and breast carcinoma.

78. The medicament of any one of clauses 73 to 77, wherein the cancer is prostate cancer.

79. The medicament of any one of clauses 73 to 78, wherein the cancer is metastatic castration-resistant prostate cancer (mCRPC).

80. The medicament of any one of clauses 73 to 79, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq.

81. The medicament of any one of clauses 73 to 80, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq.

82. The medicament of any one of clauses 73 to 81, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 7.4 GBq.

83. The medicament of any one of clauses 73 to 82, wherein the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq; or about 5 MBq to about 10 MBq; or about 5 MBq to about 7MBq.

84. The medicament of any one of clauses 73 to 83, wherein the therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq.

85. The medicament of any one of clauses 73 to 84, wherein the compound of Formula I-Lu or Ia-Lu is administered at the same time as the compound of Formula I-Ac or Ia-Ac.

86. The medicament of any one of clauses 73 to 85, wherein the compound of Formula I-Lu or Ia-Lu is administered about 1 hour prior to the compound of Formula I-Ac or Ia-Ac; or about 12 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 24 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 48 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 72 hours prior to the compound of Formula I-Ac or Ia-Ac.

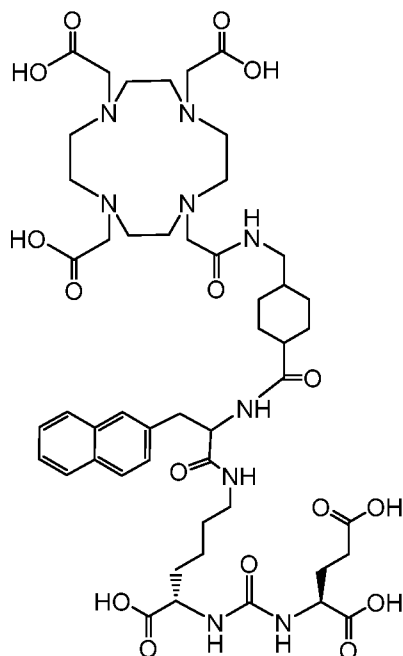
87. The medicament of any one of clauses 73 to 86, wherein the compound of Formula I-Ac or Ia-Ac is administered about 1 hour prior to the compound of Formula I-Lu or Ia-Lu; or about 12 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 24 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 48 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 72 hours prior to the compound of Formula I-Lu or Ia-Lu.

88. The medicament of clause 87, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of I-Lu or Ia-Lu and I-Ac or Ia-Ac.

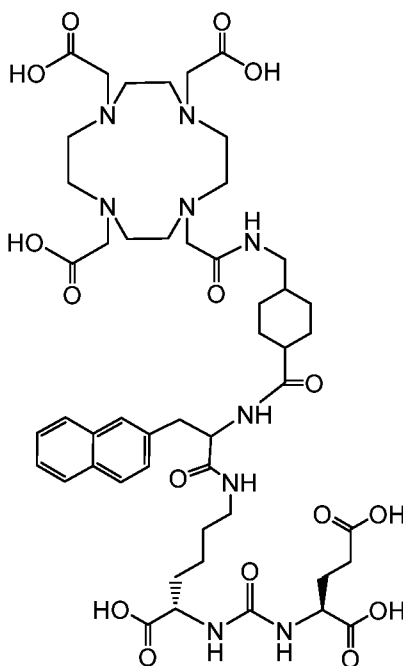
89. The medicament of clause 88, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

90. The medicament of clause 89, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

91. A synergistic composition comprising a compound of the Formula I-Lu,



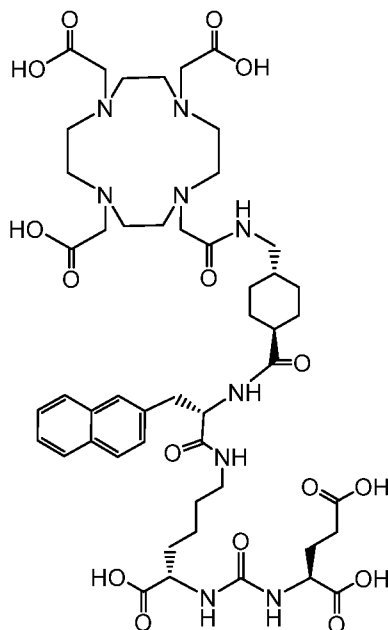
wherein the compound is complexed with ^{177}Lu , in a therapeutically effective amount, for use in the treatment of cancer in a patient, in combination with a therapeutically effective amount of a compound of the Formula I-Ac



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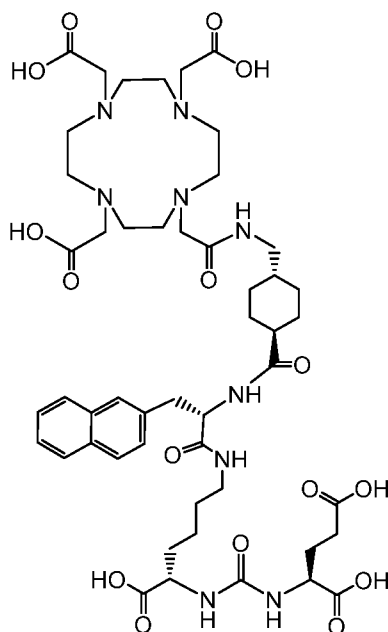
wherein the compound is complexed with ^{225}Ac .

92. The synergistic composition of clause 91, wherein the compound of the Formula I-Lu is of the Formula Ia-Lu



wherein the compound is complexed with ^{177}Lu .

93. The synergistic composition of clause 91, wherein the compound of the Formula I-Ac is of the Formula Ia-Ac



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wherein the compound is complexed with ^{225}Ac .

94. The synergistic composition of any one of clauses 91 to 93, wherein the cancer is associated with expression of prostate specific membrane antigen (PSMA).

95. The synergistic composition of any one of clauses 91 to 94, wherein the cancer is selected from the group consisting of prostate cancer, metastatic castration-resistant prostate cancer (mCRPC), thyroid cancer, renal cell carcinoma, transitional cell carcinoma of

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the bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, soft tissue sarcoma, and breast carcinoma.

5 96. The synergistic composition of any one of clauses 91 to 95, wherein the cancer is prostate cancer.

97. The synergistic composition of any one of clauses 91 to 96, wherein the cancer is metastatic castration-resistant prostate cancer (mCRPC).

98. The synergistic composition of any one of clauses 91 to 97, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq.

10 99. The synergistic composition of any one of clauses 91 to 98, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq.

100. The synergistic composition of any one of clauses 91 to 99, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 7.4 GBq.

15 101. The synergistic composition of any one of clauses 91 to 100, wherein the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq; or about 5 MBq to about 10 MBq; or about 5 MBq to about 7MBq.

102. The synergistic composition of any one of clauses 91 to 101, wherein the therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq.

20 103. The synergistic composition of any one of clauses 91 to 102, wherein the compound of Formula I-Lu or Ia-Lu is administered at the same time as the compound of Formula I-Ac or Ia-Ac.

25 104. The synergistic composition of any one of clauses 91 to 103, wherein the compound of Formula I-Lu or Ia-Lu is administered about 1 hour prior to the compound of Formula I-Ac or Ia-Ac; or about 12 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 24 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 48 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 72 hours prior to the compound of Formula I-Ac or Ia-Ac.

30 105. The synergistic composition of any one of clauses 91 to 104, wherein the compound of Formula I-Ac or Ia-Ac is administered about 1 hour prior to the compound of Formula I-Lu or Ia-Lu; or about 12 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 24 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 48 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 72 hours prior to the compound of Formula I-Lu or Ia-Lu.

106. The synergistic composition of clause 103, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of I-Lu or Ia-Lu and I-Ac or Ia-Ac.

5 107. The synergistic composition of clause 104, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

10 108. The synergistic composition of clause 105, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

DEFINITIONS

15 As used herein, “functionally active PSMA” means a cell surface membrane-bound glycoprotein that binds to a PSMA ligand. It will be appreciated that PSMA ligands are well known to those skilled in the art such as those described in US patent publication no. US 2010/0324008 A1, incorporated herein by reference.

20 As used herein, “clinical benefit” means a response of a patient to treatment with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac, where the response includes overall survival of the patient, ability to receive four or more cycles of therapy (e.g., four weeks of therapy) with Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac, inhibition of tumor growth, stable disease, a partial response, and/or a complete response, among other clinical benefits defined by the Food and Drug Administration in the United States of America.

25 As used herein, “inhibition of tumor growth” means reduction in tumor size, complete disappearance of a tumor, or growth of a patient tumor of less than 30% over the course of therapy with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac.

As used herein, “stable disease” means no material progression of disease in a patient over the course of therapy with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac.

30 As used herein, “a partial response” means a decrease in tumor size of 30% or greater in a patient treated with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac.

As used herein, “a complete response” means the disappearance of detectable disease in a patient treated with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac.

As used herein, “prior treatment” means the patient has been treated with at least one prior treatment known in the art. It will be appreciated that a prior treatment can be any treatment known to those of skill in the art, including, but not limited, chemotherapeutic agent, surgery, radiation therapy, immunotherapy, photodynamic therapy, stem cell therapy, hyperthermia, and the like. Prior treatments can include systemic treatments including, but not limited to treatment with abiraterone, orteronel, galeterone, seviteronel, apalutamide, enzalutamide, palifosfamide, 5-fluorouracil, capecitabine, pemetrexed, cisplatin, carboplatin, gemcitabine, paclitaxel, vinorelbine, eribulin, docetaxel, cyclophosphamide, doxorubicin, regorafenib, and combinations thereof.

As used herein, the term “alkyl” includes a chain of carbon atoms, which is optionally branched. It will be further understood that in certain embodiments, alkyl is advantageously of limited length, including C₁-C₂₄, C₁-C₁₂, C₁-C₈, C₁-C₆, and C₁-C₄. Illustratively, such particularly limited length alkyl groups, including C₁-C₃, C₁-C₆, and C₁-C₄ may be referred to as lower alkyl. It is appreciated herein that shorter alkyl, alkenyl, and/or alkynyl groups may add less lipophilicity to the compound and accordingly will have different pharmacokinetic behavior. In some embodiments, it will be understood, in each case, that the recitation of alkyl refers to alkyl as defined herein, and optionally lower alkyl. Illustrative alkyl groups include, but not limited to, methyl, ethyl, n-propyl, isopropyl, n-butyl, isobutyl, sec-butyl, tert-butyl, pentyl, 2-pentyl, 3-pentyl, neopentyl, hexyl, heptyl, octyl, and the like. As used herein, a “carboxyalkyl” group includes a combination of an “alkyl” group as described herein with a “carboxy” group. As used herein, a “hydroxyalkyl” group includes a combination of an “alkyl” group as described herein with a “hydroxy” group. As used herein, a “aminoalkyl” group includes a combination of an “alkyl” group as described herein with a “amino” group.

As used herein, the term “heteroalkyl” includes a chain of atoms that includes both carbon and at least one heteroatom, and is optionally branched. Illustrative heteroatoms include nitrogen, oxygen, and sulfur. In certain variations, illustrative heteroatoms also include phosphorus, and selenium.

As used herein, the term “aryl” includes monocyclic and polycyclic aromatic carbocyclic groups having from 6 to 14 ring carbon atoms, each of which may be optionally substituted. Illustrative aromatic carbocyclic groups described herein include, but are not limited to, phenyl, naphthyl, and the like. As used herein, the term “heteroaryl” includes aromatic heterocyclic groups, having from 5 to 10 ring atoms, each of which may be optionally substituted. Illustrative aromatic heterocyclic groups include, but are not limited to, pyridinyl, pyrimidinyl, pyrazinyl, triazinyl, tetrazinyl, quinolinyl, quinazolinyl, quinoxalinyl, thienyl,

pyrazolyl, imidazolyl, oxazolyl, thiazolyl, isoxazolyl, isothiazolyl, oxadiazolyl, thiadiazolyl, triazolyl, benzimidazolyl, benzoxazolyl, benzthiazolyl, benzisoxazolyl, benzisothiazolyl, and the like. As used herein, the term “heteroarylalkyl” includes a combination of an “alkyl” group as described herein with a “heteroaryl” group described herein. As used herein, the term
5 “arylalkyl” includes a combination of an “alkyl” group as described herein with a “aryl” group described herein, for example a benzyl group.

The term “optionally substituted” as used herein includes the replacement of hydrogen atoms with other functional groups on the radical that is optionally substituted. Such other functional groups illustratively include, but are not limited to, amino, hydroxyl, halo, thiol,
10 alkyl, haloalkyl, heteroalkyl, aryl, arylalkyl, arylheteroalkyl, heteroaryl, heteroarylalkyl, heteroarylheteroalkyl, nitro, sulfonic acids and derivatives thereof, carboxylic acids and derivatives thereof, and the like. Illustratively, any of amino, hydroxyl, thiol, alkyl, haloalkyl, heteroalkyl, aryl, arylalkyl, arylheteroalkyl, heteroaryl, heteroarylalkyl, heteroarylheteroalkyl, and/or sulfonic acid is optionally substituted.

As used herein, the term “administering” as used herein includes all means of introducing a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac and/or a PSMA ligand-imaging conjugate as described herein to the patient, including, but not limited to, oral (po), intravenous (iv), intramuscular (im), subcutaneous (sc), transdermal, inhalation, buccal,
15 ocular, sublingual, vaginal, rectal, and the like. A combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac and/or a PSMA ligand-imaging conjugate as described herein may be administered in unit dosage forms and/or formulations containing conventional nontoxic pharmaceutically-acceptable carriers, adjuvants, and vehicles.

As used herein, “becquerel” means a SI derived unit of radioactivity as it is commonly understood by one of skill in the art. One becquerel is defined as the activity of a quantity of
25 radioactive material in which one nucleus decays per second. A becquerel is therefore equivalent to an inverse second, s^{-1} . The becquerel is known to one of skill in the art as the successor of the curie (Ci), an older, non-SI unit of radioactivity based on the activity of 1 gram of radium-226. The curie is defined as $3.7 \cdot 10^{10} s^{-1}$, or 37 GBq.

As used herein, “curie” or “Ci” means a unit of radioactivity named after the French
30 physicist and chemist Marie Curie as commonly understood by one of skill in the art. The prefixes milli and micro are from the metric system and represent .001 and .000001, respectively. So, a millicurie (mCi) is .001 curie. A microcurie (μ Ci) is .000001 curie.

DETAILED DESCRIPTION

The embodiments of the numbered clauses provided in the summary above, or any combination thereof, are contemplated for combination with any of the embodiments described in the Detailed Description section of this patent application.

Referring to FIG. 1, the method design can be described according to the schematic
5 shown. In some embodiments, stratification factors for the design include, but are not limited to serum lactate dehydrogenase (LDH) (≤ 260 IU/L v. >260 IU/L), presence of liver metastases, ECOG score (0-1 v. 2), inclusion of NAAD in best supportive/best standard of care, and the like. In some embodiments, the primary endpoint can be overall survival. In some
10 embodiments, secondary endpoints include, but are not limited to, radiographic progression-free survival (rPFS), RECIST response, time to first symptomatic skeletal event (SSE), and the like. In some embodiments, additional secondary endpoints include, but are not limited to, safety and tolerability, heather-related quality of life (HRQoL; EQ-5D-5L, FACT-P and Brief Pain
Inventory – Short FORM [BPI-SF]), health economics, progression-free survival (PFS)
15 (radiological, clinical or PSA progression), biochemical response, such as PSA levels, alkaline phosphatase level, and/or lactate dehydrogenase level. In some embodiments, an endpoint for the treatment methods described herein can be a patient who has achieved a $\geq 50\%$ decrease from baseline that is confirmed by a second PSA measurement ≥ 4 weeks. In some
20 embodiments, an endpoint for the treatment methods described herein can be a patient who has achieved a $\geq 40\%$ decrease from baseline that is confirmed by a second PSA measurement ≥ 4 weeks. In some embodiments, an endpoint for the treatment methods described herein can be a patient who has achieved a $\geq 30\%$ decrease from baseline that is confirmed by a second PSA measurement ≥ 4 weeks.

In one embodiment, the methods described herein can be used for both human clinical
25 medicine and veterinary applications. Thus, a “patient” can be administered a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac and/or PSMA ligand-imaging conjugates described herein, and can be human or, in the case of veterinary applications, can be a laboratory, agricultural, domestic, or wild animal. In one aspect, the patient can be a human, a
laboratory animal such as a rodent (*e.g.*, mice, rats, hamsters, etc.), a rabbit, a monkey, a
chimpanzee, domestic animals such as dogs, cats, and rabbits, agricultural animals such as
30 cows, horses, pigs, sheep, goats, and wild animals in captivity such as bears, pandas, lions, tigers, leopards, elephants, zebras, giraffes, gorillas, dolphins, and whales.

In some embodiments, patients with PSMA positive scans can be randomized in a 2:1 ratio to receive either a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac plus best supportive/best standard of care or to receive best supportive/best standard of care only. In

some embodiments, best supportive/best standard of care can be determined by the treating physician/investigator. In some embodiments, best supportive/best standard of care can be determined by the treating physician/investigator, but will exclude investigational agents, cytotoxic chemotherapy, other systemic radioisotopes, and hemi-body radiotherapy. In some 5
embodiments, novel androgen axis drugs [NAADs], such as abiraterone or enzalutamide, are allowed.

In some embodiments, patients will be monitored throughout the 6 to 10-month treatment period for survival, disease progression, and adverse events. In some embodiments, a long-term follow-up period can include the collection of survival and treatment updates, 10
adverse events assessment, as well as blood for hematology and chemistry testing.

In some embodiments, the patient is 18 Years of age or older. In some embodiments, the patient is a male. In some embodiments, the patient has previously been diagnosed with prostate cancer. In some embodiments, the patient has been previously diagnosed with metastatic castration-resistant prostate cancer (mCRPC). In some embodiments, the patient meets one or 15
more criteria, selected from the group consisting of Eastern Cooperative Oncology Group (ECOG) performance status of 0 to 2; a life expectancy at least 6 months; histological, pathological, and/or cytological confirmation of prostate cancer; a positive ⁶⁸Ga-PSMA-11 PET/CT scan; prior orchiectomy and/or ongoing androgen deprivation therapy and a castrate level of serum testosterone (<50 ng/dL or <1.7 nmol/L); previously received at least one 20
NAAD, such as enzalutamide and/or abiraterone; previously treated with at least 1 or 2 previous taxane regimens, wherein a taxane regimen comprises a minimum exposure of 2 cycles of a taxane, or previously received only one taxane regimen, and a. the patient is not willing to receive a second taxane regimen, or b. The patient's physician deems him unsuitable to receive a second taxane regimen, such as due to frailty assessed by geriatric or health status evaluation 25
or intolerance; progressive mCRPC, such as documented progressive mCRPC based on at least one criteria, such as a. serum PSA progression defined as 2 consecutive increases in PSA over a previous reference value measured at least 1 week prior, where the minimal start value is 2.0 ng/mL, b. soft-tissue progression defined as an increase $\geq 20\%$ in the sum of the diameter (SOD) (short axis for nodal lesions and long axis for non-nodal lesions) of all target lesions based on 30
the smallest SOD since treatment started or the appearance of one or more new lesions, and c. progression of bone disease, such as evaluable disease or new bone lesions(s) by bone scan (2+2 PCWG3 criteria); at least one metastatic lesion that is present on baseline CT, MRI, or bone scan imaging obtained ≤ 28 days prior to beginning therapy with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac ; recovered to \leq Grade 2 from all clinically

significant toxicities related to prior therapies, such as prior chemotherapy, radiation, immunotherapy, and the like; adequate organ function, such as a. bone marrow reserve including white blood cell (WBC) count $\geq 2.5 \times 10^9/L$ ($2.5 \times 10^9/L$ is equivalent to $2.5 \times 10^3/\mu L$ and $2.5 \times K/\mu L$ and $2.5 \times 10^3/cumm$ and $2500/\mu L$) or absolute neutrophil count (ANC) $\geq 1.5 \times 10^9/L$ ($1.5 \times 10^9/L$ is equivalent to $1.5 \times 10^3/\mu L$ and $1.5 \times K/\mu L$ and $1.5 \times 10^3/cumm$ and $1500/\mu L$), platelets $\geq 100 \times 10^9/L$ ($100 \times 10^9/L$ is equivalent to $100 \times 10^3/\mu L$ and $100 \times K/\mu L$ and $100 \times 10^3/cumm$ and $100,000/\mu L$), and/or hemoglobin $\geq 9 \text{ g/dL}$ (9 g/dL is equivalent to 90 g/L and 5.59 mmol/L); b. hepatic, such as total bilirubin $\leq 1.5 \times$ the institutional upper limit of normal (ULN) (for patients with known Gilbert's Syndrome $\leq 3 \times$ ULN is permitted), alanine aminotransferase (ALT) or aspartate aminotransferase (AST) $\leq 3.0 \times$ ULN OR $\leq 5.0 \times$ ULN for patients with liver metastases, and c. renal, such as serum creatinine $\leq 1.5 \times$ ULN or creatinine clearance $\geq 50 \text{ mL/min}$; albumin $> 3.0 \text{ g/dL}$ (3.0 g/dL is equivalent to 30 g/L); and a stable bisphosphonate or denosumab regimen for ≥ 30 days prior to treatment.

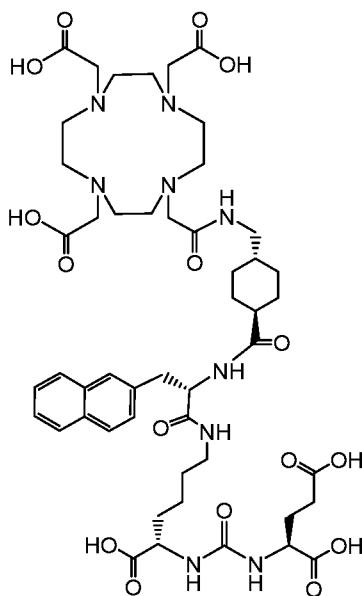
In some embodiments, a patient may not receive treatment if the patient has one of more of previous treatment with Strontium-89, Samarium-153, Rhenium-186, Rhenium-188, Radium-223 or hemi-body irradiation within about 6 months prior treatment; previous PSMA-targeted radioligand therapy; previous systemic anti-cancer therapy (e.g. chemotherapy, immunotherapy or biological therapy [including monoclonal antibodies]) within about 28 days prior to treatment; previous administration of investigational agents within about 28 days prior to treatment; a known hypersensitivity to the components of the therapy or its analogs; any other concurrent cytotoxic chemotherapy, immunotherapy, radioligand therapy, or investigational therapy; a transfusion within about 30 days of treatment; a history of CNS metastases that have received therapy (surgery, radiotherapy, gamma knife) and are neurologically stable, asymptomatic, and not receiving corticosteroids for the purposes of maintaining neurologic integrity; a superscan as seen in the baseline bone scan; a symptomatic cord compression, or clinical or radiologic findings indicative of impending cord compression; concurrent serious (as determined by a physician) medical conditions, including, but not limited to, New York Heart Association class III or IV congestive heart failure, history of congenital prolonged QT syndrome, uncontrolled infection, active hepatitis B or C, or other significant comorbid conditions that in the opinion of the investigator would impair treatment or cooperation; or been diagnosed with other malignancies that are expected to alter life expectancy or may interfere with disease assessment.

In various embodiments, the cancers described herein can be a cancer cell population that is tumorigenic, including benign tumors and malignant tumors, or the cancer can be non-

tumorigenic. The cancer can arise spontaneously or by such processes as mutations present in the germline of the patient or somatic mutations, or the cancer can be chemically-, virally-, or radiation-induced. Cancers applicable to the present disclosure described herein include, but are not limited to, a glioma, a carcinoma, a sarcoma, a lymphoma, a melanoma, a mesothelioma, a nasopharyngeal carcinoma, a leukemia, an adenocarcinoma, and a myeloma.

In some aspects the cancers can be lung cancer, bone cancer, pancreatic cancer, skin cancer, cancer of the head, cancer of the neck, cutaneous melanoma, intraocular melanoma uterine cancer, ovarian cancer, endometrial cancer, rectal cancer, stomach cancer, colon cancer, breast cancer, triple negative breast cancer, metastatic breast cancer, carcinoma of the fallopian tubes, carcinoma of the endometrium, carcinoma of the cervix, carcinoma of the vagina, carcinoma of the vulva, Hodgkin's Disease, cancer of the esophagus, cancer of the small intestine, cancer of the endocrine system, cancer of the thyroid gland, cancer of the parathyroid gland, non-small cell lung cancer, cancer of the adrenal gland, sarcoma of soft tissue, cancer of the urethra, cancer of the penis, prostate cancer, chronic leukemia, acute leukemia, lymphocytic lymphomas, pleural mesothelioma, cancer of the bladder, Burkitt's lymphoma, cancer of the ureter, cancer of the kidney, renal cell carcinoma, carcinoma of the renal pelvis, neoplasms of the central nervous system (CNS), primary CNS lymphoma, spinal axis tumors, glioma, brain stem glioma, pituitary adenoma, and adenocarcinoma of the gastroesophageal junction.

Compound Ia has the formula



Ia

wherein ^{177}Lu is complexed to the compound in Ia-Lu, and ^{225}Ac is complexed to the compound in Ia-Ac.

In other embodiments, any of a variety of PSMA ligand-imaging conjugates detectable by PET imaging, SPECT imaging, and the like can be used. The exact manner of imaging is not limited to the imaging agents described herein. Collectively, the PSMA ligand-imaging conjugates useful for imaging described herein, including those described by formulas and the agents useful for PET imaging, SPECT imaging, etc. are referred to as “PSMA ligand-imaging conjugates.”

In one embodiment, the Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac and/or PSMA ligand-imaging conjugates described herein bind to expressed PSMA on cancer cells. In one illustrative aspect, the Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac and/or PSMA ligand-imaging conjugates are capable of differentially binding to PSMA on cancer cells compared to normal cells due to preferential expression (or over-expression) of PSMA on the cancer cells.

In some embodiment, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac and PSMA ligand-imaging conjugates described herein may be administered as a formulation in association with one or more pharmaceutically acceptable carriers. In some aspects of these embodiments, the combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac will be co-formulated. In some aspects of these embodiments, the combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac will be administered as individually formulated agents. The carriers can be excipients. The choice of carrier will to a large extent depend on factors such as the particular mode of administration, the effect of the carrier on solubility and stability, and the nature of the dosage form. Pharmaceutical compositions suitable for the delivery of a combination of Compounds or Ia-Lu, and I-Ac or Ia-Ac and PSMA ligand-imaging conjugates described herein and methods for their preparation will be readily apparent to those skilled in the art. Such compositions and methods for their preparation may be found, for example, in Remington: The Science & Practice of Pharmacy, 21th Edition (Lippincott Williams & Wilkins, 2005), incorporated herein by reference.

In one illustrative aspect, a pharmaceutically acceptable carrier includes any and all solvents, dispersion media, coatings, antibacterial and antifungal agents, isotonic and absorption delaying agents, and the like, and combinations thereof, that are physiologically compatible. In some embodiments, the carrier is suitable for parenteral administration. Pharmaceutically acceptable carriers include sterile aqueous solutions or dispersions and sterile powders for the extemporaneous preparation of sterile injectable solutions or dispersions. Supplementary active compounds can also be incorporated into compositions of the present disclosure.

In various embodiments, liquid formulations may include suspensions and solutions. Such formulations may comprise a carrier, for example, water, ethanol, polyethylene glycol,

propylene glycol, methylcellulose or a suitable oil, and one or more emulsifying agents and/or suspending agents. Liquid formulations may also be prepared by the reconstitution of a solid.

In one embodiment, an aqueous suspension may contain the active materials in admixture with appropriate excipients. Such excipients are suspending agents, for example, sodium carboxymethylcellulose, methylcellulose, hydroxypropylmethylcellulose, sodium alginate, polyvinylpyrrolidone, gum tragacanth and gum acacia; dispersing or wetting agents which may be a naturally-occurring phosphatide, for example, lecithin; a condensation product of an alkylene oxide with a fatty acid, for example, polyoxyethylene stearate; a condensation product of ethylene oxide with a long chain aliphatic alcohol, for example, heptadecaethyleneoxycetanol; a condensation product of ethylene oxide with a partial ester derived from fatty acids and a hexitol such as polyoxyethylene sorbitol monooleate; or a condensation product of ethylene oxide with a partial ester derived from fatty acids and hexitol anhydrides, for example, polyoxyethylene sorbitan monooleate. The aqueous suspensions may also contain one or more preservatives, for example, ascorbic acid, ethyl, n-propyl, or p-hydroxybenzoate; or one or more coloring agents.

In one illustrative embodiment, dispersible powders and granules suitable for preparation of an aqueous suspension by the addition of water provide the active ingredient in admixture with a dispersing or wetting agent, suspending agent and one or more preservatives. Additional excipients, for example, coloring agents, may also be present.

Suitable emulsifying agents may be naturally-occurring gums, for example, gum acacia or gum tragacanth; naturally-occurring phosphatides, for example, soybean lecithin; and esters including partial esters derived from fatty acids and hexitol anhydrides, for example, sorbitan mono-oleate, and condensation products of the said partial esters with ethylene oxide, for example, polyoxyethylene sorbitan monooleate.

In other embodiments, isotonic agents, for example, sugars, polyalcohols such as mannitol, sorbitol, or sodium chloride can be included in the composition. Prolonged absorption of injectable compositions can be brought about by including in the composition an agent which delays absorption, for example, monostearate salts and gelatin.

Illustrative formats for oral administration include tablets, capsules, elixirs, syrups, and the like.

Depending upon the cancer type as described herein, the route of administration and/or whether a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac and/or PSMA ligand-imaging conjugates are administered locally or systemically, a wide range of permissible dosages are contemplated herein, including doses falling in the range from about 1 MBq to

about 4 MBq of I-Ac or Ia-Ac. In some embodiments, permissible dosages for I-Lu or Ia-Lu are contemplated herein in the units GBq, including doses falling in the range from about 2 GBq to about 13 GBq. The dosages may be single or divided, and may administered according to a wide variety of protocols, including q.d., b.i.d., t.i.d., or even every other day, biweekly
5 (b.i.w.), once a week, once a month, once a quarter, and the like. In each of these cases it is understood that the therapeutically effective amounts described herein correspond to the instance of administration, or alternatively to the total daily, weekly, monthly, or quarterly dose, as determined by the dosing protocol. In some embodiments, a combination of compounds of the formula I-Lu or Ia-Lu, and I-Ac or Ia-Ac can be administered on independent schedules of
10 once, or once per week, or once every two weeks, or once every three weeks, or once every four weeks, or once every five weeks, or once every six weeks, or once every seven weeks, or once every eight weeks, and the like

In one aspect, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac or a PSMA ligand-imaging conjugate as described herein may be administered directly into the
15 blood stream, into muscle, or into an internal organ. Suitable routes for such parenteral administration include intravenous, intraarterial, intraperitoneal, intrathecal, epidural, intracerebroventricular, intraurethral, intrasternal, intracranial, intratumoral, intramuscular and subcutaneous delivery. Suitable means for parenteral administration include needle (including microneedle) injectors, needle-free injectors and infusion techniques.

In one illustrative aspect, parenteral formulations are typically aqueous solutions which may contain carriers or excipients such as salts, carbohydrates and buffering agents (preferably at a pH of from 3 to 9), but, for some applications, they may be more suitably formulated as a sterile non-aqueous solution or as a dried form to be used in conjunction with a suitable vehicle such as sterile, pyrogen-free water. In other embodiments, any of the liquid formulations
25 described herein may be adapted for parenteral administration of the I-Lu or Ia-Lu, and I-Ac or Ia-Ac or PSMA ligand-imaging conjugates described herein. The preparation of parenteral formulations under sterile conditions, for example, by lyophilization under sterile conditions, may readily be accomplished using standard pharmaceutical techniques well known to those skilled in the art. In one embodiment, the solubility of a combination of Compounds I-Lu or Ia-
30 Lu, and I-Ac or Ia-Ac or a PSMA ligand-imaging conjugate used in the preparation of a parenteral formulation may be increased by the use of appropriate formulation techniques, such as the incorporation of solubility-enhancing agents.

In various embodiments, formulations for parenteral administration may be formulated for immediate and/or modified release. In one illustrative aspect, a combination of the active

agents of the present disclosure (i.e., Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac or PSMA ligand-imaging conjugates) may be administered in a time release formulation, for example in a composition which includes a slow release polymer. The active Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac or PSMA ligand-imaging conjugates can be prepared with carriers that will protect Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac or PSMA ligand-imaging conjugate against rapid release, such as a controlled release formulation, including implants and microencapsulated delivery systems. Biodegradable, biocompatible polymers can be used, such as ethylene vinyl acetate, polyanhydrides, polyglycolic acid, collagen, polyorthoesters, polylactic acid and poly(lactic, polyglycolic copolymers (PGLA). Methods for the preparation of such formulations are generally known to those skilled in the art. In another embodiment, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac or PSMA ligand-imaging conjugates described herein or compositions comprising the Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac or PSMA ligand-imaging conjugates may be continuously administered, where appropriate.

In one embodiment, a kit is provided. If a combination of active Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac and PSMA ligand-imaging conjugates is to be administered, two or more pharmaceutical compositions may be combined in the form of a kit suitable for sequential administration or co-administration of the compositions. Such a kit comprises two or more separate pharmaceutical compositions, at least one of which contains a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac or PSMA ligand-imaging conjugate described herein, and means for separately retaining the compositions, such as a container, divided bottle, or divided foil packet. In another embodiment, compositions comprising one or more of a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac or PSMA ligand-imaging conjugates described herein, in containers having labels that provide instructions for use of a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac or PSMA ligand-imaging conjugates for patient selection and/or treatment are provided.

In one embodiment, sterile injectable solutions can be prepared by incorporating the active agent in the required amount in an appropriate solvent with one or a combination of ingredients described above, as required, followed by filtered sterilization. Typically, dispersions are prepared by incorporating the active combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac or PSMA ligand-imaging conjugate into a sterile vehicle which contains a dispersion medium and any additional ingredients of those described above. In the case of sterile powders for the preparation of sterile injectable solutions, the preferred methods of preparation are vacuum drying and freeze-drying which yields a powder of the active ingredient

plus any additional desired ingredient from a previously sterile-filtered solution thereof, or the ingredients may be sterile-filtered together.

The composition can be formulated as a solution, microemulsion, liposome, or other ordered structure suitable to high drug concentration. The carrier can be a solvent or dispersion medium containing, for example, water, ethanol, polyol (for example, glycerol, propylene glycol, and liquid polyethylene glycol, and the like), and suitable mixtures thereof. In one embodiment, the proper fluidity can be maintained, for example, by the use of a coating such as lecithin, by the maintenance of the required particle size in the case of dispersion and by the use of surfactants.

10 Any effective regimen for administering a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac can be used. For example, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac can be administered as single doses, or the doses can be divided and administered as a multiple-dose daily regimen. Further, a staggered regimen, for example, one to five days per week can be used as an alternative to daily treatment, and for the purpose of the methods described herein, such intermittent or staggered daily regimen is considered to be equivalent to every day treatment and is contemplated. In one illustrative embodiment the patient is treated with multiple injections of a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac to treat the cancer. In one embodiment, the patient is injected multiple times (preferably about 2 up to about 50 times) with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac, for example, at 12-72 hour intervals or at 48-72 hour intervals. Additional injections of a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac can be administered to the patient at an interval of days or months after the initial injection(s) and the additional injections can prevent recurrence of the cancer. In another illustrative embodiment the patient is treated with single injections of Compound I-Lu or Ia-Lu and Compound I-Ac or Ia-Ac on the same day, in any order of injection, followed by multiple injections of I-Lu or Ia-Lu to treat the cancer. In some embodiments, the patient is injected multiple times (preferably about 2 up to about 50 times) with Compound I-Lu or Ia-Lu, after receiving an initial injection of each of Compound I-Lu or Ia-Lu and Compound I-Ac or Ia-Ac on the same day, in any order, or at the same time, for example, at 12-72 hour intervals, or at 48-72 hour intervals, or once weekly, or once every two weeks.

Any suitable course of therapy with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac can be used. In one illustrative embodiment, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac is administered in a single daily dose administered five days a week, in weeks 1, 2, and 3 of each 4 week cycle, with no dose administered in week 4. In an alternative

embodiment, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac is administered in a single daily dose administered three days a week, of weeks 1, and 3 of each 4 week cycle, with no dose administered in weeks 2 and 4. In an alternative embodiment, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac is administered biweekly on weeks 1 and 2, i.e. on days 1, 4, 8, 11, of a 3-week cycle. In an alternative embodiment, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac is administered once weekly on weeks 1 and 2, i.e. days 1 and 8 of a 3-week cycle. In an alternative embodiment, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac are administered on a single day, in any order, or at the same time, followed by administration of Compound I-Lu or Ia-Lu once weekly cycle for from about 2 to about 6-cycles. In an alternative embodiment, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac are administered on a single day, in any order, or at the same time, followed by administration of Compound I-Lu or Ia-Lu once weekly cycle for from about 2 to about 6-cycles, followed by administration of a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac on a single day, in any order, or at the same time, followed by administration of Compound I-Lu or Ia-Lu once weekly cycle for from about 2 to about 6-cycles.

In an alternative embodiment, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac are administered within about 2 to 72 hours of each other, in any order, followed by administration of Compound I-Lu or Ia-Lu once weekly cycle for from about 2 to about 6-cycles. In an alternative embodiment, a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac are administered within about 2 to 72 hours of each other, in any order, followed by administration of Compound I-Lu or Ia-Lu once weekly cycle for from about 2 to about 6-cycles, followed by administration of a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac on a single day, in any order, or at the same time, followed by administration of Compound I-Lu or Ia-Lu once weekly cycle for from about 2 to about 6-cycles.

Dose levels of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac can be measured in GBq and MBq, respectively. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 13 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from about 4 GBq to about 11 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from about 5 GBq to about 10 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 9 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from about 6.5 GBq to about 8.5 GBq. In some

embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from about 7 GBq to about 8 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is about 7.4 GBq. In some embodiments, the total dose of I-Lu or Ia-Lu ranges from about 15 GBq to about 200 GBq. In some embodiments, the total dose of I-Lu or Ia-Lu ranges from about 25 GBq to about 185 GBq. In some embodiments, the total dose of I-Lu or Ia-Lu ranges from about 35 GBq to about 150 GBq. In some embodiments, the total dose of I-Lu or Ia-Lu ranges from about 40 GBq to about 100 GBq. In some embodiments, the total dose of I-Lu, or Ia-Lu is about 44 GBq. In some embodiments, the maximum duration of treatment of a subject is about 19 to 23 months.

10 In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from 2 GBq to 20 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from 2 GBq to 13 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from 4 GBq to 11 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from 5 GBq to 10 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from 6 GBq to 9 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from 6 GBq to 8 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from 6.5 GBq to 8.5 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is from 7 GBq to 8 GBq. In some embodiments, a therapeutically effective amount of I-Lu or Ia-Lu is 7.4 GBq. In some embodiments, the total dose of I-Lu or Ia-Lu ranges from 15 GBq to 200 GBq. In some embodiments, the total dose of I-Lu or Ia-Lu ranges from 25 GBq to 185 GBq. In some embodiments, the total dose of I-Lu or Ia-Lu ranges from 35 GBq to 150 GBq. In some embodiments, the total dose of I-Lu or Ia-Lu ranges from 40 GBq to 100 GBq. In some embodiments, the total dose of I-Lu, or Ia-Lu is 44 GBq. In some embodiments, the maximum duration of treatment of a subject is 19 to 23 months.

25 In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 20 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from about 4 MBq to about 14 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from about 5 MBq to about 10 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from about 6 MBq to about 8 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from about 5 MBq to about 7 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 4 MBq. In some embodiments, a therapeutically

effective amount of I-Ac or Ia-Ac is from about 2 MBq to about 3 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is about 2.5 MBq.

In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from 1
5 MBq to 20 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from 1 MBq to 10 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from 4 MBq to 14 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from 5 MBq to 10 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from 6 MBq to 8 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from 5 MBq to 7 MBq. In some embodiments, a
10 therapeutically effective amount of I-Ac or Ia-Ac is from 1 MBq to 4 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is from 2 MBq to 3 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is 5 MBq. In some embodiments, a therapeutically effective amount of I-Ac or Ia-Ac is 2.5 MBq.

15 The PSMA ligand-imaging conjugates and Compounds I-Lu, I-Ac, Ia-Lu, and Ia-Ac described herein may contain one or more chiral centers, or may otherwise be capable of existing as multiple stereoisomers. Accordingly, it is to be understood that the present disclosure includes pure stereoisomers as well as mixtures of stereoisomers, such as enantiomers, diastereomers, and enantiomerically or diastereomerically enriched mixtures. The
20 PSMA ligand-imaging conjugates and Compounds I-Lu, I-Ac, Ia-Lu, and Ia-Ac described herein may be capable of existing as geometric isomers. Accordingly, it is to be understood that the present disclosure includes pure geometric isomers or mixtures of geometric isomers.

It is appreciated that the PSMA ligand-imaging conjugates and Compounds I-Lu, I-Ac, and Ia-Lu, Ia-Ac described herein may exist in unsolvated forms as well as solvated forms,
25 including hydrated forms. In general, the solvated forms are equivalent to unsolvated forms and are encompassed within the scope of the present disclosure. The PSMA ligand-imaging conjugates and Compounds I-Lu, I-Ac, Ia-Lu, and Ia-Ac described herein may exist in multiple crystalline or amorphous forms. In general, all physical forms are equivalent for the uses contemplated by the present disclosure and are intended to be within the scope of the present
30 disclosure.

In another embodiment, compositions and/or dosage forms for administration of a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac are prepared from Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac with a purity of at least about 90%, or about 95%, or about 96%, or about 97%, or about 98%, or about 99%, or about 99.5%. In another embodiment,

compositions and or dosage forms for administration of a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac are prepared from Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac with a purity of at least 90%, or at least 95%, or at least 96%, or at least 97%, or at least 98%, or at least 99%, or at least 99.5%.

5 In another embodiment, compositions and/or dosage forms for administration of the PSMA ligand-imaging conjugate are prepared from the PSMA ligand-imaging conjugate with a purity of at least about 90%, or about 95%, or about 96%, or about 97%, or about 98%, or about 99%, or about 99.5%. In another embodiment, compositions and or dosage forms for administration of the PSMA ligand-imaging conjugate are prepared from the PSMA ligand-
10 imaging conjugate with a purity of at least 90%, or at least 95%, or at least 97%, or at least 98%, or at least 99%, or at least 99.5%.

 In another embodiment, compositions and/or dosage forms for administration of radiolabeled PSMA ligand-imaging conjugate are prepared from the PSMA ligand-imaging conjugate with a radiochemical purity of at least about 90%, or about 95%, or about 96%, or
15 about 97%, or about 98%, or about 99%, or about 99.5%. In another embodiment, compositions and or dosage forms for administration of the PSMA ligand-imaging conjugate are prepared from the PSMA ligand-imaging conjugate with a purity of at least 90%, or at least 95%, or at least 96%, or at least 97%, or at least 98%, or at least 99%, or at least 99.5%.

 The purity of Compounds I-Lu, I-Ac, Ia-Lu, and Ia-Ac or the PSMA ligand-imaging
20 conjugates described herein may be measured using any conventional technique, including various chromatography or spectroscopic techniques, such as high pressure or high performance liquid chromatography (HPLC), nuclear magnetic resonance spectroscopy, TLC, UV absorbance spectroscopy, fluorescence spectroscopy, and the like.

 In another embodiment, Compounds I-Lu, I-Ac, Ia-Lu, and Ia-Ac or PSMA ligand-
25 imaging conjugate described herein is provided in a sterile container or package.

 In one aspect, a clinical benefit of the patient to treatment with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac can be characterized as overall survival (OS). As used herein, the term “overall survival (OS)” means the time from the date of randomization to the date of death from any cause.

30 In one aspect, a clinical benefit of the patient to treatment with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac can be characterized utilizing Response Evaluation Criteria in Solid Tumors (RECIST) criteria. Illustratively, the criteria have been adapted from the original *WHO Handbook (3)*, taking into account the measurement of the longest diameter for all target lesions: complete response, (CR) — the disappearance of all

target lesions; partial response (PR) — at least a 30% decrease in the sum of the longest diameter of target lesions, taking as reference the baseline sum longest diameter; stable disease (SD) — neither sufficient shrinkage to qualify for partial response nor sufficient increase to qualify for progressive disease, taking as reference the smallest sum longest diameter since the treatment started; progressive disease (PD) — at least a 20% increase in the sum of the longest diameter of target lesions, taking as reference the smallest sum longest diameter recorded since the treatment started or the appearance of one or more new lesions. In another aspect overall disease response rate (ORR) is a clinical benefit and is calculated as the percent of patients who achieve a best response of CR or PR. Overall disease control rate (DCR) can be another clinical benefit and is calculated as the percent of patients who achieve a best response of CR, PR, or SD. In some embodiments, the response can be disease control rate (DCR) as measured by RECIST v1.1 criteria.

In another aspect, a clinical benefit of the patient to treatment with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac can be characterized as radiographic progression-free survival (rPFS). As used herein, “radiographic progression-free survival (rPFS)” means the time from the date of randomization to the date of radiographic disease progression as outlined in Prostate Cancer Working Group 3 (PCWG3) Guidelines or death from any cause. See, for example, Scher HI, Morris MJ, Stadler WM, Higano C, Basch E, Fizazi K, et al. Trial Design and Objectives for Castration-Resistant Prostate Cancer: Updated Recommendations from the Prostate Cancer Clinical Trials Work Group 3. *J Clin Oncol* 2016;34(12):1402–18. In another aspect, a clinical benefit of the patient to treatment with I-Lu or Ia-Lu, and I-Ac or Ia-Ac can be characterized as time to a first symptomatic skeletal event (SSE). It will be appreciated that symptomatic skeletal event means a clinically significant pathological fracture, surgery or radiation to bone, or spinal cord compression. As used herein, “time to a first symptomatic skeletal event” means date of randomization to the date of first new symptomatic pathological bone fracture, spinal cord compression, tumor-related orthopedic surgical intervention, or requirement for radiation therapy to relieve bone pain, whichever occurs first.

In one illustrative example overall survival is the time to death for a given patient defined as the number of days from the first day the patient received protocol treatment (C1D1) to the date of the patient’s death. All events of death can be included, regardless of whether the event occurred while the patient was still taking the study drug or after the patient discontinued the study drug. If a patient has not died, then the data can be censored at the last study visit, or the last contact date, or the date the patient was last known to be alive, whichever is last.

Alternatively, a clinical benefit of the patient as a result of treatment with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac can be characterized as inhibition of tumor growth which can be identified in a patient through, for example, follow-up imaging of the patient's cancer after treatment with I-Lu or Ia-Lu, and I-Ac or Ia-Ac. For example, inhibition of tumor growth can be characterized by measuring the size of tumors in a patient after
5 administration of I-Lu or Ia-Lu, and I-Ac or Ia-Ac according to any of the imaging techniques described herein, where the inhibition of tumor growth is indicated by a stable tumor size, or by a reduction in tumor size. It will be appreciated that the identification of inhibition of tumor growth can be accomplished using a variety of techniques, and is not limited to the imaging
10 methods described herein (e.g CT, MRI, PET imaging, SPECT imaging or chest x-ray).

In one embodiment, a method is provided of determining whether a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac is indicated for the treatment of a patient with cancer, the method comprising the step of determining the PSMA status in a patient with cancer wherein a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac is indicated for the
15 treatment of the patient if the PSMA status of the patient is positive.

In one embodiment, a method is provided of assessing whether a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac are indicated for the treatment of a patient with one of the cancers described herein. The method comprises the steps of visually determining PSMA status in the patient wherein PSMA status is based on a imaging tumors that are PSMA
20 positive in the patient, and wherein the a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac are indicated for the treatment of the patient when the PSMA status of the patient is positive.

In the above-described embodiments, if a patient is in the group with positive PSMA status, a clinical benefit of treatment with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac is indicated. In one embodiment, the clinical benefit to the patient can be overall
25 survival of the patient, ability to receive four or more cycles of therapy with a combination of Compounds I-Lu or Ia-Lu, and I-Ac or Ia-Ac, inhibition of tumor growth, stable disease, a partial response of the patient to therapy, a complete response of the patient to therapy, disease control (i.e., the best result obtained is a complete response, a partial response, or stable
30 disease), and/or overall disease response (i.e., the best result obtained is a complete response or a partial response). In one illustrative example, the clinical benefit for a patient being treated for pleural mesothelioma or adenocarcinoma (e.g. adenocarcinoma of the gastroesophageal junction) is stable disease.

In another embodiment, the methods described herein include the following examples. The examples further illustrate additional features of the various embodiments of the present disclosure. However, it is to be understood that the examples are illustrative and are not to be construed as limiting other embodiments of the present disclosure. In addition, it is appreciated
5 that other variations of the examples are included in the various embodiments of the present disclosure. In addition, it will be appreciated that all ranges described herein, such as those described in connection with the various embodiments, are exemplary and not intended to be limiting. One of skill in the art will appreciate that all ranges described by an lower and upper
10 bound, such as about 1 to about 20, includes all possible values contained in the lower and upper bound, and includes all possible ranges of values available by the set of possible values contained in the lower and upper bound.

EXAMPLES

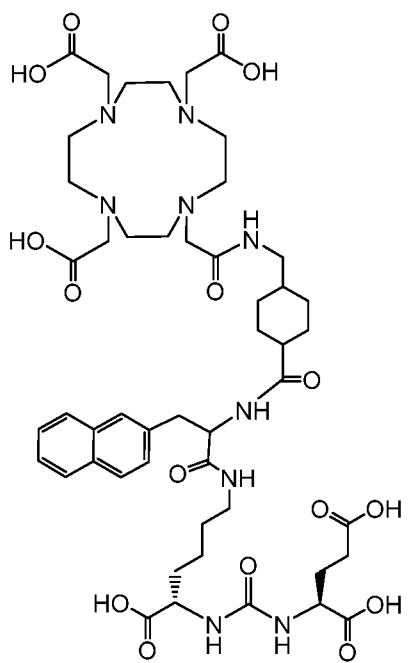
Example 1:

15 Patients with PSMA positive scans will be administered a single dose of Compound Ia-Ac on day 1, cycle 1 of the clinical regimen of 7.4GBq Compound Ia-Lu, administered 6 weekly for a maximum of 5 cycles. Subjects will be reviewed weekly for assessment of adverse events (onset, duration, grade and relatedness to treatment) during cycle 1 only. DLT will be determined by AE on cycle 1 only.

20 Subjects will be restaged at the end of every 2 cycles. At the time of each restaging, patients will be assessed by PSMA PET and Fluorinated PET/CT bone scan. Assessment of bone disease consistent with PCWG23 criteria. PSA evaluation will be measured according to institutional practice at a minimum of 2 weekly.

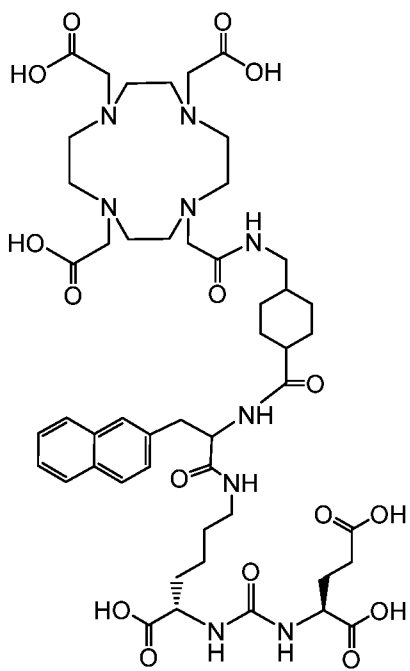
WHAT IS CLAIMED IS:

1. A method for treating cancer in a host animal, the method comprising the step of administering to the host animal a therapeutically effective amount of a first compound
- 5 having the Formula I



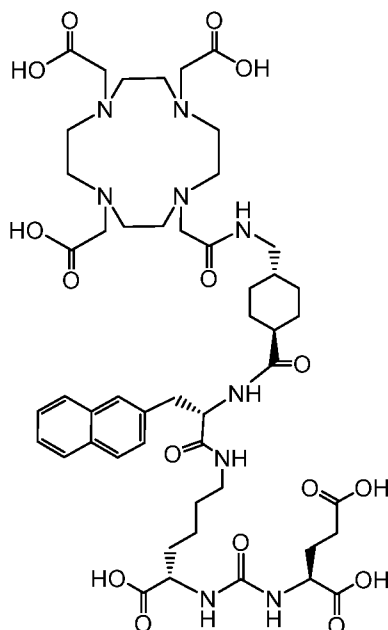
wherein the compound is complexed with ^{177}Lu (I-Lu);

in combination with a therapeutically effective amount of a second compound having the Formula I



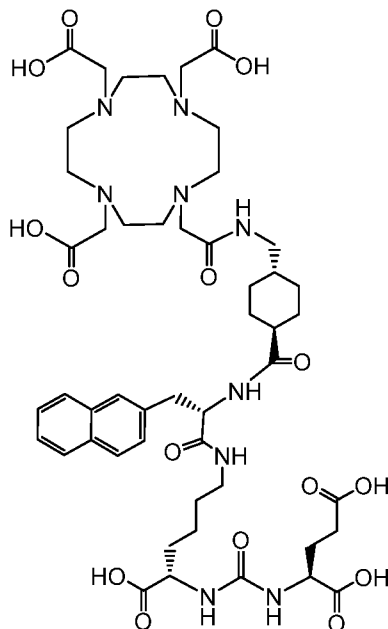
wherein the compound is complexed with ^{225}Ac (I-Ac).

2. The method of claim 1, wherein the first compound is of the Formula Ia



wherein the compound is complexed with ^{177}Lu (Ia-Lu).

- 5 3. The method of claim 2, wherein the second compound is of the Formula Ia



wherein the compound is complexed with ^{225}Ac (Ia-Ac).

- 10 4. The method of claims 1 to 3, wherein the cancer is associated with expression of prostate specific membrane antigen (PSMA).

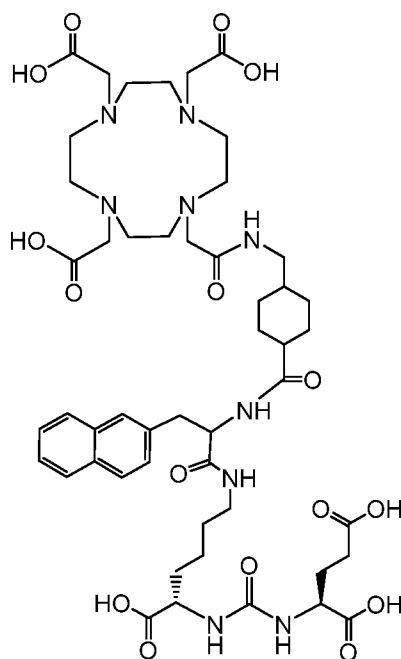
5. The method of claims 1 to 3, wherein the cancer is selected from the group consisting of prostate cancer, metastatic castration-resistant prostate cancer (mCRPC), thyroid cancer, renal cell carcinoma, transitional cell carcinoma of the bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, soft tissue sarcoma, and breast carcinoma.
6. The method of claim 5, wherein the cancer is prostate cancer.
7. The method of claim 5, wherein the cancer is metastatic castration-resistant prostate cancer (mCRPC).
8. The method of any one of claims 1-3, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq.
9. The method of any one of claims 1-3, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq.
10. The method of any one of claims 1-3, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 7.4 GBq.
11. The method of any one of claims 1-3, wherein the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq; or about 5 MBq to about 10 MBq; or about 5 MBq to about 7MBq.
12. The method of any one of claims 1-3, wherein the therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq.
13. The method of any of claims 1-3, wherein the compound of Formula I-Lu or Ia-Lu is administered at the same time as the compound of Formula I-Ac or Ia-Ac.
14. The method of any one of claims 1-3, wherein the compound of Formula I-Lu or Ia-Lu is administered about 1 hour prior to the compound of Formula I-Ac or Ia-Ac; or about 12 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 24 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 48 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 72 hours prior to the compound of Formula I-Ac or Ia-Ac.
15. The method of any one of claims 1-3, wherein the compound of Formula I-Ac or Ia-Ac is administered about 1 hour prior to the compound of Formula I-Lu or Ia-Lu; or about 12 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 24 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 48 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 72 hours prior to the compound of Formula I-Lu or Ia-Lu.
16. The method of claim 13, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly

cycle for from about 1 to about 7 cycles following the administration of I-Lu or Ia-Lu and I-Ac or Ia-Ac.

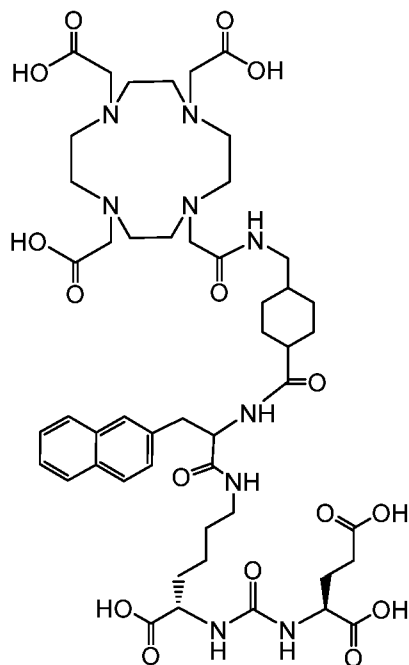
17. The method of claim 14, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

18. The method of claim 15, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

19. A compound of the Formula I-Lu

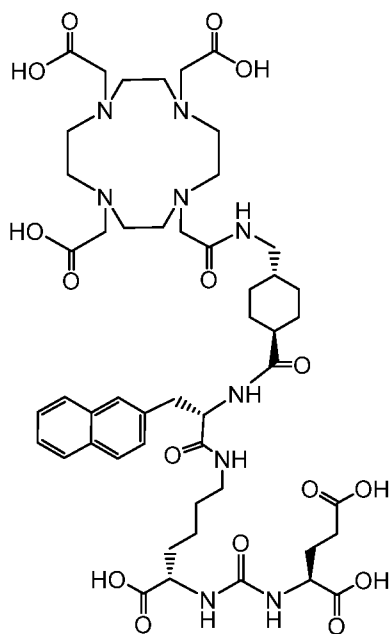


wherein the compound is complexed with ^{177}Lu , for use in the treatment of cancer in a patient, in combination with a therapeutically effective amount of a compound of the Formula I-Ac



wherein the compound is complexed with ^{225}Ac .

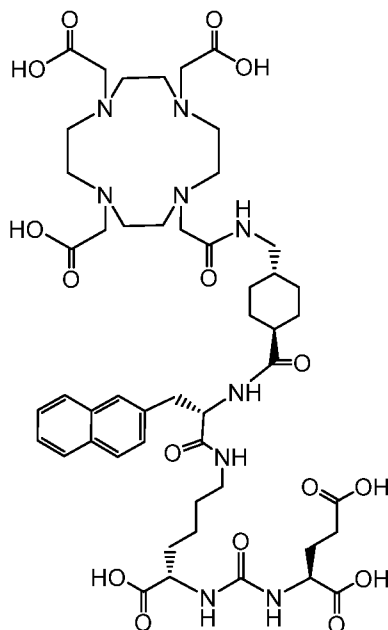
20. The compound of claim 19, wherein the compound is of the Formula Ia-Lu



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wherein the compound is complexed with ^{177}Lu .

21. The compound of claim 19, wherein the compound I-Ac is of the Formula Ia-Ac



wherein the compound is complexed with ^{225}Ac .

22. The compound of any one of claims 19 to 21, wherein the cancer is associated with expression of prostate specific membrane antigen (PSMA).

5 23. The compound of any one of claims 19 to 21, wherein the cancer is selected from the group consisting of prostate cancer, metastatic castration-resistant prostate cancer (mCRPC), thyroid cancer, renal cell carcinoma, transitional cell carcinoma of the bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, soft tissue
10 sarcoma, and breast carcinoma.

24. The compound of any one of claims 19 to 21, wherein the cancer is prostate cancer.

25. The compound of any one of claims 19 to 21, wherein the cancer is metastatic castration-resistant prostate cancer (mCRPC).

15 26. The compound of any one of claims 19 to 21, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq.

27. The compound of any one of claims 19 to 21, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq.

20 28. The compound of any one of claims 19 to 21, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 7.4 GBq.

29. The compound of any one of claims 19 to 21, wherein the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq; or about 5 MBq to about 10 MBq; or about 5 MBq to about 7MBq.

30. The compound of any one of claims 19 to 21, wherein the therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq.

31. The compound of any one of claims 19 to 21, wherein the compound of Formula I-Lu or Ia-Lu is administered at the same time as the compound of Formula I-Ac or Ia-Ac.
5

32. The compound of any one of claims 19 to 21, wherein the compound of Formula I-Lu or Ia-Lu is administered about 1 hour prior to the compound of Formula I-Ac or Ia-Ac; or about 12 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 24 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 48 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 72 hours prior to the compound of Formula I-Ac or Ia-Ac.
10

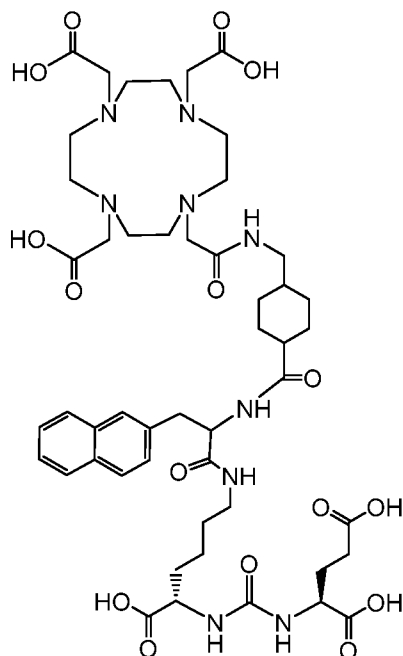
33. The compound of any one of claims 19 to 21, wherein the compound of Formula I-Ac or Ia-Ac is administered about 1 hour prior to the compound of Formula I-Lu or Ia-Lu; or about 12 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 24 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 48 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 72 hours prior to the compound of Formula I-Lu or Ia-Lu.
15

34. The compound of claim 31, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of I-Lu or Ia-Lu and I-Ac or Ia-Ac.
20

35. The compound of claim 32, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

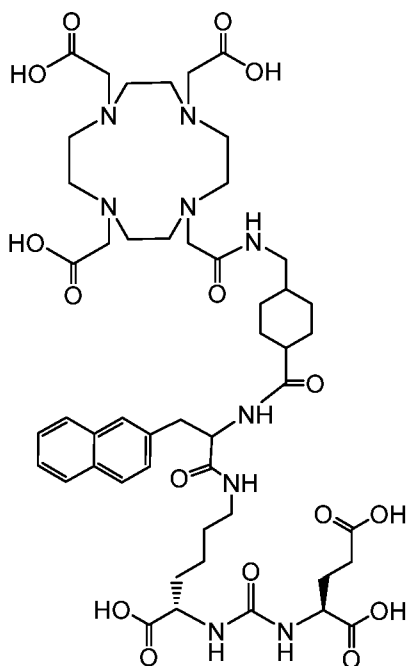
36. The compound of claim 33, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.
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37. Use of a compound of the Formula I-Lu,



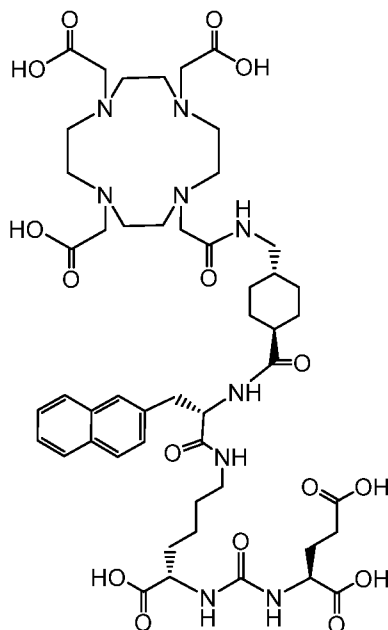
wherein the compound is complexed with ^{177}Lu , in the preparation of a medicament comprising a therapeutically effective amount of the compound of the Formula I-Lu, for treating cancer in a patient in combination with a therapeutically effective amount of a compound of the Formula I-

5 Ac



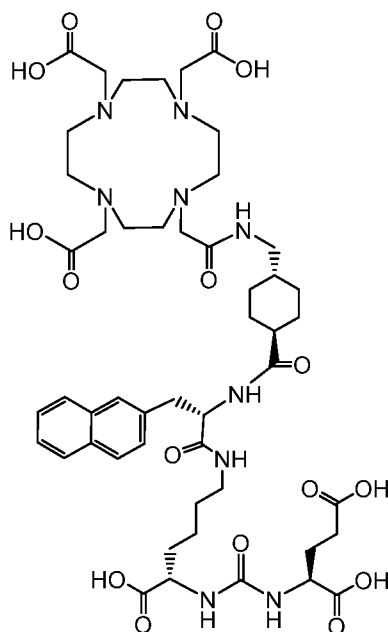
wherein the compound is complexed with ^{225}Ac .

38. The use of claim 37, wherein the compound of the Formula I-Lu is of the Formula Ia-Lu



wherein the compound is complexed with ^{177}Lu .

39. The use of claim 37, wherein the compound of the Formula I-Ac is of the Formula Ia-Ac



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wherein the compound is complexed with ^{225}Ac .

40. The use of any one of claims 37 to 39, wherein the cancer is associated with expression of prostate specific membrane antigen (PSMA).

41. The use of any one of claims 37 to 39, wherein the cancer is selected from the group consisting of prostate cancer, metastatic castration-resistant prostate cancer (mCRPC), thyroid cancer, renal cell carcinoma, transitional cell carcinoma of the bladder,

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colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, soft tissue sarcoma, and breast carcinoma.

5 42. The use of any one of claims 37 to 39, wherein the cancer is prostate cancer.

43. The use of any one of claims 37 to 39, wherein the cancer is metastatic castration-resistant prostate cancer (mCRPC).

44. The use of any one of claims 37 to 39, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq.

10 45. The use of any one of claims 37 to 39, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq.

46. The use of any one of claims 37 to 39, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 7.4 GBq.

15 47. The use of any one of claims 37 to 39, wherein the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq; or about 5 MBq to about 10 MBq; or about 5 MBq to about 7MBq.

48. The use of any one of claims 37 to 39, wherein the therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq.

20 49. The use of any one of claims 37 to 39, wherein the compound of Formula I-Lu or Ia-Lu is administered at the same time as the compound of Formula I-Ac or Ia-Ac.

25 50. The use of any one of claims 37 to 39, wherein the compound of Formula I-Lu or Ia-Lu is administered about 1 hour prior to the compound of Formula I-Ac or Ia-Ac; or about 12 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 24 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 48 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 72 hours prior to the compound of Formula I-Ac or Ia-Ac.

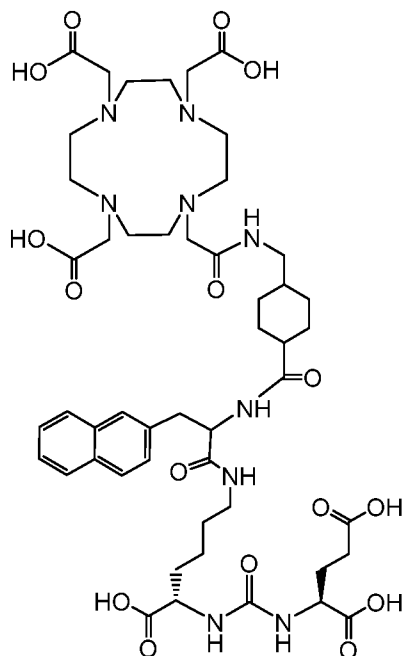
30 51. The use of any one of claims 37 to 39, wherein the compound of Formula I-Ac or Ia-Ac is administered about 1 hour prior to the compound of Formula I-Lu or Ia-Lu; or about 12 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 24 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 48 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 72 hours prior to the compound of Formula I-Lu or Ia-Lu.

52. The use of claim 49, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of I-Lu or Ia-Lu and I-Ac or Ia-Ac.

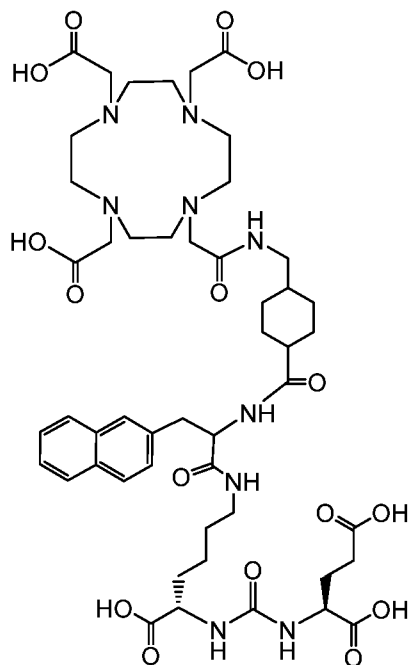
53. The use of claim 50, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

54. The use of claim 51, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

55. A composition comprising a compound of the Formula I-Lu,

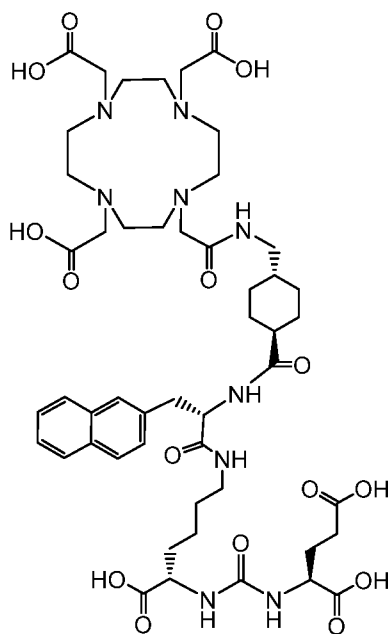


10 wherein the compound is complexed with ^{177}Lu , in a therapeutically effective amount, for use in the treatment of cancer in a patient, in combination with a therapeutically effective amount of a compound of the Formula I-Ac



wherein the compound is complexed with ^{225}Ac .

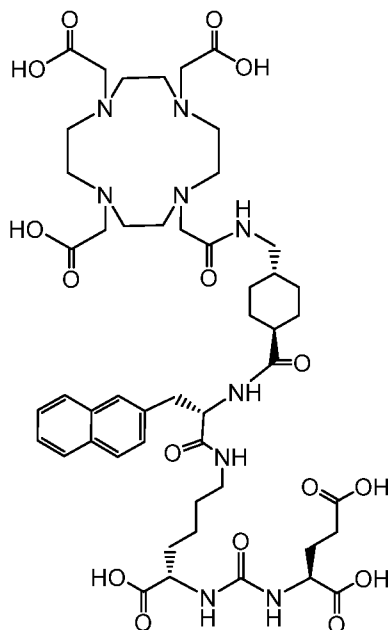
56. The composition of claim 55, wherein the compound of the Formula I-Lu is of the Formula Ia-Lu



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wherein the compound is complexed with ^{177}Lu .

57. The composition of claim 55, wherein the compound of the Formula I-Ac is of the Formula Ia-Ac



wherein the compound is complexed with ^{225}Ac .

58. The composition of any one of claims 55 to 57, wherein the cancer is associated with expression of prostate specific membrane antigen (PSMA).

5 59. The composition of any one of claims 55 to 57, wherein the cancer is selected from the group consisting of prostate cancer, metastatic castration-resistant prostate cancer (mCRPC), thyroid cancer, renal cell carcinoma, transitional cell carcinoma of the bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, soft tissue
10 sarcoma, and breast carcinoma.

60. The composition of any one of claims 55 to 57, wherein the cancer is prostate cancer.

61. The composition of any one of claims 55 to 57, wherein the cancer is metastatic castration-resistant prostate cancer (mCRPC).

15 62. The composition of any one of claims 58 to 61, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq.

63. The composition of any one of claims 58 to 61, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq.

20 64. The composition of any one of claims 58 to 61, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 7.4 GBq.

65. The composition of any one of claims 58 to 61, wherein the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq; or about 5 MBq to about 10 MBq; or about 5 MBq to about 7MBq.

66. The composition of any one of claims 58 to 61, wherein the therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq.

67. The composition of any one of claims 58 to 61, wherein the compound of Formula I-Lu or Ia-Lu is administered at the same time as the compound of Formula I-Ac or Ia-Ac.

68. The composition of any one of claims 58 to 61, wherein the compound of Formula I-Lu or Ia-Lu is administered about 1 hour prior to the compound of Formula I-Ac or Ia-Ac; or about 12 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 24 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 48 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 72 hours prior to the compound of Formula I-Ac or Ia-Ac.

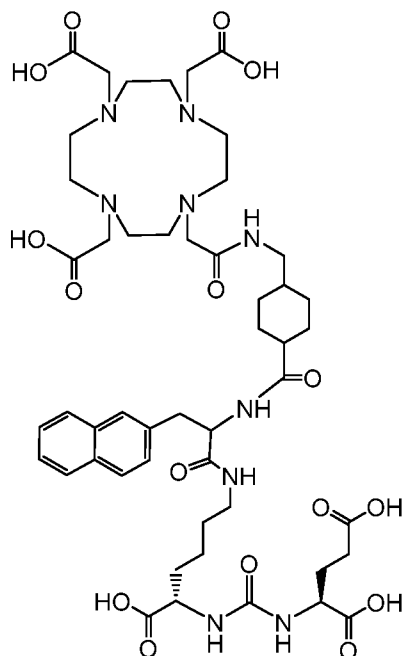
69. The composition of any one of claims 58 to 61, wherein the compound of Formula I-Ac or Ia-Ac is administered about 1 hour prior to the compound of Formula I-Lu or Ia-Lu; or about 12 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 24 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 48 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 72 hours prior to the compound of Formula I-Lu or Ia-Lu.

70. The composition of claim 67, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of I-Lu or Ia-Lu and I-Ac or Ia-Ac.

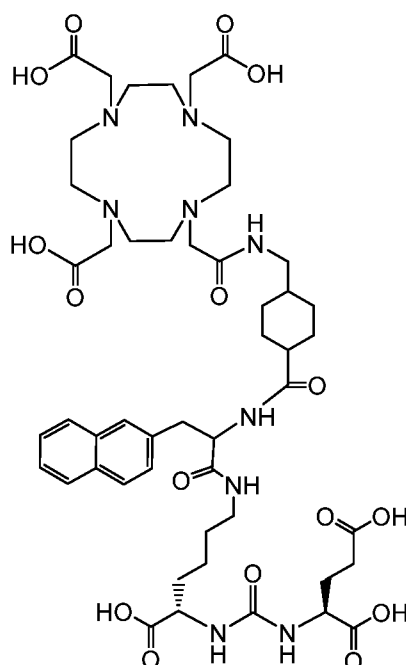
71. The composition of claim 68, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

72. The composition of claim 69, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

73. A medicament comprising a compound of the Formula I-Lu

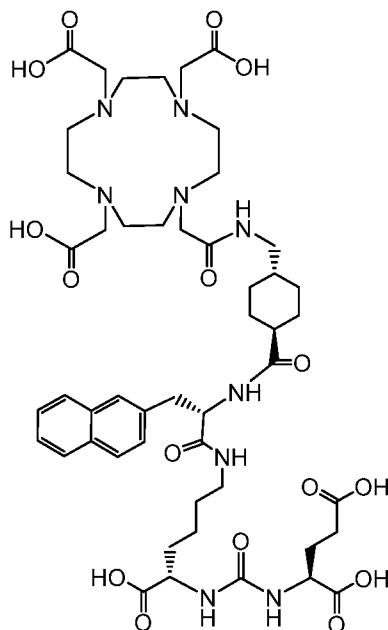


wherein the compound is complexed with ^{177}Lu , in a therapeutically effective amount, combined with a therapeutically effective amount of a compound of the Formula I-Ac



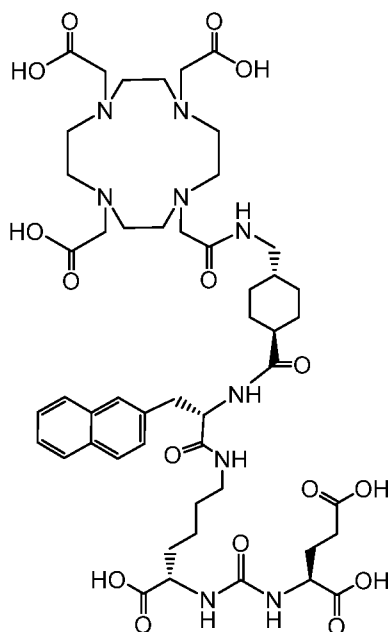
5 wherein the compound is complexed with ^{225}Ac .

74. The medicament of claim 73, wherein the compound of the Formula I-Lu is of the Formula Ia-Lu



wherein the compound is complexed with ^{177}Lu .

75. The medicament of claim 73, wherein the compound of the Formula I-Ac is of the Formula Ia-Ac



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wherein the compound is complexed with ^{225}Ac .

76. The medicament of any one of claims 73 to 75, wherein medicament provides a synergistic effect on a cancer associated with expression of prostate specific membrane antigen (PSMA).

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77. The medicament of any one of claims 73 to 75, wherein the cancer is selected from the group consisting of prostate cancer, metastatic castration-resistant prostate

cancer (mCRPC), thyroid cancer, renal cell carcinoma, transitional cell carcinoma of the bladder, colonic adenocarcinoma, neuroendocrine carcinoma, glioblastoma multiforme, malignant melanoma, pancreatic duct carcinoma, non-small cell lung carcinoma, soft tissue sarcoma, and breast carcinoma.

5 78. The medicament of any one of claims 73 to 75, wherein the cancer is prostate cancer.

 79. The medicament of any one of claims 73 to 75, wherein the cancer is metastatic castration-resistant prostate cancer (mCRPC).

10 80. The medicament of any one of claims 73 to 75, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 2 GBq to about 20 GBq.

 81. The medicament of any one of claims 73 to 75, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 6 GBq to about 8 GBq.

 82. The medicament of any one of claims 73 to 75, wherein the therapeutically effective amount of I-Lu or Ia-Lu is from about 7.4 GBq.

15 83. The medicament of any one of claims 73 to 75, wherein the therapeutically effective amount of I-Ac or Ia-Ac is from about 1 MBq to about 10 MBq; or about 5 MBq to about 10 MBq; or about 5 MBq to about 7MBq.

 84. The medicament of any one of claims 73 to 75, wherein the therapeutically effective amount of I-Ac or Ia-Ac is about 5 MBq.

20 85. The medicament of any one of claims 73 to 75, wherein the compound of Formula I-Lu or Ia-Lu is administered at the same time as the compound of Formula I-Ac or Ia-Ac.

 86. The medicament of any one of claims 73 to 75, wherein the compound of Formula I-Lu or Ia-Lu is administered about 1 hour prior to the compound of Formula I-Ac or Ia-Ac; or about 12 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 24 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 48 hours prior to the compound of Formula I-Ac or Ia-Ac; or about 72 hours prior to the compound of Formula I-Ac or Ia-Ac.

30 87. The medicament of any one of claims 73 to 75, wherein the compound of Formula I-Ac or Ia-Ac is administered about 1 hour prior to the compound of Formula I-Lu or Ia-Lu; or about 12 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 24 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 48 hours prior to the compound of Formula I-Lu or Ia-Lu; or about 72 hours prior to the compound of Formula I-Lu or Ia-Lu.

 88. The medicament of claim 87, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly

cycle for from about 1 to about 7 cycles following the administration of I-Lu or Ia-Lu and I-Ac or Ia-Ac.

89. The medicament of claim 88, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly
5 cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.

90. The medicament of claim 89, further comprising administering a therapeutically effective amount of the compound of Formula I-Lu or Ia-Lu on a once weekly
10 cycle for from about 1 to about 7 cycles following the administration of both I-Lu or Ia-Lu and I-Ac or Ia-Ac.