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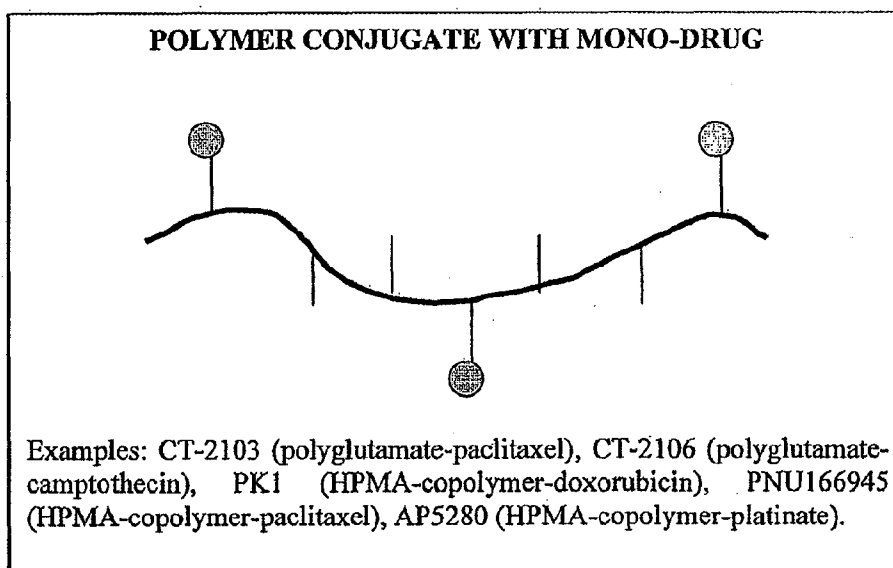
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FIGURE 1



(57) Abstract: Various biodegradable polyglutamate conjugates comprising recurring units of the general formulae (I), (II), (III), (IV), (V), and/or (VI) are prepared. The polymers are conjugated with a plurality of drugs. Such polymer conjugates are useful for variety of drug, targeting, stabilizing and/or imaging agent delivery applications.

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## **POLYGLUTAMATE CONJUGATES AND POLYGLUTAMATE-AMINO ACID CONJUGATES HAVING A PLURALITY OF DRUGS**

**[0001]** This application claims priority to U.S. Provisional Application No. 60/916,865, entitled "POLYGLUTAMATE CONJUGATES AND POLYGLUTAMATE-AMINO ACID CONJUGATES HAVING A PLURALITY OF DRUGS," filed on May 9, 2007; which is incorporated herein by reference in its entirety.

### **BACKGROUND OF THE INVENTION**

#### Field of the Invention

**[0002]** Generally disclosed herein are biocompatible polymers having a plurality of drugs conjugated thereto. The polymer conjugates described herein are useful for a variety of drug, biomolecule, and imaging agent delivery applications. Also disclosed are methods of using the polymer conjugates to treat, diagnose, and/or image a subject.

#### Description of the Related Art

**[0003]** A variety of systems have been used for the delivery of drugs, biomolecules, and imaging agents. For example, such systems include capsules, liposomes, microparticles, nanoparticles, and polymers.

**[0004]** A variety of polyester-based biodegradable systems have been characterized and studied. Polylactic acid (PLA), polyglycolic acid and their copolymers polylactic-co-glycolic acid (PLGA) are some examples of well-characterized biomaterials with regard to design and performance for drug-delivery applications. See Uhrich, K.E.; Cannizzaro, S.M.; Langer, R.S. and Shakeshelf, K.M. "Polymeric Systems for Controlled Drug Release," Chem. Rev. 1999, 99, 3181-3198 and Panyam J, Labhasetwar V. "Biodegradable nanoparticles for drug and gene delivery to cells and tissue," Adv. Drug. Deliv. Rev. 2003, 55, 329-47. Also, 2-hydroxypropyl methacrylate (HPMA) has been widely used to create a polymer for drug-delivery applications. Biodegradable systems based on polyorthoesters have also been investigated. See Heller, J.; Barr, J.; Ng, S.Y.; Abdellauoi, K.S. and Gurny, R. "Poly(ortho esters): synthesis, characterization, properties and uses." Adv. Drug Del. Rev. 2002, 54, 1015-1039. Polyanhydride systems have also been

investigated. Such polyanhydrides are typically biocompatible and may degrade in vivo into relatively non-toxic compounds that are eliminated from the body as metabolites. See Kumar, N.; Langer, R.S. and Domb, A.J. "Polyanhydrides: an overview," *Adv. Drug Del. Rev.* 2002, 54, 889-91.

**[0005]** Amino acid-based polymers have also been considered as a potential source of new biomaterials. Poly-amino acids having good biocompatibility have been investigated to deliver low molecular-weight compounds. A relatively small number of polyglutamic acids and copolymers have been identified as candidate materials for drug delivery. See Bourke, S.L. and Kohn, J. "Polymers derived from the amino acid L-tyrosine: polycarbonates, polyarylates and copolymers with poly(ethylene glycol)." *Adv. Drug Del. Rev.*, 2003, 55, 447- 466.

**[0006]** Administered hydrophobic anticancer drugs, therapeutic proteins, and polypeptides often suffer from poor bio-availability. Such poor bio-availability may be due to incompatibility of bi-phasic solutions of hydrophobic drugs and aqueous solutions and/or rapid removal of these molecules from blood circulation by enzymatic degradation. One technique for increasing the efficacy of administered proteins and other small molecule agents entails conjugating the administered agent with a polymer, such as a polyethylene glycol ("PEG") molecule, that can provide protection from enzymatic degradation in vivo. Such "PEGylation" often improves the circulation time and, hence, bio-availability of an administered agent.

**[0007]** PEG has shortcomings in certain respects, however. For example, because PEG is a linear polymer, the steric protection afforded by PEG is limited, as compared to branched polymers. Another shortcoming of PEG is that it is generally amenable to derivatization at its two terminals. This limits the number of other functional molecules (e.g. those helpful for protein or drug delivery to specific tissues) that can be conjugated to PEG.

**[0008]** Polyglutamic acid (PGA) is another polymer of choice for solubilizing hydrophobic anticancer drugs. Many anti-cancer drugs conjugated to PGA have been reported. See Chun Li. "Poly(L-glutamic acid)-anticancer drug conjugates." *Adv. Drug Del. Rev.*, 2002, 54, 695-713. However, none are currently FDA-approved.

**[0009]** Paclitaxel, extracted from the bark of the Pacific Yew tree, is a FDA-approved drug for the treatment of ovarian cancer and breast cancer. Wani et al. "Plant

antitumor agents. VI. The isolation and structure of taxol, a novel antileukemic and antitumor agent from *Taxus brevifolia*,” J. Am. Chem. Soc. 1971, 93, 2325-7. However, like other anti-cancer drugs, paclitaxel suffers from poor bio-availability due to its hydrophobicity and insolubility in aqueous solution. One way to solubilize paclitaxel is to formulate it in a mixture of Cremophor-EL and dehydrated ethanol (1:1, v/v). Sparreboom et al. “Cremophor EL-mediated Alteration of Paclitaxel Distribution in Human Blood: Clinical Pharmacokinetic Implications,” Cancer Research, 1999, 59, 1454-1457. This formulation is currently commercialized as Taxol<sup>®</sup> (Bristol-Myers Squibb). Another method of solubilizing paclitaxel is by emulsification using high-shear homogenization. Constantinides et al. “Formulation Development and Antitumor Activity of a Filter-Sterilizable Emulsion of Paclitaxel,” Pharmaceutical Research 2000, 17, 175-182. Recently, polymer-paclitaxel conjugates have been advanced in several clinical trials. Ruth Duncan “The Dawning era of polymer therapeutics,” Nature Reviews Drug Discovery 2003, 2, 347-360. More recently, paclitaxel has been formulated into nano-particles with human albumin protein and has been used in clinical studies. Damascelli et al. “Intraarterial chemotherapy with polyoxyethylated castor oil free paclitaxel, incorporated in albumin nanoparticles (ABI-007): Phase II study of patients with squamous cell carcinoma of the head and neck and anal canal: preliminary evidence of clinical activity.” Cancer, 2001, 92, 2592-602, and Ibrahim et al. “Phase I and pharmacokinetic study of ABI-007, a Cremophor-free, protein-stabilized, nanoparticle formulation of paclitaxel,” Clin. Cancer Res. 2002, 8, 1038-44. This formulation is currently commercialized as Abraxane<sup>®</sup> (American Pharmaceutical Partners, Inc.).

**[0010]** Magnetic resonance imaging (MRI) is an important tool in diagnosis and staging of disease because it is non-invasive and non-irradiating. See Bulte et al. “Magnetic resonance microscopy and histology of the CNS,” Trends in Biotechnology, 2002, 20, S24-S28). Although images of tissues can be obtained, MRI with contrast agents significantly improves its resolution. However, paramagnetic metal ions suitable for MRI contrast agents are often toxic. One of the methods to reduce toxicity is to chelate these metal ions with polydentate molecules such as diethylenetriamine pentaacetate molecules (DTPA). Gd-DTPA was approved by FDA in 1988 for clinical uses, and it is currently commercialized as Magnevist<sup>®</sup>. Other Gd-chelates were approved by FDA and commercialized, and many

others are under development. See Caravan et al. "Gadolinium(III) Chelates as MRI Contrast Agents: Structure, Dynamics, and Applications," *Chem. Rev.* 1999, 99, 2293-2352.

**[0011]** However, Gd-DTPA is not ideal for targeting tumor tissues because it lacks specificity. When Gd-DTPA is administered via IV injection, it spontaneously and rapidly diffuses into extravascular space of the tissues. Thus, large amounts of contrast agents are usually required to produce reasonable contrast images. In addition, it is quickly eliminated via kidney filtration. To avoid the diffusion and the filtration, macromolecular MRI contrast agents have been developed. See Caravan et al. "Gadolinium(III) Chelates as MRI Contrast Agents: Structure, Dynamics, and Applications," *Chem. Rev.* 1999, 99, 2293-2352. These macromolecular-MRI contrast agents include protein-MRI chelates, polysaccharide-MRI chelates, and polymer-MRI chelates. See Lauffer et al. "Preparation and Water Relaxation Properties of Proteins Labeled with Paramagnetic Metal Chelates," *Magn. Reson. Imaging* 1985, 3, 11-16; Sirlin et al. "Gadolinium-DTPA-Dextran: A Macromolecular MR Blood Pool Contrast Agent," *Acad. Radiol.* 2004, 11, 1361-1369; Lu et al. "Poly(L-glutamic acid) Gd(III)-DOTA Conjugate with a Degradable Spacer for Magnetic Resonance Imaging," *Bioconjugate Chem.* 2003, 14, 715-719; and Wen et al. "Synthesis and Characterization of Poly(L-glutamic acid) Gadolinium Chelate: A New Biodegradable MRI Contrast Agent," *Bioconjugate Chem.* 2004, 15, 1408-1415.

**[0012]** Recently, tissue-specific MRI contrast agents have been developed. See Weinmann et al. "Tissue-specific MR contrast agents," *Eur. J. Radiol.* 2003, 46, 33-44. However, tumor-specific MRI contrast agents have not been reported in clinical applications. Nano-size particles have been reported to target tumor-tissues via an enhanced permeation and retention (EPR) effect. See Brannon-Peppas et al. "Nanoparticle and targeted systems for cancer therapy," *ADDR*, 2004, 56, 1649-1659).

#### SUMMARY OF THE INVENTION

**[0013]** Relatively hydrophobic imaging agents and drugs (such as certain hydrophobic anti-cancer drugs, therapeutic proteins and polypeptides) often suffer from poor bioavailability. It is believed that this problem is due at least in part to the poor solubility of these imaging agents and drugs in aqueous systems. Certain enzymatically degradable drugs

also suffer from poor bioavailability because they are degraded relatively rapidly in the circulatory system, resulting in rapid elimination from the body.

**[0014]** The inventors have discovered a series of novel polyglutamate conjugates and/or polyglutamate-amino acid conjugates that are capable of conjugating to a number of agents, such as imaging agents, targeting agents, stabilizing agents and/or drugs. In some embodiments, the polymers and the resulting conjugates preferentially accumulate in certain tissues (e.g., tumor tissues) and/or certain receptors, and thus are useful for delivering drugs (e.g., anticancer drugs) and/or imaging agents to specific parts of the body (e.g., tumors). In an embodiment, the polymer conjugate comprises a group that comprises a first drug and a group that comprises a second drug, wherein the first drug and the second drug are not the same. In an embodiment, the polymers and/or the resulting polymer conjugates form can nanoparticles that effectively solubilize the imaging agent, targeting agent, magnetic resonance imaging agent, and/or drugs in aqueous systems by dispersing it at a molecular level, thereby increasing functionality and/or bioavailability.

**[0015]** An embodiment described herein relates to a polymer conjugate that can include a recurring unit of the formula (I), a recurring unit of the formula (II), a recurring unit of the formula (III), a recurring unit of the formula (IV), a recurring unit of the formula (V), and/or a recurring unit of the formula (VI) as set forth herein, wherein: each  $A^1$ ,  $A^2$ ,  $A^3$ ,  $A^4$ ,  $A^5$  and  $A^6$  can be independently oxygen or  $NR^7$ , wherein  $R^7$  can be hydrogen or a  $C_{1-4}$  alkyl; wherein each  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  can be independently selected from hydrogen, a  $C_{1-10}$  alkyl group, a  $C_{6-20}$  aryl group, an ammonium group, an alkali metal, a polydentate ligand, a polydentate ligand precursor with protected oxygen atoms, a group that comprises a drug, a group that comprises a targeting agent, a group that comprises an optical imaging agent, a group that comprises a magnetic resonance imaging agent, and a group that comprises a stabilizing agent; m, n, and o can be each independently 1 or 2; p, q, r, s, t, and u are each independently 0 or  $\geq 1$ , wherein the sum of p, q, r, s, t, and u is 2 or greater; and provided that at least one of  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  is a group that comprises a first drug, at least one of  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  is a group that comprises a second drug, wherein the first drug and second drug are not the same.

**[0016]** Another embodiment described herein relates to a method of making a polymer conjugate as described herein that can include dissolving or partially dissolving a

polymeric reactant comprising at least one of a recurring unit of formula (VII) and/or a recurring unit of formula (VIII), as set forth herein, in a solvent to form a dissolved or partially dissolved polymeric reactant, wherein:  $z$  can be independently 1 or 2;  $A^7$  and each  $A^8$  can be oxygen; and  $R^{10}$  and each  $R^{11}$  can be each independently selected from hydrogen, ammonium, and an alkali metal; and reacting the dissolved or partially dissolved polymeric reactant with a second reactant and a third reactant, wherein the second reactant comprises the first drug and the third reactant comprises the second drug.

[0017] Still another embodiment described herein relates to a composition that can include the polymer conjugate described herein, and further comprising at least one selected from a pharmaceutically acceptable excipient, a carrier, and a diluent.

[0018] Yet still another embodiment described herein relates to a method of treating or ameliorating a disease or condition that can include administering an effective amount of the polymer conjugate described herein to a mammal in need thereof.

[0019] Some embodiments described herein relate to a method of diagnosing a disease or condition that can include administering an effective amount of the polymer conjugate described herein to a mammal in need thereof.

[0020] Another embodiment described herein relates to a method of imaging a portion of tissue that can include contacting a portion of tissue with an effective amount of the polymer conjugate described herein.

[0021] These and other embodiments are described in greater detail below.

#### BRIEF DESCRIPTION OF THE DRAWINGS

[0022] Figure 1 schematically illustrates a polymer conjugate that includes one type of drug.

[0023] Figure 2 schematically illustrates a polymer conjugate that includes a plurality of drugs.

[0024] Figure 3 illustrates a reaction scheme for the preparation of polymer conjugates that include a plurality of drugs.

[0025] Figure 4 illustrates another reaction scheme for the preparation of polymer conjugates that include a plurality of drugs.

[0026] Figure 5 illustrates a reaction scheme for the preparation of a polyglutamic acid conjugate with paclitaxel.

[0027] Figure 6 illustrates a reaction scheme for the preparation of a polyglutamic acid conjugate with paclitaxel and doxorubicin.

[0028] Figure 7 illustrates a reaction scheme for the preparation of a polyglutamic acid conjugate with paclitaxel and camptothecin.

[0029] Figure 8 illustrates a reaction scheme for the preparation of a polyglutamic acid conjugate with paclitaxel, doxorubicin, and camptothecin.

[0030] Figure 9 illustrates a reaction scheme for the preparation of a polyglutamic acid amino acid conjugate with paclitaxel.

[0031] Figure 10 illustrates a reaction scheme for the preparation of a polyglutamic acid amino acid conjugate with paclitaxel and doxorubicin.

[0032] Figure 11 illustrates a reaction scheme for the preparation of a polyglutamic acid amino acid conjugate with paclitaxel and camptothecin.

[0033] Figure 12 illustrates a reaction scheme for the preparation of a polyglutamic acid amino acid conjugate with paclitaxel, doxorubicin, and camptothecin.

#### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

[0034] Unless defined otherwise, all technical and scientific terms used herein have the same meaning as is commonly understood by one of ordinary skill in the art. All patents, applications, published applications and other publications referenced herein are incorporated by reference in their entirety unless stated otherwise. In the event that there are a plurality of definitions for a term herein, those in this section prevail unless stated otherwise.

[0035] The term “ester” is used herein in its ordinary sense, and thus includes a chemical moiety with formula  $-(R)_n-COOR'$ , where R and R' are independently selected from alkyl, cycloalkyl, aryl, heteroaryl (bonded through a ring carbon) and heteroalicyclic (bonded through a ring carbon), and where n is 0 or 1.

[0036] The term “amide” is used herein in its ordinary sense, and thus includes a chemical moiety with formula  $-(R)_n-C(O)NHR'$  or  $-(R)_n-NHC(O)R'$ , where R and R' are independently selected from alkyl, cycloalkyl, aryl, heteroaryl (bonded through a ring



carbon) and heteroalicyclic (bonded through a ring carbon), and where n is 0 or 1. An amide may be included in an amino acid or a peptide molecule attached to drug molecule as described herein, thereby forming a prodrug.

**[0037]** Any amine, hydroxy, or carboxyl side chain on the compounds disclosed herein can be esterified or amidified. The procedures and specific groups to be used to achieve this end are known to those of skill in the art and can readily be found in reference sources such as Greene and Wuts, *Protective Groups in Organic Synthesis*, 3<sup>rd</sup> Ed., John Wiley & Sons, New York, NY, 1999, which is incorporated herein in its entirety.

**[0038]** As used herein, "alkyl" refers to a straight or branched hydrocarbon chain that comprises a fully saturated (no double or triple bonds) hydrocarbon group. The alkyl group may have 1 to 20 carbon atoms (whenever it appears herein, a numerical range such as "1 to 20" refers to each integer in the given range; *e.g.*, "1 to 20 carbon atoms" means that the alkyl group may consist of 1 carbon atom, 2 carbon atoms, 3 carbon atoms, *etc.*, up to and including 20 carbon atoms, although the present definition also covers the occurrence of the term "alkyl" where no numerical range is designated). The alkyl group may also be a medium size alkyl having 1 to 10 carbon atoms. The alkyl group could also be a lower alkyl having 1 to 5 carbon atoms. The alkyl group of the compounds may be designated as "C<sub>1</sub>-C<sub>4</sub> alkyl" or similar designations. By way of example only, "C<sub>1</sub>-C<sub>4</sub> alkyl" indicates that there are one to four carbon atoms in the alkyl chain, *i.e.*, the alkyl chain is selected from methyl, ethyl, propyl, iso-propyl, n-butyl, iso-butyl, sec-butyl, and t-butyl. Typical alkyl groups include, but are in no way limited to, methyl, ethyl, propyl, isopropyl, butyl, isobutyl, tertiary butyl, pentyl, hexyl, and the like.

**[0039]** The alkyl group may be substituted or unsubstituted. When substituted, the substituent group(s) is(are) one or more group(s) individually and independently selected from alkenyl, alkynyl, cycloalkyl, cycloalkenyl, cycloalkynyl, aryl, heteroaryl, heteroalicyclic, aralkyl, heteroaralkyl, (heteroalicyclic)alkyl, hydroxy, protected hydroxyl, alkoxy, aryloxy, acyl, ester, mercapto, alkylthio, arylthio, cyano, halogen, carbonyl, thiocarbonyl, O-carbamyl, N-carbamyl, O-thiocarbamyl, N-thiocarbamyl, C-amido, N-amido, S-sulfonamido, N-sulfonamido, C-carboxy, protected C-carboxy, O-carboxy, isocyanato, thiocyanato, isothiocyanato, nitro, silyl, sulfenyl, sulfinyl, sulfonyl, haloalkyl (*e.g.*, mono-, di- and tri-haloalkyl), haloalkoxy (*e.g.*, mono-, di- and tri-haloalkoxy),

trihalomethanesulfonyl, trihalomethanesulfonamido, and amino, including mono- and di-substituted amino groups, and the protected derivatives thereof. Wherever a substituent is described as being “optionally substituted” that substituent may be substituted with one of the above substituents.

**[0040]** As used herein, “aryl” refers to a carbocyclic (all carbon) monocyclic or multicyclic aromatic ring system that has a fully delocalized pi-electron system. Examples of aryl groups include, but are not limited to, benzene, naphthalene and azulene. An aryl group of this invention may be substituted or unsubstituted. When substituted, hydrogen atoms are replaced by substituent group(s) that is(are) one or more group(s) independently selected from alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, cycloalkynyl, aryl, heteroaryl, heteroalicycyl, aralkyl, heteroaralkyl, (heteroalicycyl)alkyl, hydroxy, protected hydroxy, alkoxy, aryloxy, acyl, ester, mercapto, cyano, halogen, thiocarbonyl, O-carbamyl, N-carbamyl, O-thiocarbamyl, N-thiocarbamyl, C-amido, N-amido, S-sulfonamido, N-sulfonamido, C-carboxy, protected C-carboxy, O-carboxy, isocyanato, thiocyanato, isothiocyanato, nitro, silyl, sulfenyl, sulfinyl, sulfonyl, haloalkyl (e.g., mono-, di- and tri-haloalkyl), haloalkoxy (e.g., mono-, di- and tri-haloalkoxy), trihalomethanesulfonyl, trihalomethanesulfonamido, and amino, including mono- and di-substituted amino groups, and the protected derivatives thereof, unless the substituent groups are otherwise indicated.

**[0041]** A “paramagnetic metal chelate” is a complex wherein a ligand is bound to a paramagnetic metal ion. Examples include, but are not limited to, 1,4,7,10-Tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA)-Gd(III), DOTA-Yttrium-88, DOTA-Indium-111, diethylenetriaminepentaacetic acid (DTPA)-Gd(III), DTPA-yttrium-88, DTPA-Indium-111.

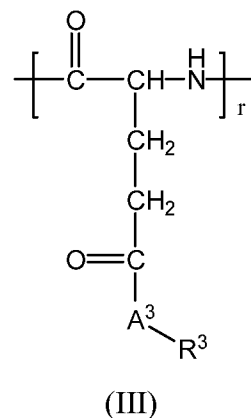
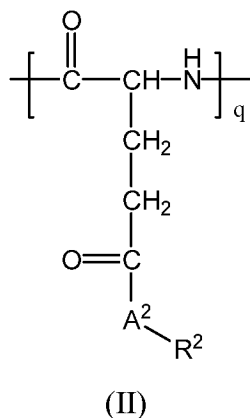
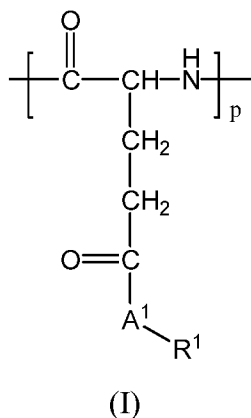
**[0042]** A “polydentate ligand” is a ligand that can bind itself through two or more points of attachment to a metal ion through, for example, coordinate covalent bonds. Examples of polydentate ligands include, but are not limited to, diethylenetriaminepentaacetic acid (DTPA), tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA), (1,2-ethanediyldinitrilo)tetraacetate (EDTA), ethylenediamine, 2,2'-bipyridine (bipy), 1,10-phenanthroline (phen), 1,2-bis(diphenylphosphino)ethane (DPPE), 2,4-pentanedione (acac), and ethanedioate (ox).

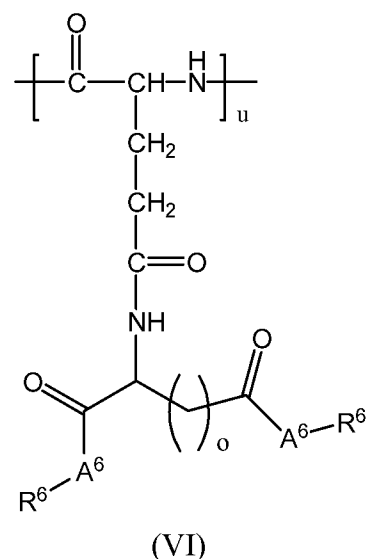
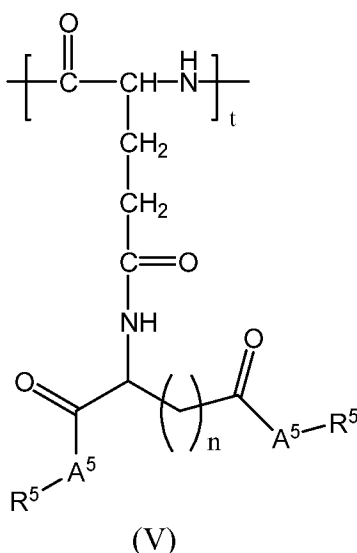
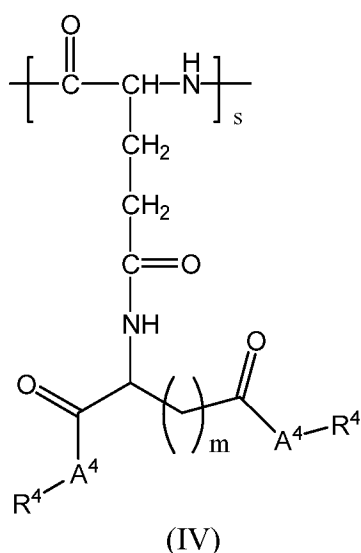
**[0043]** A “polydentate ligand precursor with protected oxygen atoms” is a polydentate ligand comprising oxygen atoms, such as the single-bonded oxygen atoms of carboxyl groups, that are protected with suitable protecting groups. Suitable protecting groups include, but are not limited to, lower alkyls, benzyls, and silyl groups.

**[0044]** A “stabilizing agent” is a substituent that enhances bioavailability and/or prolongs the half-life of a carrier-drug conjugate in vivo by rendering it more resistant to hydrolytic enzymes and less immunogenic. An exemplary stabilizing agent is polyethylene glycol (PEG).

**[0045]** It is understood that, in any compound described herein having one or more chiral centers, if an absolute stereochemistry is not expressly indicated, then each center may independently be of R-configuration or S-configuration or a mixture thereof. Thus, the compounds provided herein may be enantiomerically pure or be stereoisomeric mixtures. In addition it is understood that, in any compound described herein having one or more double bond(s) generating geometrical isomers that can be defined as E or Z each double bond may independently be E or Z a mixture thereof. Likewise, all tautomeric forms are also intended to be included.

**[0046]** An embodiment provides a polymer conjugate that can include at least one recurring unit selected from Formulae (I), (II), (III), (IV), (V) and (VI):





wherein each A<sup>1</sup>, A<sup>2</sup>, A<sup>3</sup>, A<sup>4</sup>, A<sup>5</sup> and A<sup>6</sup> can be independently oxygen or NR<sup>7</sup>, wherein R<sup>7</sup> can be hydrogen or a C<sub>1-4</sub> alkyl; each R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup> and R<sup>6</sup> are independently selected from hydrogen, a C<sub>1-10</sub> alkyl group, a C<sub>6-20</sub> aryl group, an ammonium group, an alkali metal, a polydentate ligand, a polydentate ligand precursor with protected oxygen atoms, a group that comprises a drug, a group that comprises a targeting agent, a group that comprises an optical imaging agent, a group that comprises a magnetic resonance imaging agent, and a group that comprises a stabilizing agent; m, n, and o can be each independently 1 or 2; p, q, r, s, t, and u can be each independently 0 or ≥ 1, wherein the sum of p, q, r, s, t, and u is 2 or greater; and provided that at least one of R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup> and R<sup>6</sup> is a group that comprises a first drug, at least one of R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup> and R<sup>6</sup> is a group that comprises a second drug, wherein the first drug and second drug are not the same.

**[0047]** The relative proportions of the recurring units, e.g., of the Formula (I), (II), (III), (IV), (V) and (VI), that are present in the polymer conjugates described herein can vary over a wide range. In an embodiment, p + q is 2 or greater; and r, s, t and u are 0. In an embodiment, s + t is 2 or greater; and p, q, r and u are 0. In an embodiment, p + q + r is 3 or greater; and s, t and u are 0. In an embodiment, s + t + u is 3 or greater; and q, r and u are 0. In an embodiment, p + s is 2 or greater; and q, r, t and u are 0. In an embodiment, p + q + s is 3 or greater; and r, t and u are 0. In an embodiment, p + s + t is 3 or greater; and q, r and t are 0. In an embodiment, p + q + s + t is 4 or greater; and r and u are 0. In an embodiment, p + q + r + s + t is 5 or greater; and u is 0. In an embodiment, p + q + s + t + u is 5 or greater; and r is 0. In an embodiment, p + q + r + s + t + u is 6 or greater.

**[0048]** Many different types of drugs may be used for the first drug. In an embodiment, the first drug can be a first hydrophobic drug. In an embodiment, the first hydrophobic drug can include an anticancer drug. In an embodiment, the anticancer drug can be selected from a taxane, a camptotheca and an anthracycline. In an embodiment, the taxane can be paclitaxel or docetaxel. In an embodiment, the taxane can be paclitaxel. In one embodiment wherein the first hydrophobic drug comprises paclitaxel, the paclitaxel can be attached or conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C2'-carbon of the paclitaxel. In another embodiment, the paclitaxel can be attached or conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C7-carbon of the paclitaxel. In an embodiment, the camptotheca can be camptothecin. In an embodiment, the anthracycline can be doxorubicin.

**[0049]** Many different types of drugs may be used for the second drug. In an embodiment, the second drug can be a second hydrophobic drug. In an embodiment, the second hydrophobic drug can include an anticancer drug. In an embodiment, the anticancer drug can be selected from a taxane, a camptotheca and an anthracycline. In an embodiment, the taxane can be selected from paclitaxel and docetaxel. In an embodiment, the taxane can be paclitaxel. In one embodiment wherein the second hydrophobic drug comprises paclitaxel, the paclitaxel can be attached or conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C2'-carbon of the paclitaxel. In another embodiment, the paclitaxel can be attached or conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C7-carbon of the paclitaxel. In an embodiment, the camptotheca can be camptothecin. In an embodiment, the anthracycline can be doxorubicin.

**[0050]** Figure 1 schematically illustrates an embodiment wherein a polymer conjugate includes a single type of drug attached thereto. The polymer conjugate, which may be represented by numerous types of polymeric material, contains numerous side branches to which a drug, for example, paclitaxel, doxorubicin, or camptothecin, can be conjugated.

**[0051]** Figure 2 schematically illustrates an embodiment wherein the polymer conjugate includes up to three types of drugs attached thereto. The polymer conjugate may be represented by numerous types of polymeric materials. For example, polyamino acids,

such as polyglutamic acid, and their associated salts may be used to form the polymer conjugates described herein. Additionally, polyamino amino acids, such as polyglutamic glutamic acid, and their associated salts may be used to form the polymer conjugates described herein. Additionally, copolymers of polyamino acids and polyamino amino acids, and their associated salts may be used to form the polymer conjugates described herein. Attachment of a plurality of drugs can allow for combination therapy of a disease or illness, such as cancer. For example, any combination of taxanes, such as paclitaxel and docetaxel, camptothecas, such as camptothecin, and anthracyclines, such as doxorubicin, may be conjugated to the polymers described herein.

**[0052]** The amount of first drug conjugated to the polymer may vary over a wide range. In an embodiment, the polymer conjugate can include an amount of the first drug in the range of about 0.5% to about 50% (weight/weight) based on the mass ratio of the first drug to the polymer conjugate (the weight of the first drug is accounted for in the polymer conjugate). In an embodiment, the polymer conjugate can include an amount of the first drug in the range of about 1% to about 40% (weight/weight) based on the mass ratio of the first drug to the polymer conjugate. In an embodiment, the polymer conjugate can include an amount of the first drug in the range of about 1% to about 30% (weight/weight) based on the mass ratio of the first drug to the polymer conjugate. In an embodiment, the polymer conjugate can include an amount of the first drug in the range of about 1% to about 20% (weight/weight) based on the mass ratio of the first drug to the polymer conjugate. In an embodiment, the polymer conjugate can include an amount of the first drug in the range of about 1% to about 10% (weight/weight) based on the mass ratio of the first drug to the polymer conjugate.

**[0053]** The amount of second drug conjugated to the polymer may also vary over a wide range. In an embodiment, the polymer conjugate can include an amount of the second drug in the range of about 0.5% to about 50% (weight/weight) based on the mass ratio of the second drug to the polymer conjugate (the weight of the second drug is accounted for in the polymer conjugate). In an embodiment, the polymer conjugate can include an amount of the second drug in the range of about 1% to about 40% (weight/weight) based on the mass ratio of the second drug to the polymer conjugate. In an embodiment, the polymer conjugate can include an amount of the second drug in the range of about 1% to about 30% (weight/weight)

based on the mass ratio of the second drug to the polymer conjugate. In an embodiment, the polymer conjugate can include an amount of the second drug in the range of about 1% to about 20% (weight/weight) based on the mass ratio of the second drug to the polymer conjugate. In an embodiment, the polymer conjugate can include an amount of the second drug in the range of about 1% to about 10% (weight/weight) based on the mass ratio of the second drug to the polymer conjugate.

**[0054]** The total amount of first drug and second drug conjugated to the polymer may vary over a wide range. In an embodiment, the polymer conjugate can include a total amount of the first drug and the second drug in the range of about 1% to about 50% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate (the weight of the drugs is accounted for in the polymer conjugate). In an embodiment, the polymer conjugate can include a total amount of the first drug and the second drug in the range of about 1% to about 40% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate. In an embodiment, the polymer conjugate can include a total amount of the first drug and the second drug in the range of about 1% to about 30% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate. In an embodiment, the polymer conjugate can include a total amount of the first drug and the second drug in the range of about 1% to about 20% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate. In an embodiment, the polymer conjugate can include a total amount of the first drug and the second drug in the range of about 1% to about 10% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate.

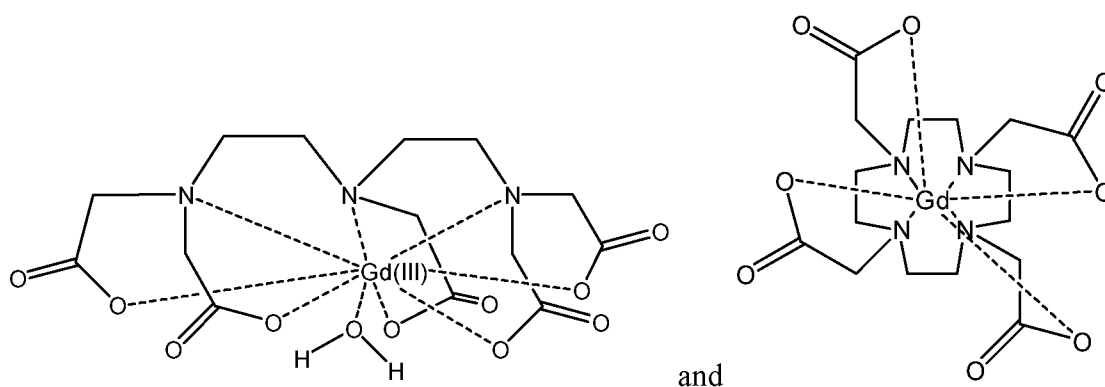
**[0055]** In an embodiment, each of  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  can be independently a group that comprises an agent. Many types of agents may be used. For example, the agent(s) may be selected from a targeting agent, an optical imaging agent, a magnetic resonance imaging agent, and a stabilizing agent.

**[0056]** The agent may comprise any type of active compound. In an embodiment, the agent can comprise an optical imaging agent. In an embodiment, the optical imaging agent can be one or more selected from an acridine dye, a coumarine dye, a rhodamine dye, a xanthene dye, a cyanine dye, and a pyrene dye. For instance, specific optical imaging agents may include Texas Red, Alexa Fluor® dye, BODIPY® dye, Fluorescein, Oregon Green®

dye, and Rhodamine Green<sup>TM</sup> dye, which are commercially available or readily prepared by methods known to those skilled in the art.

**[0057]** In an embodiment, the agent can comprise a targeting agent. In an embodiment, the targeting agent can be one or more selected from an arginine-glycine-aspartate (RGD) peptide, fibronectin, folate, galactose, an apolipoprotein, insulin, transferrin, a fibroblast growth factor (FGF), an epidermal growth factor (EGF), and an antibody. In an embodiment, the targeting agent can interact with a receptor selected from  $\alpha_v\beta_3$ -integrin, folate, asialoglycoprotein, a low-density lipoprotein (LDL), an insulin receptor, a transferrin receptor, a fibroblast growth factor (FGF) receptor, an epidermal growth factor (EGF) receptor, and an antibody receptor. In an embodiment, the arginine-glycine-aspartate (RGD) peptide can be cyclic(fKRGD).

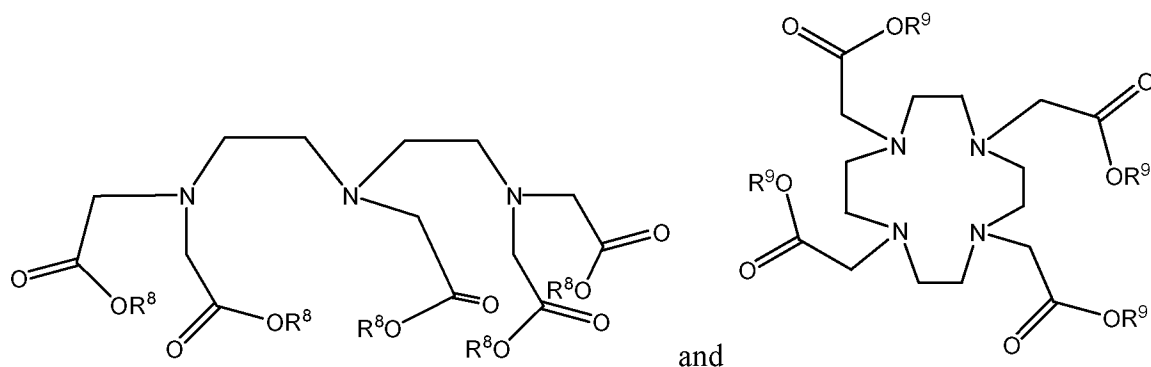
**[0058]** In an embodiment, the agent can comprise a magnetic resonance imaging agent. In an embodiment, the magnetic resonance imaging agent can include a paramagnetic metal compound. For example, the magnetic resonance imaging agent may include a Gd(III) compound. In an embodiment, the Gd(III) compound can be selected from:



**[0059]** In an embodiment, the agent can comprise a stabilizing agent. In a preferred embodiment, the stabilizing agent is polyethylene glycol.

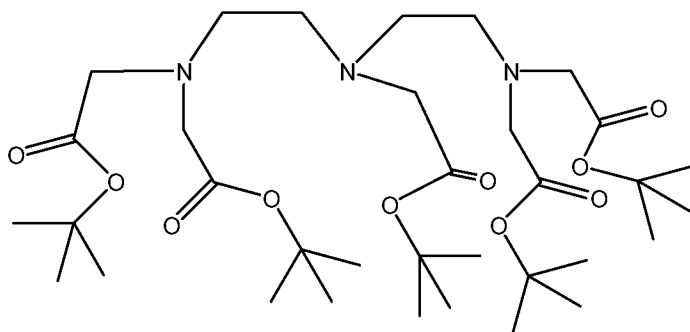
**[0060]** In an embodiment, the polymer conjugate can comprise a polydentate ligand. In an embodiment, each of  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  can be independently selected to comprise a group that includes a polydentate ligand. In an embodiment, the polydentate ligand may be capable of reaction with a paramagnetic metal to form a magnetic resonance imaging agent. The polydentate ligand may comprise several carboxylic acid and/or carboxylate groups. In an embodiment, the polydentate ligand can be selected from:





wherein each  $R^8$  and each  $R^9$  can be independently selected from hydrogen, ammonium, and an alkali metal.

**[0061]** In an embodiment, the polymer conjugate comprises a polydentate ligand precursor. In an embodiment, each of  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  can be independently selected to comprise a group that includes a polydentate ligand precursor. In such an embodiment, the oxygen atoms of the polydentate ligand may be protected by a suitable protecting group. Suitable protecting groups include, but are not limited to, lower alkyls, benzyls, and silyl groups. One example of a polydentate ligand precursor having protecting groups is provided as follows:



**[0062]** In some embodiments, the polymers and/or polymer conjugates described herein comprise an alkali metal. In an embodiment, each of  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  can be independently selected to comprise an alkali metal, such as lithium (Li), sodium (Na), potassium (K), rubidium (Rb), and cesium (Cs). In an embodiment, the alkali metal can be sodium. In an embodiment, each of  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  of the polymers and/or polymer conjugates described herein can comprise hydrogen, a  $C_{1-10}$  alkyl group, a  $C_{6-20}$  aryl group or an ammonium group.

**[0063]** The amount of agent(s), such as a targeting agent, an optical imaging agent, a magnetic resonance imaging agent, and/or a stabilizing agent, present in the polymer

can vary over a wide range. Additionally, the amount of a ligand or a ligand precursor present in the polymer can vary over a wide range. In an embodiment, the polymer conjugate comprises an amount of an agent(s), a ligand, and/or a ligand precursor in the range of about 0.1% to about 50% (weight/weight) based on the mass ratio of the agent(s), ligand, and/or ligand precursor to the polymer conjugate (the weight of the agent(s), ligand, and/or ligand precursor, along with the weight of conjugated drugs, is accounted for in the polymer conjugate). In an embodiment, the polymer conjugate comprises an amount of an agent(s), a ligand, and/or a ligand precursor in the range of about 1% to about 40% (weight/weight) based on the mass ratio of the agent(s), ligand, and/or ligand precursor to the polymer conjugate. In an embodiment, the polymer conjugate comprises an amount of an agent(s), a ligand, and/or a ligand precursor in the range of about 1% to about 30% (weight/weight) based on the mass ratio of the agent(s), ligand, and/or ligand precursor to the polymer conjugate. In an embodiment, the polymer conjugate comprises an amount of an agent(s), a ligand, and/or a ligand precursor in the range of about 1% to about 20% (weight/weight) based on the mass ratio of the agent(s), ligand, and/or ligand precursor to the polymer conjugate. In an embodiment, the polymer conjugate comprises an amount of an agent(s), a ligand, and/or a ligand precursor in the range of about 1% to about 10% (weight/weight) based on the mass ratio of the agent(s), ligand, and/or ligand precursor to the polymer conjugate. In an embodiment, the polymer conjugate comprises an amount of an agent(s), a ligand, and/or a ligand precursor in the range of about 5% to about 40% (weight/weight) based on the mass ratio of the agent(s), ligand, and/or ligand precursor to the polymer conjugate. In an embodiment, the polymer conjugate comprises an amount of an agent(s), a ligand, and/or a ligand precursor in the range of about 10% to about 30% (weight/weight) based on the mass ratio of the agent(s), ligand, and/or ligand precursor to the polymer conjugate. In an embodiment, the polymer conjugate comprises an amount of an agent(s), a ligand, and/or a ligand precursor in the range of about 20% to about 40% (weight/weight) based on the mass ratio of the agent(s), ligand, and/or ligand precursor to the polymer conjugate. In an embodiment, the polymer conjugate comprises an amount of an agent(s), a ligand, and/or a ligand precursor in the range of about 30% to about 50% (weight/weight) based on the mass ratio of the agent(s), ligand, and/or ligand precursor to the polymer conjugate.

**[0064]** In an embodiment, at least one of  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  can be a group that comprises a third drug. In an embodiment, the third drug can be different from the first drug and the second drug. Many different types of drugs may be used for the third drug. In an embodiment, the third drug can be a third hydrophobic drug. In an embodiment, the third hydrophobic drug can comprise an anticancer drug. In an embodiment, the anticancer drug can be selected from a taxane, a camptotheca, and an anthracycline. In an embodiment, the taxane can be selected from paclitaxel and docetaxel. In an embodiment, the taxane can be paclitaxel. In one embodiment wherein the third hydrophobic drug comprises paclitaxel, the paclitaxel can be attached or conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C2'-carbon of the paclitaxel. In another embodiment, the paclitaxel can be attached or conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C7-carbon of the paclitaxel. In an embodiment, the camptotheca can be camptothecin. In an embodiment, the anthracycline can be doxorubicin.

**[0065]** The amount of third drug conjugated to the polymer may also vary over a wide range. In an embodiment, the polymer conjugate can include an amount of the third drug in the range of about 0.5% to about 50% (weight/weight) based on the mass ratio of the third drug to the polymer conjugate (the weight of the third drug is accounted for in the polymer conjugate). In an embodiment, the polymer conjugate can include an amount of the third drug in the range of about 1% to about 40% (weight/weight) based on the mass ratio of the third drug to the polymer conjugate. In an embodiment, the polymer conjugate can include an amount of the third drug in the range of about 1% to about 30% (weight/weight) based on the mass ratio of the third drug to the polymer conjugate. In an embodiment, the polymer conjugate can include an amount of the third drug in the range of about 1% to about 20% (weight/weight) based on the mass ratio of the third drug to the polymer conjugate. In an embodiment, the polymer conjugate can include an amount of the third drug in the range of about 1% to about 10% (weight/weight) based on the mass ratio of the third drug to the polymer conjugate.

**[0066]** The total amount of first drug, second drug, and third drug conjugated to the polymer may vary over a wide range. In an embodiment, the polymer conjugate can include a total amount of the first drug, second drug, and the third drug in the range of about

1% to about 50% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate (the weight of the drugs is accounted for in the polymer conjugate). In an embodiment, the polymer conjugate can include a total amount of the first drug, second drug, and the third drug in the range of about 1% to about 40% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate. In an embodiment, the polymer conjugate can include a total amount of the first drug, second drug, and the third drug in the range of about 1% to about 30% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate. In an embodiment, the polymer conjugate can include a total amount of the first drug, second drug, and the third drug in the range of about 1% to about 20% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate. In an embodiment, the polymer conjugate can include a total amount of the first drug, second drug, and the third drug in the range of about 1% to about 10% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate.

**[0067]** In an embodiment, at least one of m, n, or o can be 1. In an embodiment, at least one of m, n, or o can be 2. In some embodiments, m can be 1. In other embodiments, m can be 2. In some embodiments, n can be 1. In other embodiments, n can be 2. In some embodiments, o can be 1. In other embodiments, o can be 2.

**[0068]** One or more of a group that comprises a drug, a group that comprises a targeting agent, a group that comprises an optical imaging agent, a group that comprises a magnetic resonance imaging agent, a group that comprises a polydentate ligand, a group that comprises a polydentate ligand precursor, and a group that comprises a stabilizing agent may be conjugated to the polymer in many different ways. In some embodiments, the aforementioned compounds can be directly attached to the polymer, e.g., to a recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI). In one embodiment, one or more of a group that comprises a drug, a group that comprises a targeting agent, a group that comprises an optical imaging agent, a group that comprises a magnetic resonance imaging agent, a group that comprises a polydentate ligand, a group that comprises a polydentate ligand precursor, and a group that comprises a stabilizing agent can be directly attached to the polymer through an oxygen, a sulfur, a nitrogen and/or carbon atom of the agent or drug.

**[0069]** In other embodiments, one or more of a group that comprises a drug, a group that comprises a targeting agent, a group that comprises an optical imaging agent, a

group that comprises a magnetic resonance imaging agent, a group that comprises a polydentate ligand, a group that comprises a polydentate ligand precursor, and a group that comprises a stabilizing agent can further include a linker group. In an embodiment, the group that comprises the first drug further can include a linker group. In an embodiment, the group that comprises the second drug further can include a linker group. In an embodiment, the group that comprises the third drug further can include a linker group. In an embodiment, the group that comprises a targeting agent, the group that comprises an optical imaging agent, the group that comprises a magnetic resonance imaging agent, the group that comprises a polydentate ligand, the group that comprises a polydentate ligand precursor, and/or the group that comprises a stabilizing agent can further include a linker group.. A linker group is a group that attaches, for example, the agent (or the compound that comprises the agent) to the polymer. In an embodiment, one or more of the aforementioned compounds can be attached to the polymer, e.g., to a recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI), through a linker group. The linker group may be relatively small. For instance, the linker group may comprise an amine, an amide, an ether, an ester, a hydroxyl group, a carbonyl group, or a thioether group. Alternatively, the linker group may be relatively large. For instance, the linker group may comprise an alkyl group, an ether group, an aryl group, an aryl(C<sub>1-6</sub> alkyl) group (e.g., phenyl-(CH<sub>2</sub>)<sub>1-4</sub>-), a heteroaryl group, or a heteroaryl(C<sub>1-6</sub> alkyl) group. In one embodiment, the linker can be -NH(CH<sub>2</sub>)<sub>1-4</sub>-NH-. In another embodiment, the linker can be -(CH<sub>2</sub>)<sub>1-4</sub>-aryl-NH-. The linker group can be attached to one or more of a group that comprises a drug, a group that comprises a targeting agent, a group that comprises an optical imaging agent, a group that comprises a magnetic resonance imaging agent, a group that comprises a polydentate ligand, a group that comprises a polydentate ligand precursor, or a group that comprises a stabilizing agent at any suitable position. For example, the linker group can be attached in place of a hydrogen at a carbon of one of the aforementioned compounds. The linker group can be added to the compounds using methods known to those skilled in the art.

**[0070]** Polymers comprising a recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) can be copolymers comprising two or more different recurring units of the formulae (I), (II), (III), (IV), (V), and/or (VI). Further, polymers comprising a recurring unit of the formulae (I), (II), (III), (IV), (V), and/or (VI) can be copolymers that comprise other

recurring units that are not of the formulae (I), (II), (III), (IV), (V), and/or (VI). A broad variety of other recurring units may be included in the polymer conjugates described herein. The number of recurring units of the formulae (I), (II), (III), (IV), (V), and/or (VI) in the polymer can vary over a broad range, but is preferably in the range of from about 50 to about 5,000, and more preferably from about 100 to about 2,000.

**[0071]** The percentage of recurring units of formula (I) in the polymer conjugate, based on the total number of recurring units, may vary over a wide range. In an embodiment, the polymer conjugate may comprise up to about 99 mole % of the recurring unit of formula (I), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 99 mole % of the recurring unit of formula (I), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer may comprise about 1 mole % to about 50 mole % of the recurring unit of formula (I) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 30 mole % of the recurring unit of formula (I) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 20 mole % of the recurring unit of formula (I) based on the total moles of recurring units of the polymer conjugate. In another embodiment, the polymer conjugate may comprise about 1 mole % to about 10 mole % of the recurring unit of formula (I) based on the total moles of recurring units of the polymer conjugate.

**[0072]** The percentage of recurring units of formula (II) in the polymer conjugate, based on the total number of recurring units, may vary over a wide range. In an embodiment, the polymer conjugate may comprise up to about 99 mole % of the recurring unit of formula (II), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 99 mole % of the recurring unit of formula (II), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 50 mole % of the recurring unit of formula (II) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 30 mole % of the recurring unit of formula (II) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer may comprise about 1 mole % to

about 20 mole % of the recurring unit of formula (II) based on the total moles of recurring units of the polymer conjugate. In another embodiment, the polymer conjugate may comprise about 1 mole % to about 10 mole % of the recurring unit of formula (II) based on the total moles of recurring units of the polymer conjugate.

**[0073]** The percentage of recurring units of formula (III) in the polymer conjugate, based on the total number of recurring units, may vary over a wide range. In an embodiment, the polymer conjugate may comprise up to about 99 mole % of the recurring unit of formula (III), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 99 mole % of the recurring unit of formula (III), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 50 mole % of the recurring unit of formula (III) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 30 mole % of the recurring unit of formula (III) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 20 mole % of the recurring unit of formula (III) based on the total moles of recurring units of the polymer conjugate. In another embodiment, the polymer conjugate may comprise about 1 mole % to about 10 mole % of the recurring unit of formula (III) based on the total moles of recurring units of the polymer conjugate.

**[0074]** The percentage of recurring units of formula (IV) in the polymer conjugate, based on the total number of recurring units, may vary over a wide range. In an embodiment, the polymer conjugate may comprise up to about 99 mole % of the recurring unit of formula (IV), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 99 mole % of the recurring unit of formula (IV), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 50 mole % of the recurring unit of formula (IV) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 30 mole % of the recurring unit of formula (IV) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 20 mole % of the recurring unit of formula (IV) based on

the total moles of recurring units of the polymer conjugate. In another embodiment, the polymer conjugate may comprise about 1 mole % to about 10 mole % of the recurring unit of formula (IV) based on the total moles of recurring units of the polymer conjugate.

**[0075]** The percentage of recurring units of formula (V) in the polymer conjugate, based on the total number of recurring units, may vary over a wide range. In an embodiment, the polymer conjugate may comprise up to about 99 mole % of the recurring unit of formula (V), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 99 mole % of the recurring unit of formula (V), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 50 mole % of the recurring unit of formula (V) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 30 mole % of the recurring unit of formula (V) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 20 mole % of the recurring unit of formula (V) based on the total moles of recurring units of the polymer conjugate. In another embodiment, the polymer conjugate may comprise about 1 mole % to about 10 mole % of the recurring unit of formula (V) based on the total moles of recurring units of the polymer conjugate.

**[0076]** The percentage of recurring units of formula (VI) in the polymer conjugate, based on the total number of recurring units, may vary over a wide range. In an embodiment, the polymer conjugate may comprise up to about 99 mole % of the recurring unit of formula (VI), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 99 mole % of the recurring unit of formula (VI), based on the total moles of recurring units in the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 50 mole % of the recurring unit of formula (VI) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 30 mole % of the recurring unit of formula (VI) based on the total moles of recurring units of the polymer conjugate. In an embodiment, the polymer conjugate may comprise about 1 mole % to about 20 mole % of the recurring unit of formula (VI) based on the total moles of recurring units of the polymer conjugate. In another embodiment, the



polymer conjugate may comprise about 1 mole % to about 10 mole % of the recurring unit of formula (VI) based on the total moles of recurring units of the polymer conjugate.

**[0077]** In an embodiment, the polymer conjugate can include two or more recurring units selected from a recurring unit of the formula (I), a recurring unit of the formula (II), a recurring unit of the formula (III), a recurring unit of the formula (IV), a recurring unit of the formula (V), and a recurring unit of the formula (VI). In an embodiment, the polymer conjugate can include three or more recurring units selected from a recurring unit of the formula (I), a recurring unit of the formula (II), a recurring unit of the formula (III), a recurring unit of the formula (IV), a recurring unit of the formula (V), and a recurring unit of the formula (VI). In an embodiment, the polymer conjugate can include four or more recurring units selected from a recurring unit of the formula (I), a recurring unit of the formula (II), a recurring unit of the formula (III), a recurring unit of the formula (IV), a recurring unit of the formula (V), and a recurring unit of the formula (VI). In an embodiment, the polymer conjugate can include five or more recurring units selected from a recurring unit of the formula (I), a recurring unit of the formula (II), a recurring unit of the formula (III), a recurring unit of the formula (IV), a recurring unit of the formula (V), and a recurring unit of the formula (VI). In an embodiment, the polymer conjugate can include six different recurring units of the formulae (I), (II), (III), (IV), (V), and (VI).

**[0078]** The amount of each recurring unit (e.g., mole percent) present in the polymer conjugate can vary greatly, as set forth above. In an embodiment, selection of an amount of any one recurring unit of the formulae (I), (II), (III), (IV), (V), and/or (VI) can be independent of the selection of an amount of a different recurring unit of the formulae (I), (II), (III), (IV), (V), and/or (VI).

**[0079]** The polymer conjugate can contain one or more chiral carbon atoms. The chiral carbon (which may be indicated by an asterisk \*) can have the *rectus* (right handed) or the *sinister* (left handed) configuration, and thus the recurring unit may be racemic, enantiomeric or enantiomerically enriched. The symbols “n” and “\*” (designating a chiral carbon), as used elsewhere herein, have the same meaning as specified above, unless otherwise stated.

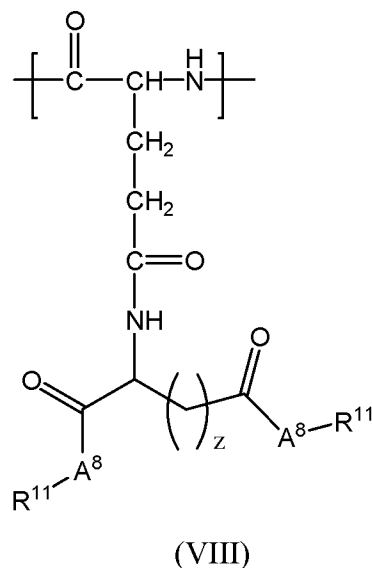
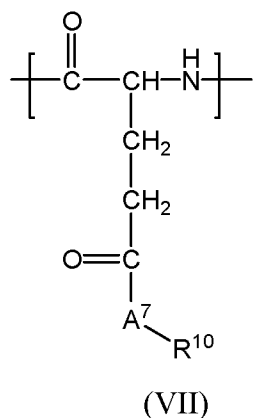
**[0080]** In an embodiment, the amounts of the agent(s), the amount of first, second, and/or third drug, and the percentage of the recurring unit of the formulae (I), (II),

(III), (IV), (V), and/or (VI) in the polymer conjugate can be selected to provide a polymer conjugate solubility that is greater than that of a comparable polyglutamic acid conjugate that comprises substantially the same amount of the agent(s) and/or drugs. The range of pH values over which the polymer conjugate, comprising recurring units of the formulae (I), (II), (III), (IV), (V), and/or (VI), has greater solubility than that of a comparable polyglutamic acid conjugate may be narrow or broad. Solubility is measured by forming a polymer conjugate solution comprising at least 5 mg/mL of the polymer conjugate in 0.9 wt. % aqueous NaCl at about 22°C, and determining the optical clarity. In an embodiment, the polymer conjugate is soluble over a pH range of at least about three pH units. In another embodiment, the polymer conjugate is soluble over a pH range of at least about 8 pH units. In another embodiment, the polymer conjugate is soluble over a pH range of at least about 9 pH units. In another embodiment, the pH range over which the polymer conjugate is soluble includes at least one pH value in the range of about 2 to about 5, e.g., at pH = 2, pH = 3, pH = 4 and/or pH = 5. Preferably, the pH range over which the polymer conjugate is soluble is broader than the pH range over which the comparable polyglutamic acid conjugate is soluble. For example, in an embodiment, the polymer conjugate is soluble over a pH range that is at least about one pH unit broader, preferably at least about two pH units broader, than the pH range over which the comparable polyglutamic acid conjugate is soluble.

**[0081]** The amount of polymer conjugate placed in solution to measure solubility can also vary greatly. In one embodiment, solubility is measured when the tested polymer conjugate solution comprises at least about 5 mg/mL of the polymer conjugate. In another embodiment, solubility is measured when the tested polymer conjugate solution comprises at least about 10 mg/mL of the polymer conjugate. In another embodiment, solubility is measured when the tested polymer conjugate solution comprises at least about 25 mg/mL of the polymer conjugate. In another embodiment, solubility is measured when the tested polymer conjugate solution comprises at least about 100 mg/mL of the polymer conjugate. In another embodiment, solubility is measured when the tested polymer conjugate solution comprises at least about 150 mg/mL of the polymer conjugate. Those skilled in the art will understand that the comparable polyglutamic acid conjugate is tested at about the same concentration as that of the tested polymer conjugate.

**[0082]** Polymers comprising a recurring unit of the formulae (I), (II), (III), (IV), (V), and/or (VI) may be prepared in various ways. In an embodiment, a polymeric reactant can be dissolved or partially dissolved in a solvent to form a dissolved or partially dissolved polymeric reactant. The dissolved or partially dissolved polymeric reactant can be then reacted with a second reactant and third reactant to form an intermediate product or, in some embodiments, a polymer comprising a recurring units of the formulae (I), (II), (III), (IV), (V), and/or (VI). In an embodiment, the second reactant can include a first drug. In an embodiment, the third reactant can include a second drug.

**[0083]** The polymeric reactant may comprise any suitable material capable of forming a polymer comprising a recurring unit of the formulae (I), (II), (III), (IV), (V), and/or (VI). In an embodiment, the polymeric reactant can include a recurring unit selected formula (VII) and formula (VIII):



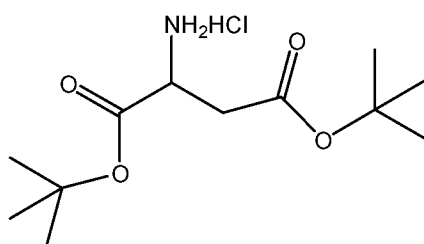
wherein  $z$  can be independently 1 or 2;  $A^7$  and each  $A^8$  can be oxygen; and each  $R^{10}$  and  $R^{11}$  can be independently selected from hydrogen, ammonium, and an alkali metal, for example, lithium (Li), sodium (Na), potassium (K), rubidium (Rb), and cesium (Cs).

**[0084]** In an embodiment, a polymer reactant comprising a recurring unit of the formula (VII) can be produced starting with polyglutamic acid. Alternatively, in another embodiment, the polymer may be created by first converting the starting polyglutamic acid material into its salt form. The salt form of polyglutamic can be obtained by reacting polyglutamic acid with a suitable base, e.g., sodium bicarbonate. The weight average

molecular weight of the polyglutamic acid is not limited, but is preferably from about 10,000 to about 500,000 daltons, and more preferably from about 25,000 to about 300,000 daltons.

**[0085]** In an embodiment, a polymer reactant comprising a recurring unit of the formula (VIII) can be produced starting with polyglutamic acid and an amino acid such as aspartic and/or glutamic acid. Alternatively, in another embodiment, the polymer may be created by first converting the starting polyglutamic acid material into its salt form. The salt form of polyglutamic can be obtained by reacting polyglutamic acid with a suitable base, e.g., sodium bicarbonate. An amino acid moiety can be attached to the pendant carboxylic acid group of the polyglutamic acid. The weight average molecular weight of the polyglutamic acid is not limited, but is preferably from about 10,000 to about 500,000 daltons, and more preferably from about 25,000 to about 300,000 daltons. Such a reaction may be used to create poly-( $\gamma$ -L-aspartyl-glutamine) or poly-( $\gamma$ -L-glutamyl-glutamine).

**[0086]** In an embodiment, the amino acid can be protected by a protecting group before attachment to the polyglutamic acid. One example of a protected amino acid moiety suitable for this reaction is L-aspartic acid di-t-butyl ester hydrochloride, shown below:



L-aspartic acid di-t-butyl ester hydrochloride

**[0087]** Reaction of the polyglutamic acid with the amino acid may take place in the presence of any suitable solvent. In an embodiment, the solvent can be an aprotic solvent. In a preferred embodiment, the solvent is N,N'-dimethylformamide. In an embodiment, a coupling agent such as EDC, DCC, CDI, DSC, HATU, HBTU, HCTU, PyBOP®, PyBroP®, TBTU, and BOP can be used in the reaction between the polyglutamic acid and the amino acid. In other embodiments, polyglutamic acid and an amino acid can be reacted using a catalyst (e.g., DMAP).

**[0088]** The polymer may be recovered and/or purified by methods known to those skilled in the art. For example, the solvent may be removed by suitable methods, for instance, rotary evaporation. Additionally, the reaction mixture may be filtered into an acidic water

solution to induce precipitation. The resultant precipitate can then be filtered, and washed with water.

**[0089]** In an embodiment, a polymer reactant comprising a recurring unit of the formula (VII) can also include a recurring unit of formula (VIII). One method for forming a polymer reactant comprising a recurring unit of the formula (VII) and a recurring unit of formula (VIII) is by starting with polyglutamic acid and reacting it with an amino acid such as aspartic and/or glutamic acid, in an amount that is less than 1.0 equivalents of the amino acid based on polyglutamic acid. For example, in one embodiment, 0.7 equivalents of an amino acid based on the polyglutamic acid can be reacted with polyglutamic acid, so that about 70% of the recurring units of the resulting polymer include the amino acid. As discussed above, the oxygen atoms of the amino acid can be protected using a suitable protecting group. In an embodiment, the amino acid may be L-aspartic acid or L-glutamic acid. In another embodiment, the oxygen atoms of the amino acid can be protected with t-butyl groups. If the oxygen atoms of the amino acid are protected, the protecting groups can be removed using known methods such as a suitable acid (e.g., trifluoroacetic acid).

**[0090]** In an embodiment, the polymeric reactant can be dissolved or partially dissolved with a second reactant and a third reactant, wherein the second reactant comprises the first drug and the third reactant comprises the second drug.

**[0091]** The second reactant may comprise many different types of drugs. In an embodiment, the first drug can include an anticancer drug. In an embodiment, the anticancer drug can be selected from a taxane, a camptotheca, and an anthracycline. In an embodiment, the taxane can be selected from paclitaxel and docetaxel. In an embodiment, the taxane can be paclitaxel. In one embodiment wherein the first hydrophobic drug comprises paclitaxel, the paclitaxel can be attached or conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C2'-carbon of the paclitaxel. In another embodiment, the paclitaxel can be attached or conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C7-carbon of the paclitaxel. In an embodiment, the camptotheca can be camptothecin. In an embodiment, the anthracycline can be doxorubicin.

**[0092]** The third reactant may comprise many different types of drugs. In an embodiment, the second drug can include an anticancer drug. In an embodiment, the

anticancer drug can be selected from a taxane, a camptotheca, and an anthracycline. In an embodiment, the taxane can be selected from paclitaxel and docetaxel. In an embodiment, the taxane can be paclitaxel. In one embodiment wherein the second hydrophobic drug comprises paclitaxel, the paclitaxel can be attached or conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C2'-carbon of the paclitaxel. In another embodiment, the paclitaxel can be attached or conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C7-carbon of the paclitaxel. In an embodiment, the camptotheca can be camptothecin. In an embodiment, the anthracycline can be doxorubicin.

**[0093]** In an embodiment, the second reactant can include a substituent selected from hydroxy and amine. In an embodiment, the third reactant can include a substituent selected from hydroxy and amine.

**[0094]** In an embodiment, the dissolved or partially dissolved polymer reactant can be reacted with at least a portion of the second reactant before the dissolved or partially dissolved reactant is reacted with at least a portion of the third reactant. In another embodiment, the dissolved or partially dissolved polymer reactant can be reacted with at least a portion of the second reactant after the dissolved partially dissolved reactant is reacted with at least a portion of the third reactant. In an embodiment, the dissolved or partially dissolved polymer reactant can be reacted with at least a portion of the second reactant at about the same time as the dissolved or partially dissolved polymer reactant is reacted with at least a portion of the third reactant.

**[0095]** Figure 3 illustrates a non-limiting example of a reaction scheme for the preparation of various polyamino polymer conjugates. The illustrated reaction scheme shows reaction steps for conjugating a plurality of drugs to polyglutamic acid. Other forms of polyglutamic acid may also be used in the reaction scheme illustrated by Figure 3. For example, alkali salts or ammonium salts of polyglutamic acid may be used. In an embodiment, a polymer reactant comprising a recurring unit of formula (VII) can be used in the reaction scheme illustrated by Figure 3.

**[0096]** As illustrated in Figure 3, a dissolved or partially dissolved polyglutamic acid (PGA) is reacted with paclitaxel in the presence of a coupling agent to form a polyglutamic acid-paclitaxel conjugate (PGA-paclitaxel). The dissolved or partially

dissolved PGA-paclitaxel may then further be reacted with a second drug in the presence of a coupling agent. The second drug may be doxorubicin, resulting in a PGA-(paclitaxel)-doxorubicin conjugate. Alternatively, the second drug may be camptothecin, resulting in a PGA-(paclitaxel)-camptothecin conjugate. In an embodiment that is shown in Figure 3, PGA-(paclitaxel)-camptothecin is dissolved or partially dissolved and reacted with a third drug, doxorubicin, to form a PGA-(paclitaxel)-(camptothecin)-doxorubicin conjugate.

**[0097]** While Figure 3 displays embodiments where the order of drug conjugation is described, the order should not be construed as limiting. The first drug conjugated to the polymer can be any one of a taxane, a camptotheca, or an anthracycline, as described herein. The second drug conjugated to the polymer can be any one of a taxane, a camptotheca, or an anthracycline, as described herein. Additionally, the third drug conjugated to the polymer can be any one of a taxane, a camptotheca, or an anthracycline, as described herein. Furthermore, any one of the first, second, and/or third drugs may be conjugated to the polymer at about the same time as any one of the other first, second, and/or third drugs.

**[0098]** Figure 4 illustrates a non-limiting example of a reaction scheme for the preparation of various polyamino amino acid polymer conjugates. The illustrated reaction scheme shows reaction steps for conjugating a plurality of drugs to poly-( $\gamma$ -glutamyl-glutamine). Various forms of the polyamino amino acid conjugate may be used in the reaction scheme illustrated by Figure 4. For example, alkali salts or ammonium salts of poly-( $\gamma$ -glutamyl-glutamine) may be used as the polymer conjugate. In an embodiment, a polymer reactant comprising a recurring unit of formula (VIII) is used in the reaction scheme illustrated by Figure 4.

**[0099]** As illustrated in Figure 4, a dissolved or partially dissolved poly-( $\gamma$ -glutamyl-glutamine) (PGGA) is reacted with paclitaxel in the presence of a coupling agent to form a PGGA-paclitaxel conjugate. The dissolved or partially dissolved PGGA-paclitaxel may then further be reacted with a second drug in the presence of a coupling agent. The second drug may be doxorubicin, resulting in a PGGA-(paclitaxel)-doxorubicin conjugate. Alternatively, the second drug may be camptothecin, resulting in a PGGA-(paclitaxel)-camptothecin conjugate. In an embodiment that is shown in Figure 4, PGGA-(paclitaxel)-camptothecin is dissolved or partially dissolved and reacted with a third drug, doxorubicin, to form a PGGA-(paclitaxel)-(camptothecin)-doxorubicin conjugate.

**[0100]** While Figure 4 displays embodiments where the order of drug conjugation is described, the order should not be construed as limiting. The first drug conjugated to the polymer can be any one of a taxane, a camptotheca, or an anthracycline, as described herein. Additionally, the second drug conjugated to the polymer can be any one of a taxane, a camptotheca, or an anthracycline, as described herein. Furthermore, the third drug conjugated to the polymer can be any one of a taxane, a camptotheca, or an anthracycline, as described herein. Furthermore, any one of the first, second, and/or third drugs may be conjugated to the polymer at about the same time as any one of the other first, second, and/or third drugs.

**[0101]** In an embodiment, the method of making the polymer conjugate further can include reacting the dissolved or partially dissolved polymeric reactant with a fourth reactant, wherein the fourth reactant comprises at least one selected from a polydentate ligand, a polydentate ligand precursor with protected oxygen atoms, a group that comprises a third drug, a group that comprises a targeting agent, a group that comprises an optical imaging agent, a group that comprises a magnetic resonance imaging agent, and a group that comprises a stabilizing agent. In an embodiment, the fourth reactant may further include a substituent. The substituent may be selected from a hydroxy and an amine.

**[0102]** In an embodiment, the fourth reactant can include a compound that comprises an agent. The agent may be any active compound. For instance, the compound that comprises the agent may be selected from a compound that comprises a drug, a compound that comprises a targeting agent, a compound that comprises an optical imaging agent, a compound that comprises a magnetic resonance imaging agent, and a compound that comprises stabilizing agent.

**[0103]** In some embodiments, the fourth reactant can include a compound that includes a third drug such as an anticancer drug. In an embodiment, the anticancer drug can be selected from a taxane, camptotheca, and anthracycline. In an embodiment, the taxane may be selected from paclitaxel and docetaxel. Paclitaxel may be conjugated to the polymer in a number of ways. In an embodiment, paclitaxel can be conjugated to the recurring unit of formula (I) at the oxygen atom attached to the C2'-carbon. In another embodiment, paclitaxel can be conjugated to the recurring unit of formula (I) at the oxygen atom attached to the C7-carbon. In an embodiment, the camptotheca can be camptothecin. In an



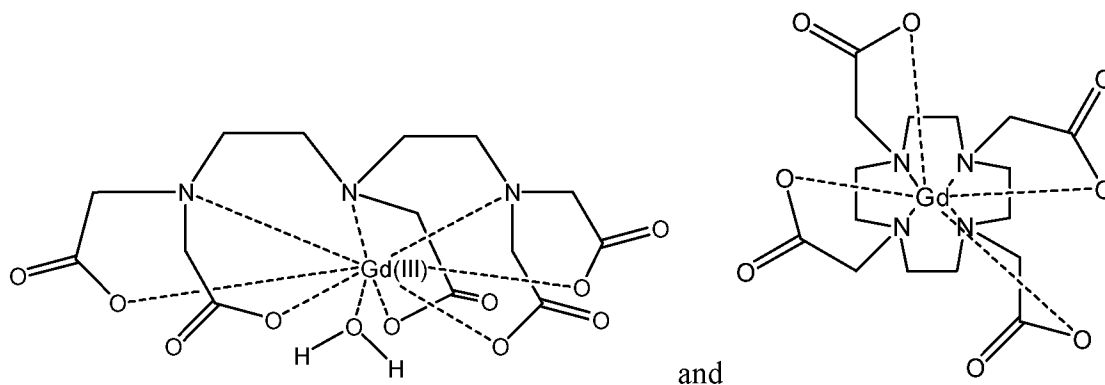
embodiment, the anthracycline can be doxorubicin. In an embodiment, the third drug can be different from the first drug and the second drug.

**[0104]** In an embodiment, the fourth reactant can include a group that comprises a targeting agent. In an embodiment, the targeting agent can be selected from an arginine-glycine-aspartate (RGD) peptide, fibronectin, folate, galactose, an apolipoprotein, insulin, transferrin, a fibroblast growth factor (FGF), an epidermal growth factor (EGF), and an antibody. In an embodiment, the targeting agent can interact with a receptor selected from  $\alpha_v\beta_3$ -integrin, folate, asialoglycoprotein, a low-density lipoprotein (LDL), an insulin receptor, a transferrin receptor, a fibroblast growth factor (FGF) receptor, an epidermal growth factor (EGF) receptor, and an antibody receptor. In some embodiments, the arginine-glycine-aspartate (RGD) peptide can be cyclic (fKRGD).

**[0105]** In an embodiment, the fourth reactant can include a group that comprises an optical imaging agent. In an embodiment, the optical imaging agent may be selected from an acridine dye, a coumarine dye, a rhodamine dye, a xanthene dye, a cyanine dye, and a pyrene dye.

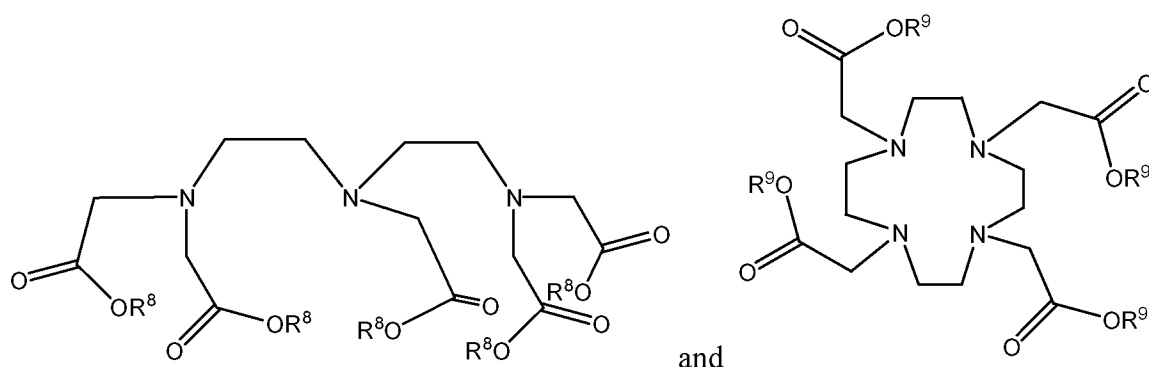
**[0106]** In an embodiment, the fourth reactant can include a group that comprises a stabilizing agent. In an embodiment, the stabilizing agent can be polyethylene glycol.

**[0107]** In an embodiment, the fourth reactant can include a group that comprises a magnetic resonance imaging agent. In an embodiment, the magnetic resonance imaging agent can include a paramagnetic metal compound. Preferably, the compound that comprises the agent comprises a Gd(III) compound. Exemplary Gd(III) compounds include the following:



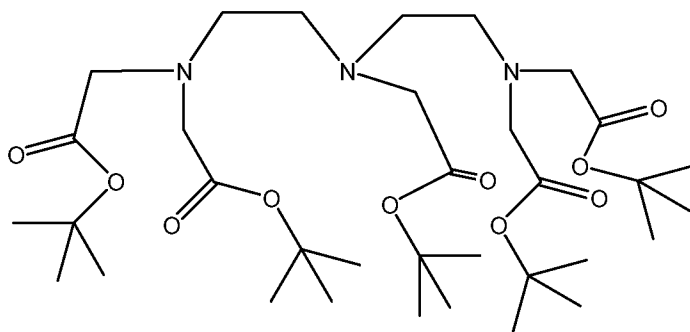
**[0108]** In an embodiment, the fourth reactant can include a polydentate ligand. Any suitable polydentate ligand may be used. In an embodiment, the polydentate ligand may

be capable of reaction with a paramagnetic metal to form a magnetic resonance imaging agent. For example, the polydentate ligand may comprise several carboxylic acid and/or carboxylate groups. For example, polydentate ligands of the following structures may be conjugated to the polymer:



wherein each  $R^8$  and each  $R^9$  can be independently hydrogen, ammonium, or an alkali metal.

**[0109]** In an embodiment, the fourth reactant can include a polydentate ligand precursor. In another embodiment, a polydentate ligand precursor having protecting groups may be conjugated to the polymer. Such a precursor has its oxygen atoms protected by a suitable protecting group(s). Suitable protecting groups include, but are not limited to, lower alkyls, benzyls, and silyl groups. One example of a polydentate ligand precursor having protecting groups is provided as follows:



**[0110]** As previously mentioned, in some embodiments, the dissolved or partially dissolved polymer reactant can be reacted with at least a portion of the second reactant before reacting with the third reactant. In an embodiment, the intermediate compound that forms after the addition of at least a portion of the second reactant can be isolated before adding the third reactant. In another embodiment, the third reactant can be added without isolating the intermediate compound that forms after the addition of the second reactant. In other

embodiments, the dissolved or partially dissolved polymer reactant is reacted with at least a portion of the second reactant at about the same time as reacting with the third reactant. In an embodiment, the dissolved or partially dissolved polymer reactant is reacted with at least a portion of the second reactant after reacting with at least a portion of the third reactant.

**[0111]** In an embodiment, the dissolved or partially dissolved polymer reactant can be reacted with at least a portion of the second reactant and/or at least a portion of the third reactant before reacting with at least a portion of a fourth reactant. In an embodiment, the dissolved or partially dissolved polymer reactant is reacted with at least a portion of a fourth reactant before reacting with at least a portion of the second reactant and/or at least a portion of the third reactant. In an embodiment, the dissolved or partially dissolved polymer reactant is reacted with at least a portion of the fourth reactant at about the same time it is reacted with at least a portion of the second reactant and/or at least a portion of the third reactant.

**[0112]** In an embodiment, a method of making the polymer conjugate can include reacting the dissolved or partially dissolved polymeric reactant with the second reactant and/or third reactant in the presence of a coupling agent. A coupling reagent may also be present for reaction with the fourth reactant. Any suitable coupling agent may be used. In an embodiment, the coupling agent can be selected from 1-ethyl-3-(3-dimethylaminopropyl)-carbodiimide (EDC), 1,3-dicyclohexyl carbodiimide (DCC), 1,1'-carbonyl-diimidazole (CDI), N,N'-disuccinimidyl carbonate (DSC), N-[(dimethylamino)-1H-1,2,3-triazolo-[4,5-b]pyridine-1-yl-methylene]-N-methylmethanaminium hexafluorophosphate N-oxide (HATU), 2-[(1H-benzotriazol-1-yl)-1,1,3,3-tetramethylaminium hexafluorophosphate (HBTU), 2-[(6-chloro-1H-benzotriazol-1-yl)-1,1,3,3-tetramethylaminium hexafluorophosphate (HCTU), benzotriazole-1-yl-oxy-tris-pyrrolidino-phosphonium hexafluorophosphate (PyBOP®), bromo-tris-pyrrolidino-phosphonium hexafluorophosphate (PyBroP®), 2-[(1H-benzotriazol-1-yl)-1,1,3,3-tetramethylaminium tetrafluoroborate (TBTU), and benzotriazol-1-yl-oxy-tris-(dimethylamino)phosphonium hexafluorophosphate (BOP).

**[0113]** Any suitable solvent that allows the reaction to take place may be used. In an embodiment, the solvent may be a polar aprotic solvent. For instance, the solvent may be

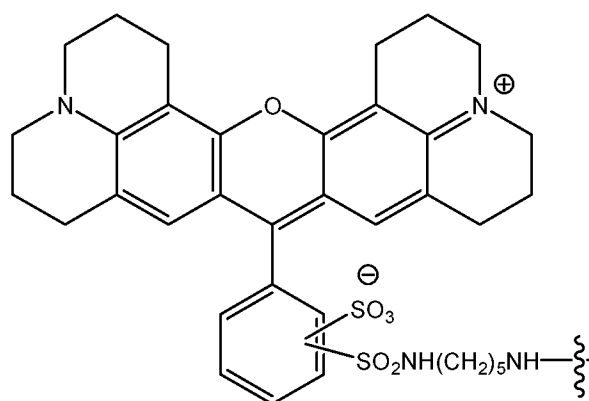
selected from N,N-dimethylformamide (DMF), dimethyl sulfoxide (DMSO), N-methyl-2-pyrindone (NMP), and N,N-dimethylacetamide (DMAc).

**[0114]** In another embodiment, the reaction may further include reacting the dissolved or partially dissolved polymeric reactant in the presence of a catalyst. Any catalyst that promotes the reaction may be used. In an embodiment, the catalyst may comprise 4-dimethylaminopyridine (DMAP).

**[0115]** Conjugation of a group that comprises a drug, a group that comprises a targeting agent, a group that comprises an optical imaging agent, a group that comprises a magnetic resonance imaging agent, a group that comprises a polydentate ligand, a group that comprises a polydentate ligand precursor and/or a group that comprises a stabilizing agent to the polymer acid or its salt form may be carried out in various ways, e.g., by covalently bonding the group comprising an agent, a polydentate ligand, and/or a polydentate ligand precursor with protected oxygen atoms to various polymers. One method for conjugating the aforementioned groups to the polymer is by using heat (e.g., heat from using a microwave method). Alternatively, conjugation may take place at room temperature. Appropriate solvents, coupling agents, catalysts, and/or buffers as generally known to those skilled in the art and/or as described herein may be used to form the polymer conjugate. As with polyglutamic acid, both the salt or acid form of the polymer obtained from polyglutamic acid and/or salt and an amino acid can be used as starting material for forming the polymer conjugate.

**[0116]** Suitable agents that can be conjugated to the polymers described herein include but are not limited to drugs, optical agents, targeting agents, magnetic resonance imaging agents (e.g., paramagnetic metal compounds), stabilizing agents, polydentate ligands, and polydentate ligand precursors with protected oxygen atoms.

**[0117]** As an example, in an embodiment, the polymer can be conjugated to an optical imaging agent such as those described herein. In an embodiment, the optical agent can be Texas Red-NH<sub>2</sub>.



Texas Red-NH--

**[0118]** In one particular embodiment, a suitable polymer reactant capable of forming a polymer comprising at least one recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) (e.g., a polymer obtained from polyglutamic acid and/or salt and an amino acid) may be reacted with DCC, Texas Red-NH<sub>2</sub> dye, pyridine, and 4-dimethylaminopyridine. The mixture can be heated using a microwave method. In an embodiment, the reaction can be heated up to a temperature in the range of about 100° to about 150°C. In another embodiment, the time the materials are heated ranges from about 5 to about 40 minutes. If desired, the reaction mixture can be cooled to room temperature. Suitable methods known to those skilled in the art can be used to isolate and/or purify the polymer conjugate. For instance, reaction mixture can be filtered into an acidic water solution. Any precipitate that forms can then be filtered and washed with water. Optionally, the precipitate can be purified by any suitable method. For example, the precipitate can be transferred into acetone and dissolved, and the resulting solution can be filtered again into a sodium bicarbonate solution. If desired, the resulting reaction solution can be dialyzed in water using a cellulose membrane and the polymer can be lyophilized and isolated.

**[0119]** In an embodiment, a suitable polymer reactant capable of forming the polymer comprising at least one recurring unit of the formulae (I), (II), (III), (IV), (V), and/or (VI) can be conjugated to a drug (e.g., an anticancer drug). In an embodiment, the anticancer drug can be a taxane, camptotheca, and/or anthracycline. In an embodiment, the anticancer drug can be a taxane such as paclitaxel or docetaxel. In other embodiments, the anticancer drug can be a camptotheca such as camptothecin. In yet still other embodiments, the anticancer drug can be an anthracycline such as doxorubicin. In some embodiments, the

anticancer drug conjugated to the polymer can be doxorubicin. In other embodiments, the anticancer drug conjugated to the polymer can be paclitaxel. In an embodiment, paclitaxel may be joined to the polymer at the C2'-oxygen atom. In another embodiment, the paclitaxel may be joined to the polymer at the C7-oxygen atom. In yet another embodiment, the polymer can include both C2'-conjugated paclitaxel groups and C7-conjugated paclitaxel groups.

**[0120]** The anti-cancer drug can be conjugated to the suitable polymer reactant using the methods described above with respect to Texas-Red.

**[0121]** In an embodiment, paclitaxel, preferably in the presence of a coupling agent (e.g, EDC and/or DCC) and a catalyst (e.g, DMAP), can be reacted with a suitable polymer reactant capable of forming a polymer comprising at least one recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) in a solvent (e.g, an aprotic solvent such as DMF). Additional agents, such as pyridine or hydroxybenzotriazole may be used. In one embodiment, the reaction may take place over the period of 0.5-2 days. Suitable methods known to those skilled in the art can be used to isolate and/or purify the polymer conjugate. For example, the reaction mixture can be poured into an acidic solution to form a precipitate. Any precipitate that forms can then be filtered and washed with water. Optionally, the precipitate can be purified by any suitable method. For example, the precipitate can be transferred into acetone and dissolved, and the resulting solution can be filtered again into a sodium bicarbonate solution. If desired, the resulting reaction solution can be dialyzed in water using a cellulose membrane and the polymer can be lyophilized and isolated. The content of paclitaxel in the resulting polymer may be determined by UV spectrometry.

**[0122]** In some embodiments, the compound comprising the agent can be reacted with an amino acid such as glutamic and/or aspartic acid in which the compound comprising the agent is coupled (e.g., covalently bonded) to the amino acid. The amino acid-agent compound can then be reacted with polyglutamic acid or its salt to form one of the polymer conjugates described herein. In one embodiment, paclitaxel can be reacted with glutamic acid to form a compound in which the paclitaxel is covalently bonded to the pendant carboxylic acid group of the glutamic acid. The glutamic acid-paclitaxel compound can then be reacted with polyglutamic acid or its salt to form one of the polymer conjugates described herein. In one embodiment, paclitaxel can be reacted with aspartic acid to form a compound in which

the paclitaxel is covalently bonded to the pendant carboxylic acid group of the aspartic acid. The aspartic acid-paclitaxel compound can then be reacted with polyglutamic acid or its salt to form the polymer conjugate. If desired, the paclitaxel coupled to the amino acid by the C2'-oxygen can be separated from the paclitaxel coupled to the amino acid by the C7-oxygen using known separation methods (e.g., HPLC).

**[0123]** After formation of the polymer conjugate, any free amount of agent not covalently bonded to the polymer may also be measured. For example, thin layer chromatography (TLC) may be used to confirm the substantial absence of free paclitaxel remaining in the compositions of polymers conjugated to paclitaxel.

**[0124]** In one embodiment, a suitable polymer reactant capable of forming a polymer comprising at least one recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) can be conjugated to a polydentate ligand. Suitable polydentate ligands include but are not limited to diethylenetriaminepentacetic acid (DTPA), tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA), (1,2-ethanediyldinitrilo)tetraacetate (EDTA), ethylenediamine, 2,2'-bipyridine (bipy), 1,10-phenanthroline (phen), 1,2-bis(diphenylphosphino)ethane (DPPE), 2,4-pentanedione (acac), and ethanedioate (ox). Appropriate solvents, coupling agents, catalysts, and/or buffers as generally known to those skilled in the art and/or described herein may be used to form the polymer conjugate. In another embodiment, the polymer comprising at least one recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) can be conjugated to a polydentate ligand precursor with protected oxygen atoms. As with polyglutamic acid, both the salt or acid form of the polymer obtained from polyglutamic acid and/or salt and an amino acid can be used as starting material for forming the polymer conjugate.

**[0125]** In an embodiment, the polydentate ligand can be DTPA. In another embodiment, the polydentate ligand can be DOTA. In one embodiment, the polydentate ligand such as DTPA (with or without protected oxygen atoms), preferably in the presence of a coupling agent (e.g., DCC) and a catalyst (e.g., DMAP), can be reacted in a solvent (e.g., an aprotic solvent such as DMF). If protecting groups are present, removal can be achieved using suitable methods. For example, the polymer conjugate with the polydentate ligand precursor with protected oxygen atoms such as DTPA with oxygen atoms protected by t-butyl groups can be treated with acid such as trifluoroacetic acid. After removal of the protecting groups,

the acid can be removed by rotary evaporation. In one embodiment, DTPA can be treated with a suitable base to remove the hydrogen atoms on the carboxylic acid –OH groups. In some embodiments, the base is sodium bicarbonate.

**[0126]** In one embodiment, a suitable polymer reactant capable of forming a polymer comprising at least one recurring unit of the formulae (I), (II), (III), (IV), (V), and/or (VI) can be conjugated to a targeting agent. Exemplary targeting agents include, but are not limited to, arginine-glycine-aspartate (RGD) peptides, fibronectin, folate, galactose, apolipoprotein, insulin, transferrin, fibroblast growth factors (FGF), epidermal growth factors (EGF), and antibodies. Targeting agents can be chosen such that they interact with particular receptors. For example, a targeting agent can be chosen so that it interacts with one or more of the following receptors:  $\alpha_v\beta_3$ -integrin, folate, asialoglycoprotein, a low-density lipoprotein (LDL), an insulin receptor, a transferrin receptor, a fibroblast growth factor (FGF) receptor, an epidermal growth factor (EGF) receptor, and an antibody receptor. In one embodiment, the arginine-glycine-aspartate (RGD) peptide is cyclic(fKRGD).

**[0127]** Both the salt or acid form of the polymer reactant capable of forming a polymer comprising at least one recurring unit of the formulae (I), (II), (III), (IV), (V), and/or (VI) can be used as starting material for forming the polymer conjugate with a targeting agent. In one embodiment, the targeting agent preferably in the presence of a coupling agent (e.g., DCC) and a catalyst (e.g., DMAP), can be reacted with the polymer obtained from polyglutamic acid and/or salt and an amino acid in a solvent (e.g., an aprotic solvent such as DMF). After formation of the polymer conjugate, any free amount of agent not covalently bonded to the polymer may also be measured. For example, thin layer chromatography (TLC) may be used to confirm the substantial absence of any free targeting agent. Suitable methods known to those skilled in the art can be used to isolate and/or purify the polymer conjugate (e.g., lyophilization).

**[0128]** In an embodiment, a suitable polymer reactant capable of forming a polymer comprising at least one recurring unit of the formulae (I), (II), (III), (IV), (V), and/or (VI) can be conjugated to a magnetic resonance imaging agent. In an embodiment, the magnetic resonance imaging agent can comprise a Gd(III) compound. One method for forming the magnetic resonance imaging agent is by reacting a paramagnetic metal with the polymer conjugate comprising a polydentate ligand. Suitable paramagnetic metals include



but are not limited to Gd(III), Indium-111, and Yttrium-88. For example, a polymer conjugate comprising DTPA can be treated with Gd(III) in a buffer solution for a period of several hours. Suitable methods known to those skilled in the art can be used to isolate and/or purify the polymer conjugate. For instance, the resulting reaction solution can be dialyzed in water using a cellulose membrane and the polymer can be lyophilized and isolated. The amount of paramagnetic metal may be quantified by inductively coupled plasma-optical emission spectroscopy (ICP-OES) measurement.

**[0129]** In one embodiment, a suitable polymer reactant capable of forming a polymer comprising at least one recurring unit of the formulae (I), (II), (III), (IV), (V), and/or (VI) can be conjugated to a stabilizing agent. In some embodiments, the stabilizing agent can be polyethylene glycol. In one method, the stabilizing agent, preferably in the presence of a coupling agent (e.g., DCC) and a catalyst (e.g., DMAP), can be reacted with the polymer obtained from polyglutamic acid and/or salt and an amino acid in a solvent (e.g., an aprotic solvent such as DMF). Progress of the reaction can be measured by any suitable method such as TLC. The resulting polymer conjugate can be purified using methods known to those skilled in the art such as dialysis.

**[0130]** The polymer conjugates may be used to deliver an imaging agent, targeting agent, magnetic resonance imaging agent and/or a drug to a selected tissue. For example, polymer conjugates comprising the Texas Red dye may be used to deliver an imaging agent to a selected tissue. In one embodiment, the polymer conjugates comprising at least one recurring unit of the formulae (I), (II), (III), (IV), (V), and/or (VI) can be used to treat or ameliorate a disease or condition such as cancer. In an embodiment, the polymer conjugates described herein can be used to diagnose a disease or condition (e.g., cancer). In yet one more embodiment, the polymer conjugates described herein can be used to image a portion of tissue. In some embodiments, the disease or condition can be a cancer such as lung cancer, breast cancer, colon cancer, ovarian cancer, prostate cancer, and melanoma. In an embodiment, the disease or condition can be a tumor selected from lung tumor, breast tumor, colon tumor, ovarian tumor, prostate tumor, and melanoma tumor. In some embodiments, the tissue being imaged can be tissue from lung tumor, breast tumor, colon tumor, ovarian tumor, prostate tumor, and/or melanoma tumor.

**[0131]** The polymers described above may be formed into nanoparticles in aqueous solution. Conjugates comprising a polymer and a drug may be formed into nanoparticles in a similar manner. Such nanoparticles may be used to preferentially deliver a drug to a selected tissue.

**[0132]** An embodiment provides a composition comprising one or more polymer conjugates described herein and at least one selected from a pharmaceutically acceptable excipient, a carrier, and a diluent. In some embodiments, prodrugs, metabolites, stereoisomers, hydrates, solvates, polymorphs, and pharmaceutically acceptable salts of the compounds disclosed herein (e.g., the polymer conjugate and/or the agent(s) that it comprises) are provided.

**[0133]** A “prodrug” refers to an agent that is converted into the parent drug *in vivo*. Prodrugs are often useful because, in some situations, they may be easier to administer than the parent drug. They may, for instance, be bioavailable by oral administration whereas the parent is not. The prodrug may also have improved solubility in pharmaceutical compositions over the parent drug. An example, without limitation, of a prodrug would be a compound which is administered as an ester (the “prodrug”) to facilitate transmittal across a cell membrane where water solubility is detrimental to mobility but which then is metabolically hydrolyzed to the carboxylic acid, the active entity, once inside the cell where water-solubility is beneficial. A further example of a prodrug might be a short peptide (polyaminoacid) bonded to an acid group where the peptide is metabolized to reveal the active moiety. Conventional procedures for the selection and preparation of suitable prodrug derivatives are described, for example, in *Design of Prodrugs*, (ed. H. Bundgaard, Elsevier, 1985), which is hereby incorporated herein by reference in its entirety.

**[0134]** The term “pro-drug ester” refers to derivatives of the compounds disclosed herein formed by the addition of any of several ester-forming groups that are hydrolyzed under physiological conditions. Examples of pro-drug ester groups include pivaloyloxymethyl, acetoxymethyl, phthalidyl, indanyl and methoxymethyl, as well as other such groups known in the art, including a (5-R-2-oxo-1,3-dioxolen-4-yl)methyl group. Other examples of pro-drug ester groups can be found in, for example, T. Higuchi and V. Stella, in “Pro-drugs as Novel Delivery Systems”, Vol. 14, A.C.S. Symposium Series, American Chemical Society (1975); and “Bioreversible Carriers in Drug Design: Theory and

Application", edited by E. B. Roche, Pergamon Press: New York, 14-21 (1987) (providing examples of esters useful as prodrugs for compounds containing carboxyl groups). Each of the above-mentioned references is herein incorporated by reference in their entirety.

**[0135]** The term "pharmaceutically acceptable salt" refers to a salt of a compound that does not cause significant irritation to an organism to which it is administered and does not abrogate the biological activity and properties of the compound. In some embodiments, the salt is an acid addition salt of the compound. Pharmaceutical salts can be obtained by reacting a compound with inorganic acids such as hydrohalic acid (e.g., hydrochloric acid or hydrobromic acid), sulfuric acid, nitric acid, phosphoric acid and the like. Pharmaceutical salts can also be obtained by reacting a compound with an organic acid such as aliphatic or aromatic carboxylic or sulfonic acids, for example acetic, succinic, lactic, malic, tartaric, citric, ascorbic, nicotinic, methanesulfonic, ethanesulfonic, p-toluenesulfonic, salicylic or naphthalenesulfonic acid. Pharmaceutical salts can also be obtained by reacting a compound with a base to form a salt such as an ammonium salt, an alkali metal salt, such as a sodium or a potassium salt, an alkaline earth metal salt, such as a calcium or a magnesium salt, a salt of organic bases such as dicyclohexylamine, N-methyl-D-glucamine, tris(hydroxymethyl)methylamine, C<sub>1</sub>-C<sub>7</sub> alkylamine, cyclohexylamine, triethanolamine, ethylenediamine, and salts with amino acids such as arginine, lysine, and the like.

**[0136]** If the manufacture of pharmaceutical formulations involves intimate mixing of the pharmaceutical excipients and the active ingredient in its salt form, then it may be desirable to use pharmaceutical excipients which are non-basic, that is, either acidic or neutral excipients.

**[0137]** In various embodiments, the compounds disclosed herein (e.g., the polymer conjugate and/or the agent(s) that it comprises) can be used alone, in combination with other compounds disclosed herein, or in combination with one or more other agents active in the therapeutic areas described herein.

**[0138]** In another aspect, the present disclosure relates to a pharmaceutical composition comprising one or more physiologically acceptable surface active agents, carriers, diluents, excipients, smoothing agents, suspension agents, film forming substances, and coating assistants, or a combination thereof; and a compound (e.g., the polymer conjugate and/or the agent(s) that it comprises) disclosed herein. Acceptable carriers or

diluents for therapeutic use are well known in the pharmaceutical art, and are described, for example, in Remington's Pharmaceutical Sciences, 18th Ed., Mack Publishing Co., Easton, PA (1990), which is incorporated herein by reference in its entirety. Preservatives, stabilizers, dyes, sweeteners, fragrances, flavoring agents, and the like may be provided in the pharmaceutical composition. For example, sodium benzoate, ascorbic acid and esters of p-hydroxybenzoic acid may be added as preservatives. In addition, antioxidants and suspending agents may be used. In various embodiments, alcohols, esters, sulfated aliphatic alcohols, and the like may be used as surface active agents; sucrose, glucose, lactose, starch, crystallized cellulose, mannitol, light anhydrous silicate, magnesium aluminate, magnesium metasilicate aluminate, synthetic aluminum silicate, calcium carbonate, sodium acid carbonate, calcium hydrogen phosphate, calcium carboxymethyl cellulose, and the like may be used as excipients; magnesium stearate, talc, hardened oil and the like may be used as smoothing agents; coconut oil, olive oil, sesame oil, peanut oil, soya may be used as suspension agents or lubricants; cellulose acetate phthalate as a derivative of a carbohydrate such as cellulose or sugar, or methylacetate-methacrylate copolymer as a derivative of polyvinyl may be used as suspension agents; and plasticizers such as ester phthalates and the like may be used as suspension agents.

**[0139]** The term “pharmaceutical composition” refers to a mixture of a compound disclosed herein (e.g., the polymer conjugate and/or the agent(s) that it comprises) with other chemical components, such as diluents or carriers. The pharmaceutical composition facilitates administration of the compound to an organism. Multiple techniques of administering a compound exist in the art including, but not limited to, oral, injection, aerosol, parenteral, and topical administration. Pharmaceutical compositions can also be obtained by reacting compounds with inorganic or organic acids such as hydrochloric acid, hydrobromic acid, sulfuric acid, nitric acid, phosphoric acid, methanesulfonic acid, ethanesulfonic acid, p-toluenesulfonic acid, salicylic acid and the like.

**[0140]** The term “carrier” refers to a chemical compound that facilitates the incorporation of a compound into cells or tissues. For example dimethyl sulfoxide (DMSO) is a commonly utilized carrier as it facilitates the uptake of many organic compounds into the cells or tissues of an organism.

**[0141]** The term “diluent” refers to chemical compounds diluted in water that will dissolve the compound of interest (e.g., the polymer conjugate and/or the agent(s) that it comprises) as well as stabilize the biologically active form of the compound. Salts dissolved in buffered solutions are utilized as diluents in the art. One commonly used buffered solution is phosphate buffered saline because it mimics the salt conditions of human blood. Since buffer salts can control the pH of a solution at low concentrations, a buffered diluent rarely modifies the biological activity of a compound. The term “physiologically acceptable” refers to a carrier or diluent that does not abrogate the biological activity and properties of the compound.

**[0142]** The pharmaceutical compositions described herein can be administered to a human patient *per se*, or in pharmaceutical compositions where they are mixed with other active ingredients, as in combination therapy, or suitable carriers or excipient(s). Techniques for formulation and administration of the compounds of the instant application may be found in “Remington’s Pharmaceutical Sciences,” Mack Publishing Co., Easton, PA, 18th edition, 1990.

**[0143]** Suitable routes of administration may, for example, include oral, rectal, transmucosal, topical, or intestinal administration; parenteral delivery, including intramuscular, subcutaneous, intravenous, intramedullary injections, as well as intrathecal, direct intraventricular, intraperitoneal, intranasal, or intraocular injections. The compounds (e.g., the polymer conjugate and/or the agent(s) that it comprises) can also be administered in sustained or controlled release dosage forms, including depot injections, osmotic pumps, pills, transdermal (including electrotransport) patches, and the like, for prolonged and/or timed, pulsed administration at a predetermined rate.

**[0144]** The pharmaceutical compositions of the present invention may be manufactured in a manner that is itself known, e.g., by means of conventional mixing, dissolving, granulating, dragee-making, levigating, emulsifying, encapsulating, entrapping or tableting processes.

**[0145]** Pharmaceutical compositions for use in accordance with the present invention thus may be formulated in conventional manner using one or more physiologically acceptable carriers comprising excipients and auxiliaries which facilitate processing of the active compounds into preparations which can be used pharmaceutically. Proper formulation

is dependent upon the route of administration chosen. Any of the well-known techniques, carriers, and excipients may be used as suitable and as understood in the art; *e.g.*, in Remington's Pharmaceutical Sciences, above.

[0146] Injectables can be prepared in conventional forms, either as liquid solutions or suspensions, solid forms suitable for solution or suspension in liquid prior to injection, or as emulsions. Suitable excipients are, for example, water, saline, dextrose, mannitol, lactose, lecithin, albumin, sodium glutamate, cysteine hydrochloride, and the like. In addition, if desired, the injectable pharmaceutical compositions may contain minor amounts of nontoxic auxiliary substances, such as wetting agents, pH buffering agents, and the like. Physiologically compatible buffers include, but are not limited to, Hanks's solution, Ringer's solution, or physiological saline buffer. If desired, absorption enhancing preparations (for example, liposomes), may be utilized.

[0147] For transmucosal administration, penetrants appropriate to the barrier to be permeated may be used in the formulation.

[0148] Pharmaceutical formulations for parenteral administration, *e.g.*, by bolus injection or continuous infusion, include aqueous solutions of the active compounds (*e.g.*, the polymer conjugate and/or the agent(s) that it comprises) in water-soluble form. Additionally, suspensions of the active compounds may be prepared as appropriate oily injection suspensions. Suitable lipophilic solvents or vehicles include fatty oils such as sesame oil, or other organic oils such as soybean, grapefruit or almond oils, or synthetic fatty acid esters, such as ethyl oleate or triglycerides, or liposomes. Aqueous injection suspensions may contain substances which increase the viscosity of the suspension, such as sodium carboxymethyl cellulose, sorbitol, or dextran. Optionally, the suspension may also contain suitable stabilizers or agents that increase the solubility of the compounds to allow for the preparation of highly concentrated solutions. Formulations for injection may be presented in unit dosage form, *e.g.*, in ampoules or in multi-dose containers, with an added preservative. The compositions may take such forms as suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as suspending, stabilizing and/or dispersing agents. Alternatively, the active ingredient may be in powder form for constitution with a suitable vehicle, *e.g.*, sterile pyrogen-free water, before use.

**[0149]** For oral administration, the compounds (e.g., the polymer conjugate and/or the agent(s) that it comprises) can be formulated readily by combining the active compounds with pharmaceutically acceptable carriers well known in the art. Such carriers enable the compounds of the invention to be formulated as tablets, pills, dragees, capsules, liquids, gels, syrups, slurries, suspensions and the like, for oral ingestion by a patient to be treated. Pharmaceutical preparations for oral use can be obtained by combining the active compounds with solid excipient, optionally grinding a resulting mixture, and processing the mixture of granules, after adding suitable auxiliaries, if desired, to obtain tablets or dragee cores. Suitable excipients are, in particular, fillers such as sugars, including lactose, sucrose, mannitol, or sorbitol; cellulose preparations such as, for example, maize starch, wheat starch, rice starch, potato starch, gelatin, gum tragacanth, methyl cellulose, hydroxypropylmethylcellulose, sodium carboxymethylcellulose, and/or polyvinylpyrrolidone (PVP). If desired, disintegrating agents may be added, such as the cross-linked polyvinyl pyrrolidone, agar, or alginic acid or a salt thereof such as sodium alginate. Dragee cores are provided with suitable coatings. For this purpose, concentrated sugar solutions may be used, which may optionally contain gum arabic, talc, polyvinyl pyrrolidone, carbopol gel, polyethylene glycol, and/or titanium dioxide, lacquer solutions, and suitable organic solvents or solvent mixtures. Dyestuffs or pigments may be added to the tablets or dragee coatings for identification or to characterize different combinations of active compound doses. For this purpose, concentrated sugar solutions may be used, which may optionally contain gum arabic, talc, polyvinyl pyrrolidone, carbopol gel, polyethylene glycol, and/or titanium dioxide, lacquer solutions, and suitable organic solvents or solvent mixtures. Dyestuffs or pigments may be added to the tablets or dragee coatings for identification or to characterize different combinations of active compound doses.

**[0150]** Pharmaceutical preparations which can be used orally include push-fit capsules made of gelatin, as well as soft, sealed capsules made of gelatin and a plasticizer, such as glycerol or sorbitol. The push-fit capsules can contain the active ingredients (e.g., the polymer conjugate and/or the agent(s) that it comprises) in admixture with filler such as lactose, binders such as starches, and/or lubricants such as talc or magnesium stearate and, optionally, stabilizers. In soft capsules, the active compounds may be dissolved or suspended in suitable liquids, such as fatty oils, liquid paraffin, or liquid polyethylene glycols. In

addition, stabilizers may be added. All formulations for oral administration should be in dosages suitable for such administration.

**[0151]** For buccal administration, the compositions (e.g., the polymer conjugate and/or the agent(s) that it comprises) may take the form of tablets or lozenges formulated in conventional manner.

**[0152]** For administration by inhalation, the compounds (e.g., the polymer conjugate and/or the agent(s) that it comprises) for use according to the present invention are conveniently delivered in the form of an aerosol spray presentation from pressurized packs or a nebulizer, with the use of a suitable propellant, *e.g.*, dichlorodifluoromethane, trichlorofluoromethane, dichlorotetrafluoroethane, carbon dioxide or other suitable gas. In the case of a pressurized aerosol the dosage unit may be determined by providing a valve to deliver a metered amount. Capsules and cartridges of, *e.g.*, gelatin for use in an inhaler or insufflator may be formulated containing a powder mix of the compound and a suitable powder base such as lactose or starch.

**[0153]** Further disclosed herein are various pharmaceutical compositions well known in the pharmaceutical art for uses that include intraocular, intranasal, and intraauricular delivery. Suitable penetrants for these uses are generally known in the art. Pharmaceutical compositions for intraocular delivery include aqueous ophthalmic solutions of the active compounds (e.g., the polymer conjugate and/or the agent(s) that it comprises) in water-soluble form, such as eyedrops, or in gellan gum (Shedden et al., Clin. Ther., 23(3):440-50 (2001)) or hydrogels (Mayer et al., Ophthalmologica, 210(2):101-3 (1996)); ophthalmic ointments; ophthalmic suspensions, such as microparticulates, drug-containing small polymeric particles that are suspended in a liquid carrier medium (Joshi, A., J. Ocul. Pharmacol., 10(1):29-45 (1994)), lipid-soluble formulations (Alm et al., Prog. Clin. Biol. Res., 312:447-58 (1989)), and microspheres (Mordenti, Toxicol. Sci., 52(1):101-6 (1999)); and ocular inserts. All of the above-mentioned references, are incorporated herein by reference in their entireties. Such suitable pharmaceutical formulations are most often and preferably formulated to be sterile, isotonic and buffered for stability and comfort. Pharmaceutical compositions for intranasal delivery may also include drops and sprays often prepared to simulate in many respects nasal secretions to ensure maintenance of normal ciliary action. As disclosed in Remington's Pharmaceutical Sciences, 18th Ed., Mack



Publishing Co., Easton, PA (1990), which is incorporated herein by reference in its entirety, and well-known to those skilled in the art, suitable formulations are most often and preferably isotonic, slightly buffered to maintain a pH of 5.5 to 6.5, and most often and preferably include antimicrobial preservatives and appropriate drug stabilizers. Pharmaceutical formulations for intraauricular delivery include suspensions and ointments for topical application in the ear. Common solvents for such aural formulations include glycerin and water.

**[0154]** The compounds (e.g., the polymer conjugate and/or the agent(s) that it comprises) may also be formulated in rectal compositions such as suppositories or retention enemas, *e.g.*, containing conventional suppository bases such as cocoa butter or other glycerides.

**[0155]** In addition to the formulations described previously, the compounds (e.g., the polymer conjugate and/or the agent(s) that it comprises) may also be formulated as a depot preparation. Such long acting formulations may be administered by implantation (for example subcutaneously or intramuscularly) or by intramuscular injection. Thus, for example, the compounds may be formulated with suitable polymeric or hydrophobic materials (for example as an emulsion in an acceptable oil) or ion exchange resins, or as sparingly soluble derivatives, for example, as a sparingly soluble salt.

**[0156]** For hydrophobic compounds, a suitable pharmaceutical carrier may be a cosolvent system comprising benzyl alcohol, a nonpolar surfactant, a water-miscible organic polymer, and an aqueous phase. A common cosolvent system used is the VPD co-solvent system, which is a solution of 3% w/v benzyl alcohol, 8% w/v of the nonpolar surfactant Polysorbate 80™, and 65% w/v polyethylene glycol 300, made up to volume in absolute ethanol. Naturally, the proportions of a co-solvent system may be varied considerably without destroying its solubility and toxicity characteristics. Furthermore, the identity of the co-solvent components may be varied: for example, other low-toxicity nonpolar surfactants may be used instead of POLYSORBATE 80™; the fraction size of polyethylene glycol may be varied; other biocompatible polymers may replace polyethylene glycol, *e.g.*, polyvinyl pyrrolidone; and other sugars or polysaccharides may substitute for dextrose.

**[0157]** Alternatively, other delivery systems for hydrophobic pharmaceutical compounds may be employed. Liposomes and emulsions are well known examples of

delivery vehicles or carriers for hydrophobic drugs. Certain organic solvents such as dimethylsulfoxide also may be employed, although usually at the cost of greater toxicity. Additionally, the compounds may be delivered using a sustained-release system, such as semipermeable matrices of solid hydrophobic polymers containing the therapeutic agent. Various sustained-release materials have been established and are well known by those skilled in the art. Sustained-release capsules may, depending on their chemical nature, release the compounds for a few hours or weeks up to over 100 days. Depending on the chemical nature and the biological stability of the therapeutic reagent, additional strategies for protein stabilization may be employed.

**[0158]** Agents intended to be administered intracellularly may be administered using techniques well known to those of ordinary skill in the art. For example, such agents may be encapsulated into liposomes. All molecules present in an aqueous solution at the time of liposome formation are incorporated into the aqueous interior. The liposomal contents are both protected from the external micro-environment and, because liposomes fuse with cell membranes, are efficiently delivered into the cell cytoplasm. The liposome may be coated with a tissue-specific antibody. The liposomes will be targeted to and taken up selectively by the desired organ. Alternatively, small hydrophobic organic molecules may be directly administered intracellularly.

**[0159]** Additional therapeutic or diagnostic agents may be incorporated into the pharmaceutical compositions. Alternatively or additionally, pharmaceutical compositions may be combined with other compositions that contain other therapeutic or diagnostic agents.

**[0160]** The compounds (e.g., the polymer conjugate and/or the agent(s) that it comprises) or pharmaceutical compositions may be administered to the patient by any suitable means. Non-limiting examples of methods of administration include, among others, (a) administration through oral pathways, which administration includes administration in capsule, tablet, granule, spray, syrup, or other such forms; (b) administration through non-oral pathways such as rectal, vaginal, intraurethral, intraocular, intranasal, or intraauricular, which administration includes administration as an aqueous suspension, an oily preparation or the like or as a drip, spray, suppository, salve, ointment or the like; (c) administration via injection, subcutaneously, intraperitoneally, intravenously, intramuscularly, intradermally, intraorbitally, intracapsularly, intraspinally, intrasternally, or the like, including infusion

pump delivery; (d) administration locally such as by injection directly in the renal or cardiac area, e.g., by depot implantation; as well as (e) administration topically; as deemed appropriate by those of skill in the art for bringing the active compound into contact with living tissue.

**[0161]** Compositions suitable for administration (e.g., the polymer conjugate and/or the agent(s) that it comprises) include compositions where the active ingredients are contained in an amount effective to achieve its intended purpose. The effective amount of the compounds disclosed herein required as a dose will depend on the route of administration, the type of animal, including human, being treated, and the physical characteristics of the specific animal under consideration. The dose can be tailored to achieve a desired effect, but will depend on such factors as weight, diet, concurrent medication and other factors which those skilled in the medical arts will recognize. More specifically, an effective amount means an amount of compound effective to prevent, alleviate or ameliorate symptoms of disease or prolong the survival of the subject being treated. Determination of an effective amount is well within the capability of those skilled in the art, especially in light of the detailed disclosure provided herein.

**[0162]** As will be readily apparent to one skilled in the art, the useful *in vivo* dosage to be administered and the particular mode of administration will vary depending upon the age, weight and mammalian species treated, the particular compounds employed, and the specific use for which these compounds are employed. The determination of effective dosage levels, that is the dosage levels necessary to achieve the desired result, can be accomplished by one skilled in the art using routine pharmacological methods. Typically, human clinical applications of products are commenced at lower dosage levels, with dosage level being increased until the desired effect is achieved. Alternatively, acceptable *in vitro* studies can be used to establish useful doses and routes of administration of the compositions identified by the present methods using established pharmacological methods.

**[0163]** In non-human animal studies, applications of potential products are commenced at higher dosage levels, with dosage being decreased until the desired effect is no longer achieved or adverse side effects disappear. The dosage may range broadly, depending upon the desired effects and the therapeutic indication. Typically, dosages may be between about 10 microgram/kg and 100 mg/kg body weight, preferably between about 100

microgram/kg and 10 mg/kg body weight. Alternatively dosages may be based and calculated upon the surface area of the patient, as understood by those of skill in the art.

**[0164]** The exact formulation, route of administration and dosage for the pharmaceutical compositions of the present invention can be chosen by the individual physician in view of the patient's condition. (See *e.g.*, Fingl et al. 1975, in "The Pharmacological Basis of Therapeutics", which is hereby incorporated herein by reference in its entirety, with particular reference to Ch. 1, p. 1). Typically, the dose range of the composition administered to the patient can be from about 0.5 to 1000 mg/kg of the patient's body weight. The dosage may be a single one or a series of two or more given in the course of one or more days, as is needed by the patient. In instances where human dosages for compounds have been established for at least some condition, the present invention will use those same dosages, or dosages that are between about 0.1% and 500%, more preferably between about 25% and 250% of the established human dosage. Where no human dosage is established, as will be the case for newly-discovered pharmaceutical compositions, a suitable human dosage can be inferred from ED<sub>50</sub> or ID<sub>50</sub> values, or other appropriate values derived from *in vitro* or *in vivo* studies, as qualified by toxicity studies and efficacy studies in animals.

**[0165]** It should be noted that the attending physician would know how to and when to terminate, interrupt, or adjust administration due to toxicity or organ dysfunctions. Conversely, the attending physician would also know to adjust treatment to higher levels if the clinical response were not adequate (precluding toxicity). The magnitude of an administered dose in the management of the disorder of interest will vary with the severity of the condition to be treated and to the route of administration. The severity of the condition may, for example, be evaluated, in part, by standard prognostic evaluation methods. Further, the dose and perhaps dose frequency, will also vary according to the age, body weight, and response of the individual patient. A program comparable to that discussed above may be used in veterinary medicine.

**[0166]** Although the exact dosage will be determined on a drug-by-drug basis, in most cases, some generalizations regarding the dosage can be made. The daily dosage regimen for an adult human patient may be, for example, an oral dose of between 0.1 mg and 2000 mg of each active ingredient, preferably between 1 mg and 500 mg, *e.g.* 5 to 200 mg.

In other embodiments, an intravenous, subcutaneous, or intramuscular dose of each active ingredient of between 0.01 mg and 100 mg, preferably between 0.1 mg and 60 mg, e.g. 1 to 40 mg is used. In cases of administration of a pharmaceutically acceptable salt, dosages may be calculated as the free base. In some embodiments, the composition (e.g., the polymer conjugate and/or the agent(s) that it comprises) is administered 1 to 4 times per day. Alternatively the composition be administered by continuous intravenous infusion, preferably at a dose of each active ingredient up to 1000 mg per day. As will be understood by those of skill in the art, in certain situations it may be necessary to administer the compounds disclosed herein in amounts that exceed, or even far exceed, the above-stated, preferred dosage range in order to effectively and aggressively treat particularly aggressive diseases or infections. In some embodiments, the compounds will be administered for a period of continuous therapy, for example for a week or more, or for months or years.

[0167] Dosage amount and interval may be adjusted individually to provide plasma levels of the active moiety which are sufficient to maintain the modulating effects, or minimal effective concentration (MEC). The MEC will vary for each compound but can be estimated from *in vitro* data. Dosages necessary to achieve the MEC will depend on individual characteristics and route of administration. However, HPLC assays or bioassays can be used to determine plasma concentrations.

[0168] Dosage intervals can also be determined using MEC value. Compositions should be administered using a regimen which maintains plasma levels above the MEC for 10-90% of the time, preferably between 30-90% and most preferably between 50-90%.

[0169] In cases of local administration or selective uptake, the effective local concentration of the drug may not be related to plasma concentration.

[0170] The amount of composition administered may be dependent on the subject being treated, on the subject's weight, the severity of the affliction, the manner of administration and the judgment of the prescribing physician.

[0171] Compounds disclosed herein (e.g., the polymer conjugate and/or the agent(s) that it comprises) can be evaluated for efficacy and toxicity using known methods. For example, the toxicology of a particular compound, or of a subset of the compounds, sharing certain chemical moieties, may be established by determining *in vitro* toxicity towards a cell line, such as a mammalian, and preferably human, cell line. The results of

such studies are often predictive of toxicity in animals, such as mammals, or more specifically, humans. Alternatively, the toxicity of particular compounds in an animal model, such as mice, rats, rabbits, or monkeys, may be determined using known methods. The efficacy of a particular compound may be established using several recognized methods, such as *in vitro* methods, animal models, or human clinical trials. Recognized *in vitro* models exist for nearly every class of condition, including but not limited to cancer, cardiovascular disease, and various immune dysfunction. Similarly, acceptable animal models may be used to establish efficacy of chemicals to treat such conditions. When selecting a model to determine efficacy, the skilled artisan can be guided by the state of the art to choose an appropriate model, dose, and route of administration, and regime. Of course, human clinical trials can also be used to determine the efficacy of a compound in humans.

**[0172]** The compositions may, if desired, be presented in a pack or dispenser device which may contain one or more unit dosage forms containing the active ingredient. The pack may for example comprise metal or plastic foil, such as a blister pack. The pack or dispenser device may be accompanied by instructions for administration. The pack or dispenser may also be accompanied with a notice associated with the container in form prescribed by a governmental agency regulating the manufacture, use, or sale of pharmaceuticals, which notice is reflective of approval by the agency of the form of the drug for human or veterinary administration. Such notice, for example, may be the labeling approved by the U.S. Food and Drug Administration for prescription drugs, or the approved product insert. Compositions comprising a compound of the invention formulated in a compatible pharmaceutical carrier may also be prepared, placed in an appropriate container, and labeled for treatment of an indicated condition.

**[0173]** Polymers and copolymers comprising a recurring unit of the formula (I) may have many different uses. An embodiment provides a method of treating or ameliorating a disease or condition comprising administering an effective amount of one or more polymer conjugates described herein or the pharmaceutical composition described herein to a mammal in need thereof. Another embodiment provides a use an effective amount of one or more polymer conjugates described herein or the pharmaceutical composition described herein for treating or ameliorating a disease or condition. In an embodiment, the disease or condition is selected from lung tumor, breast tumor, colon tumor,

ovarian tumor, prostate tumor, and melanoma tumor. In an embodiment, the disease or condition is selected from lung cancer, breast cancer, colon cancer, ovarian cancer, prostate cancer, and melanoma.

[0174] An embodiment provides a method of diagnosing a disease or condition comprising administering an effective amount of one or more polymer conjugates described herein or the pharmaceutical composition described herein to a mammal in need thereof. Another embodiment provides a use an effective amount of one or more polymer conjugates described herein or the pharmaceutical composition described herein for diagnosing a disease or condition. In an embodiment, the disease or condition is selected from lung tumor, breast tumor, colon tumor, ovarian tumor, prostate tumor, and melanoma tumor. In an embodiment, the disease or condition is selected from lung cancer, breast cancer, colon cancer, ovarian cancer, prostate cancer, and melanoma.

[0175] An embodiment provides a method of imaging a portion of tissue comprising contacting a portion of tissue with an effective amount of one or more polymer conjugates described herein or the pharmaceutical composition described herein. Another embodiment provides a use an effective amount of one or more polymer conjugates described herein or the pharmaceutical composition described herein for imaging a portion of tissue. In some embodiments, the tissue being imaged can be tissue from lung tumor, breast tumor, colon tumor, ovarian tumor, prostate tumor, and/or melanoma tumor.

### EXAMPLES

[0176] The following examples are provided for the purposes of further describing the embodiments described herein, and do not limit the scope of the invention.

#### Materials:

[0177] Poly-L-glutamate sodium salts with different molecular weights (average molecular weights of 41,400 (PGA(97k)), 17,600 (PGA(44k)), 16,000 (PGA(32k)), and 10,900 (PGA(21k)) daltons based on multi-angle light scattering (MALS)); N-(3-dimethylaminopropyl)-N'-ethylcarbodiimide hydrochloride (EDC); hydroxybenzotriazole (HOBt); pyridine; 4-dimethylaminopyridine (DMAP); N,N'-dimethylformamide (DMF); gadolinium-acetate; chloroform; camptothecin, and sodium bicarbonate were purchased from

Sigma-Aldrich Chemical company. Poly-L-glutamate was converted into poly-L-glutamic acid using 2 N hydrochloric acid solution. Trifluoroacetic acid (TFA) was purchased from Bioscience. L-glutamic acid di-*t*-butyl ester hydrochloride (H-Glu(OtBu)-OtBu-HCl), N- $\alpha$ -CBZ-L-glutamic acid  $\alpha$ -benzyl ester (Z-Glu-OBzl) were purchased from Novabiochem (La Jolla, CA). Paclitaxel and doxorubicin was purchased from PolyMed (Houston, Texas). The chemical p-NH<sub>2</sub>-Bn-DPTA-penta-(*t*-Bu ester) was purchased from Macrocyclics (Dallas, Texas). <sup>1</sup>H NMR was obtained from Joel (400 MHz), and particle sizes were measured by ZetalPals (Brookhaven Instruments Corporation). Microwave chemistry was carried out in Biotage. Molecular weights of polymers were determined by size exclusion chromatography (SEC) combined with a multi-angle light scattering (MALS) (Wyatt Corporation) detector.

**[0178]** A poly-( $\gamma$ -L-glutamyl-glutamine) was prepared from a polyglutamate sodium salt, according to the procedures described in U.S. Patent Publication No. 2007-0128118, filed December 1, 2006, which is hereby incorporated by reference in its entirety, and particularly for the purpose of describing the synthesis of the polymer described therein (e.g. poly-( $\gamma$ -L-glutamyl-glutamine), poly-( $\gamma$ -L-aspartyl-glutamine), poly-( $\gamma$ -L-glutamyl-glutamine)-poly-L-glutamic acid, and poly-( $\gamma$ -L-aspartyl-glutamine)-poly-L-glutamic acid. Average molecular weights of the polymers were determined using the system and conditions described below (hereinafter, referred to as the Heleos system with MALS detector).

#### SEC-MALS Analysis Conditions:

- HPLC system: Agilent 1200
- Column: Shodex SB 806M HQ  
(exclusion limit for Pullulan is 20,000,000, particle size: 13 micron, size (mm) IDxLength; 8.0 x300)
- Mobile Phase: 1xDPBS or 1% LiBr in DPBS (pH7.0)
- Flow Rate: 1 ml/min
- MALS detector: DAWN HELEOS from Wyatt
- DRI detector: Optilab rEX from Wyatt
- On-line Viscometer: ViscoStar from Wyatt
- Software: ASTRA 5.1.9 from Wyatt
- Sample Concentration: 1-2 mg/ml
- Injection volume: 100  $\mu$ l

dn/dc value of polymer: 0.185 was used in the measurement.  
BSA was used as a control before actual samples are run.



[0179] Synthesis of fKRGD-protected was carried out by a standard Fmoc-solid phase using 2-chlorotriyl chloride resin, HBTU and HOBt coupling agents with diisopropylethylamine (DIPEA). Deprotection of Fmoc group was carried out in 20% piperidine in DMF. Cleavage of fKRGD-protected from the resin was carried out in acetic acid: trifluoroethanol:dichloromethane (1:1:3). Cyclization of fKRGD-protected was carried out using NaHCO<sub>3</sub> and DPPA in DMF. Deprotection of cyclic (fKRGD) was carried out in 95% TFA. Purification of cyclic(fKRGD)-protected and cyclic(fKRGD) was carried out in HPLC system and purity of the products were confirmed with LC-MS.

#### EXAMPLE 1

[0180] A PGA-PTX polymer conjugate was prepared according to the general scheme illustrated in Figure 5 as follows:

[0181] Synthesis of poly-L-glutamic acid-paclitaxel conjugate (PGA-PTX), **1**, was carried out as reported in previous literature. *See* Li et al. "Complete Regression of Well-established tumors using a novel water-soluble poly(L-glutamic acid)-paclitaxel conjugate." *Cancer Research* **1998**, 58, 2404-2409, the contents of which are herein incorporated by reference in its entirety.

#### EXAMPLE 2

[0182] A PGA-PTX-DOX polymer conjugate was prepared according to the general scheme illustrated in Figure 6 as follows:

[0183] Poly-L-glutamic acid-paclitaxel conjugate (50 mg), **1**, was dissolved in DMF (5 mL). Doxorubicin (7 mg), EDC (16 mg), and HOBt (10 mg) were then added. The mixture was stirred for 24 hours. Completion of the reaction was monitored by TLC and the absence of free doxorubicin. A solution of diluted HCl (0.2M) was added to induce precipitation. The mixture was stirred for 2 minutes and centrifuged at 10,000 rpm for 15 minutes. A red-orange solid precipitate was collected, washed with water, and freeze-dried. The product (40 mg) was obtained and was confirmed by <sup>1</sup>H-NMR. The content of doxorubicin was measured by UV-Vis spectroscopy.

#### EXAMPLE 3

[0184] A PGA-PTX-CPT polymer conjugate was prepared according to the general scheme illustrated in Figure 7 as follows:

[0185] Poly-L-glutamic acid-paclitaxel conjugate (50 mg), **1**, was dissolved in DMF (5 mL). Then, camptothecin (8 mg), EDC (16 mg), and HOBt (10 mg) were added. The mixture was stirred for 24 hours. Completion of the reaction was monitored by TLC and the absence of free doxorubicin. A solution of diluted HCl (0.2M) was added to induce precipitation. The mixture was stirred for 2 minutes and centrifuged at 10,000 rpm for 15 minutes. A red-orange solid precipitate was collected, washed with water, and freeze-dried. The product (35 mg) was obtained and was confirmed by <sup>1</sup>H-NMR. The content of camptothecin was measured by UV-Vis spectroscopy.

#### EXAMPLE 4

[0186] A PGA-PTX-CPT-DOX polymer conjugate was prepared according to the general scheme illustrated in Figure 8 as follows:

[0187] Poly-L-glutamic acid-paclitaxel-camptothecin conjugate (30 mg), **3**, was dissolved in DMF (5 mL). Doxorubicin (5 mg), EDC (16 mg), and HOBt (10 mg) were added. The mixture was stirred for 24 hours. Completion of the reaction was monitored by TLC and the absence of free doxorubicin. A solution of diluted HCl (0.2M) was added to induce precipitation. The mixture was stirred for 2 minutes and centrifuged at 10,000 rpm for 15 minutes. A red-orange solid precipitate was collected, washed with water, and freeze-dried. The product (29 mg) was obtained and was confirmed by <sup>1</sup>H-NMR, and the content of doxorubicin was measured by UV-Vis spectroscopy.

#### EXAMPLE 5

[0188] A PGGA-PTX polymer conjugate was prepared according to the general scheme illustrated in Figure 9 as follows:

[0189] Synthesis of poly( $\gamma$ -glutamyl-glutamine)-paclitaxel conjugate, **5**, was carried out as described in U.S. Patent Publication No. 2007-0128118, filed December 1, 2006, which is hereby incorporated by reference in its entirety.

#### EXAMPLE 6

[0190] A PGGA-PTX-DOX polymer conjugate was prepared according to the general scheme illustrated in Figure 10 as follows:

[0191] Poly( $\gamma$ -glutamyl-glutamine)-paclitaxel conjugate (50 mg), **5**, was dissolved in DMF (5 mL). Doxorubicin (7 mg), EDC (16 mg), and HOBt (10 mg) were added. The mixture was stirred for 24 hours. Completion of the reaction was monitored by TLC and the absence of free doxorubicin. A solution of diluted HCl (0.2M) was added to induce precipitation. The mixture was stirred for 2 minutes and centrifuged at 10,000 rpm for 15 minutes. A red-orange solid precipitate was collected, washed with water, and freeze-dried. The product (45 mg) was obtained and was confirmed by  $^1\text{H-NMR}$ , and the content of doxorubicin was measured by UV-Vis spectroscopy.

#### EXAMPLE 7

[0192] A PGGA-PTX-CPT polymer conjugate was prepared according to the general scheme illustrated in Figure 11 as follows:

[0193] Poly( $\gamma$ -glutamyl-glutamine)-paclitaxel conjugate (50 mg), **5**, was dissolved in DMF (5 mL). Camptothecin (8 mg), EDC (16 mg), and HOBt (10 mg) were added. The mixture was stirred for 24 hours. Completion of the reaction was monitored by TLC and the absence of free doxorubicin. A solution of diluted HCl (0.2M) was added to induce precipitation. The mixture was stirred for 2 minutes and centrifuged at 10,000 rpm for 15 minutes. A red-orange solid precipitate was collected, washed with water, and freeze-dried. The product (42 mg) was obtained and was confirmed by  $^1\text{H-NMR}$ . The content of camptothecin was measured by UV-Vis spectroscopy.

#### EXAMPLE 8

[0194] A PGGA-PTX-CAMP-DOX polymer conjugate was prepared according to the general scheme illustrated in Figure 12 as follows:

[0195] Poly( $\gamma$ -glutamyl-glutamine)-paclitaxel-camptothecin conjugate (30 mg), **7**, was dissolved in DMF (5 mL). Doxorubicin (5 mg), EDC (16 mg), and HOBt (10 mg) were added. The mixture was stirred for 24 hours. Completion of the reaction was monitored by TLC and the absence of free doxorubicin. A solution of diluted HCl (0.2M) was added to induce precipitation. The mixture was stirred for 2 minutes and centrifuged at 10,000 rpm for 15 minutes. A red-orange solid precipitate was collected, washed with water, and freeze-

dried. The product (25 mg) was obtained and was confirmed by  $^1\text{H-NMR}$ , and the content of doxorubicin was measured by UV-Vis spectroscopy.

#### EXAMPLE 9

##### Cell culture and preparation:

[0196] B16F0 cells were purchased from ATCC (CRL-6322, ATCC American Type Culture Collection, Rockville, MD) and were grown in Dulbecco's modified Eagle's medium (DMEM) with 10% fetal bovine serum and 100 units/mL penicillin. The cells were grown at 37°C in 5% CO<sub>2</sub> environment. The culture medium was removed and discarded. The cells were rinsed with Dulbecco Phosphate Buffer Solution (DPBS), Trypsin-ethylenediaminetetra-acetic acid (EDTA) solution (0.5ml) was added, and the cells were observed under an inverted microscope to make sure that they were dispersed. Complete growth medium (6.0 to 8.0ml) was added, and the cells were aspirated by gently pipetting. The cell suspension in appropriate aliquots was transferred to new culture plates. The cells were allowed to grow at 37°C in 5% CO<sub>2</sub> for 24 hours before further experiments.

#### EXAMPLE 10

##### In vitro Cytotoxicity MTT Studies

[0197] Polymer conjugates described herein containing an anti-cancer drug are evaluated for their effect on the proliferation of B16F0 melanoma cells at several different concentrations of the drug. Cytotoxic MTT assay is carried out as reported in Monks et al. *JNCI* **1991**, 83, 757-766, which is hereby incorporated by reference in its entirety. Polymer conjugates are prepared as described in Examples 1-8.

#### EXAMPLE 11

##### Binding Studies

[0198] The binding assays are carried out as described in Line et al, [*Journal of Nuclear Medicine*], **2005**, 46, 1552-1560; and Mitra et al., [*Journal of Controlled Release*], **2006**, 114, 175-183, both of which are hereby incorporated by reference in their entireties. Polymers conjugates described herein are prepared as described in Examples 1-8.

### EXAMPLE 12

#### Animals and Tumor Models

[0199] Nude mice (6-7 week old, body weight 25-30g, male) are purchased from Charles River Lab (Willington, MA). B16 cell line is purchased from ATCC (CRL-6322, ATCC American Type Culture Collection, Rockville, MD). The B16 cells are cultured in RMPI 1640 supplemented with 10% Fetal bovine serum, 2 $\mu$ M Glutamine, 1mM non-essential amino acids, 1mM sodium pyruvate, 100U/ml penicillin and 100ug/ml streptomycin. The B16 cells harvested from tissue culture is counted and re-suspended to a concentration of  $5 \times 10^6$  per mL. Using a TB syringe, 0.2 mL (a total of  $1 \times 10^6$  cells) is administered via subcutaneous injection into each mouse. One tumor is inoculated per animal at the right hip. The site of tumor inoculation is shaved prior to inoculation to make it easier to measure the tumor as it grows.

### EXAMPLE 13

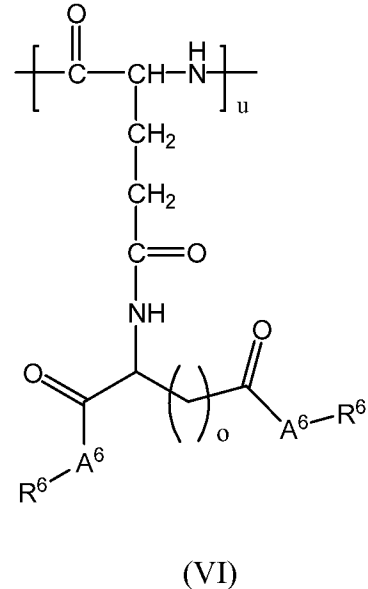
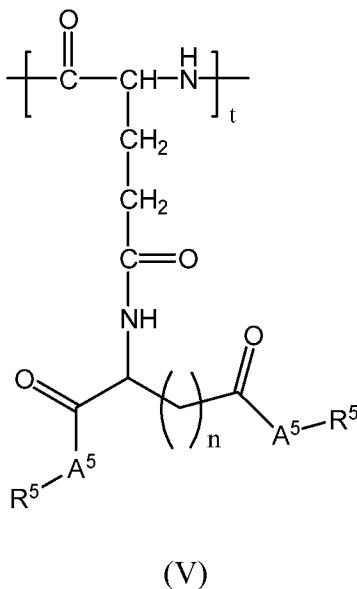
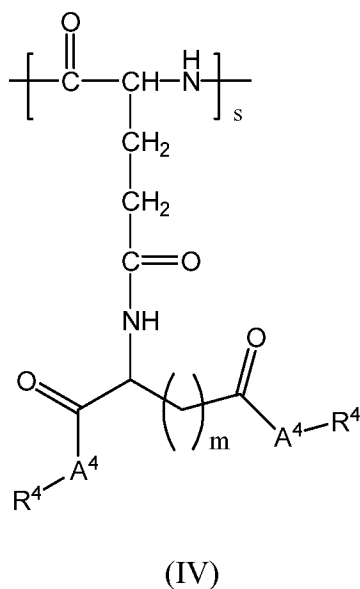
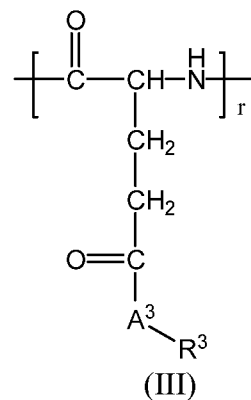
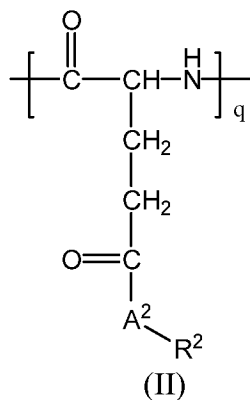
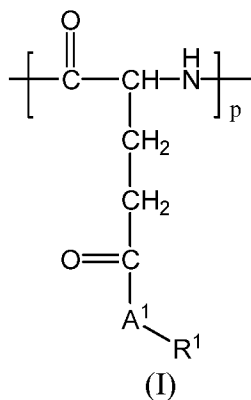
#### Magnetic resonance imaging for tumor accumulation

[0200] Images of mice is acquired on a GE 3T MR scanner using a knee coil pre- and post-contrast. The following imaging parameters are TE: minful, TR= 250 ms, FOV: 8 and 24 slices/slab, and 1.0 mm coronal slice thickness. Polymer conjugates with a compound comprising a magnetic resonance imaging agent, such as Gd(III), and Omniscan-Gd(III)-(DTPA-BMA (0.1 mmol Gd(III)/ kg), a control, are injected via a tail vein into anesthetized mice. The dose of injection of the polymer conjugate and Omniscan<sup>TM</sup> is 0.1 mmol Gd(III)/kg. Images are acquired at pre-injection and at 6 minutes to 4 hours post-injection of the contrast agents.

[0201] It will be understood by those of skill in the art that numerous and various modifications can be made without departing from the spirit of the present invention. Therefore, it should be clearly understood that the forms of the present invention are illustrative only and not intended to limit the scope of the present invention.

WHAT IS CLAIMED IS:

1. A polymer conjugate comprising at least one recurring unit selected from Formulae (I), (II), (III), (IV), (V) and (VI):



wherein:

each A<sup>1</sup>, each A<sup>2</sup>, each A<sup>3</sup>, each A<sup>4</sup>, each A<sup>5</sup> and each A<sup>6</sup> are independently oxygen or NR<sup>7</sup>, wherein R<sup>7</sup> is hydrogen or C<sub>1-4</sub> alkyl;

each R<sup>1</sup>, each R<sup>2</sup>, each R<sup>3</sup>, each R<sup>4</sup>, each R<sup>5</sup> and each R<sup>6</sup> are independently selected from the group consisting of a hydrogen, a C<sub>1-10</sub> alkyl group, a C<sub>6-20</sub> aryl group, an ammonium group, an alkali metal, a polydentate ligand, a polydentate ligand precursor with protected oxygen atoms, a group that comprises a drug, a group that comprises a targeting agent, a group that comprises an optical imaging agent, a

group that comprises a magnetic resonance imaging agent, and a group that comprises a stabilizing agent;

m, n, and o are each independently 1 or 2;

p, q, r, s, t, and u are each independently 0 or  $\geq 1$ , wherein the sum of p, q, r, s, t, and u is 2 or greater; and

provided that at least one of  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  is a group that comprises a first drug, at least one of  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  is a group that comprises a second drug, wherein the first drug and second drug are not the same.

2. The polymer conjugate of Claim 1, wherein  $p + q$  is 2 or greater; and r, s, t and u are 0.
3. The polymer conjugate of Claim 1, wherein  $s + t$  is 2 or greater; and p, q, r and u are 0.
4. The polymer conjugate of Claim 1, wherein  $p + q + r$  is 3 or greater; and s, t and u are 0.
5. The polymer conjugate of Claim 1, wherein  $s + t + u$  is 3 or greater; and q, r and u are 0.
6. The polymer conjugate of Claim 1, wherein  $p + s$  is 2 or greater; and q, r, t and u are 0.
7. The polymer conjugate of Claim 1, wherein  $p + q + s$  is 3 or greater; and r, t and u are 0.
8. The polymer conjugate of Claim 1, wherein  $p + s + t$  is 3 or greater; and q, r and t are 0.
9. The polymer conjugate of Claim 1, wherein  $p + q + s + t$  is 4 or greater; and r and u are 0.
10. The polymer conjugate of Claim 1, wherein  $p + q + r + s + t$  is 5 or greater; and u is 0.
11. The polymer conjugate of Claim 1, wherein  $p + q + s + t + u$  is 5 or greater; and r is 0.
12. The polymer conjugate of Claim 1, wherein  $p + q + r + s + t + u$  is 6 or greater.

13. The polymer conjugate of any one of Claims 1 to 12, wherein the polymer conjugate comprises a total amount of the first drug and the second drug in the range of about 1% to about 50% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate.

14. The polymer conjugate of any one of Claims 1 to 12, wherein the polymer conjugate comprises a total amount of the first drug and the second drug in the range of about 1% to about 40% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate.

15. The polymer conjugate of any one of Claims 1 to 12, wherein the polymer conjugate comprises a total amount of the first drug and the second drug in the range of about 1% to about 30% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate.

16. The polymer conjugate of any one of Claims 1 to 12, wherein the polymer conjugate comprises a total amount of the first drug and the second drug in the range of about 1% to about 20% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate.

17. The polymer conjugate of any one of Claims 1 to 12, wherein the polymer conjugate comprises a total amount of the first drug and the second drug in the range of about 1% to about 10% (weight/weight) based on the mass ratio of the drugs to the polymer conjugate.

18. The polymer conjugate of any one of Claims 1 to 17, wherein the first drug is an anticancer drug.

19. The polymer conjugate of any one of Claims 1 to 18, wherein the second drug is an anticancer drug.

20. The polymer conjugate of any one of Claims 18 to 19, wherein the anticancer drug is selected from the group consisting of a taxane, a camptotheca, and an anthracycline.

21. The polymer conjugate of Claim 20, wherein the taxane is selected from the group consisting of paclitaxel and docetaxel.

22. The polymer conjugate of Claim 21, wherein the paclitaxel is conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C2'-carbon.



23. The polymer conjugate of Claim 21, wherein the paclitaxel is conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C7-carbon.

24. The polymer conjugate of Claim 20, wherein the camptotheca is camptothecin.

25. The polymer conjugate of Claim 20, wherein the anthracycline is doxorubicin.

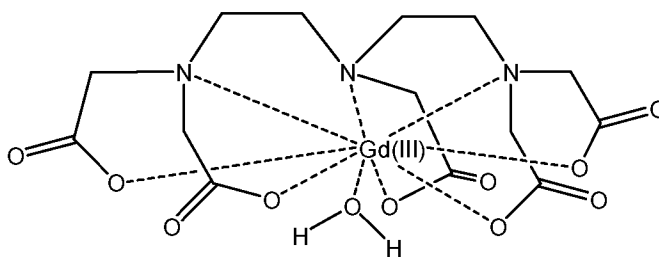
26. The polymer conjugate of any one of Claims 1 to 25, wherein the targeting agent is selected from the group consisting of an arginine-glycine-aspartate (RGD) peptide, fibronectin, folate, galactose, an apolipoprotein, insulin, transferrin, a fibroblast growth factor (FGF), an epidermal growth factor (EGF), and an antibody.

27. The polymer conjugate of any one of Claims 1 to 25, wherein the targeting agent interacts with a receptor selected from the group consisting of  $\alpha_v\beta_3$ -integrin, folate, asialoglycoprotein, a low-density lipoprotein (LDL), an insulin receptor, a transferrin receptor, a fibroblast growth factor (FGF) receptor, an epidermal growth factor (EGF) receptor, and an antibody receptor.

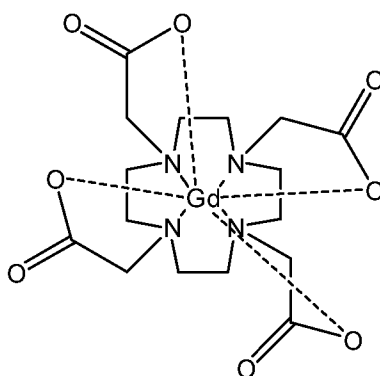
28. The polymer conjugate of any one of Claims 1 to 27, wherein the optical imaging agent is selected from the group consisting of an acridine dye, a coumarine dye, a rhodamine dye, a xanthene dye, a cyanine dye, and a pyrene dye.

29. The polymer conjugate of any one of Claims 1 to 28, wherein the magnetic resonance imaging agent comprises a Gd(III) compound.

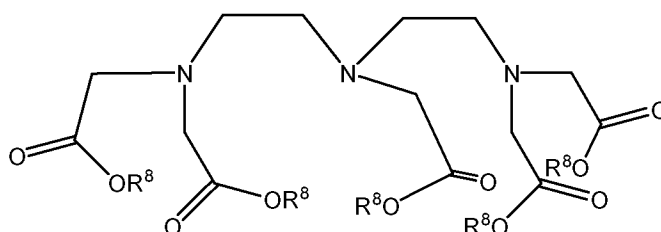
30. The polymer conjugate of Claim 29, wherein the Gd(III) compound comprises:



31. The polymer conjugate of Claim 29, wherein the Gd(III) compound comprises:

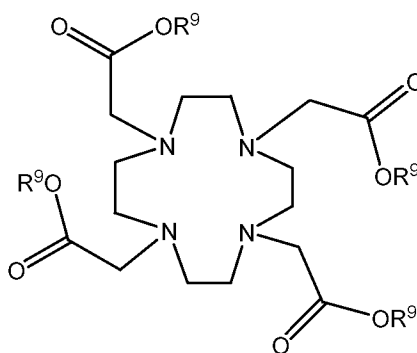


32. The polymer conjugate of any one of Claims 1 to 31, wherein the polydentate ligand comprises:



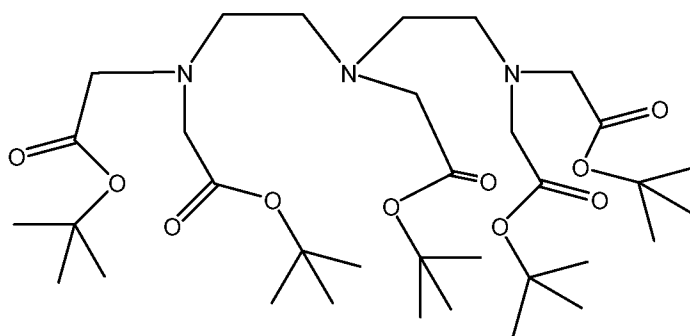
wherein each R<sup>8</sup> is independently hydrogen, ammonium, or an alkali metal.

33. The polymer conjugate of any one of Claims 1 to 31, wherein the polydentate ligand comprises:



wherein each R<sup>9</sup> is independently hydrogen, ammonium, or an alkali metal.

34. The polymer conjugate of any one of Claims 1 to 33, wherein the polydentate ligand precursor with protected oxygen atoms comprises:



35. The polymer conjugate of any one of Claims 1 to 34, wherein the stabilizing agent is polyethylene glycol.

36. The polymer conjugate of any one of Claims 1 to 35, further provided that the at least one of  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$  and  $R^6$  is a group that comprises a third drug, wherein the third drug is different from the first drug and the second drug.

37. The polymer conjugate of any one of Claims 1, 3, or 5 to 36, wherein at least one of m, n, or o is 1.

38. The polymer conjugate of any one of Claims 1, 3, or 5 to 37, wherein at least one of m, n, or o is 2.

39. The polymer conjugate of any one of Claims 1 to 38, wherein the alkali metal is sodium.

40. The polymer conjugate of any one of Claims 1 to 39, wherein any one or more selected from the group consisting of a polydentate ligand, a polydentate ligand precursor with protected oxygen atoms, a group that comprises a drug, a group that comprises a targeting agent, a group that comprises an optical imaging agent, a group that comprises a magnetic resonance imaging agent, and a group that comprises a stabilizing agent, further comprises a linker group.

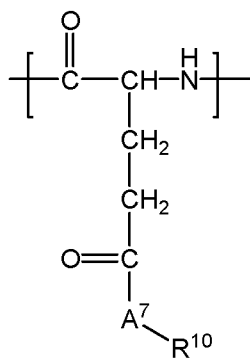
41. The polymer conjugate of any one of Claims 1 to 40, wherein the group that comprises the first drug further comprises a linker group.

42. The polymer conjugate of any one of Claims 1 to 41, wherein the group that comprises the second drug further comprises a linker group.

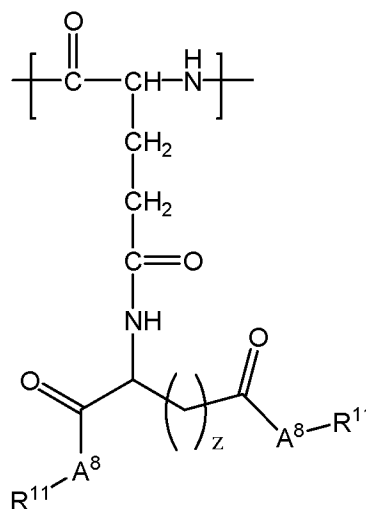
43. A composition comprising the polymer conjugate of any one of Claims 1 to 42 and at least one selected from the group consisting of a pharmaceutically acceptable excipient, a carrier, and a diluent.

44. A method of making the polymer conjugate of any one of Claims 1 to 42, comprising:

dissolving or partially dissolving a polymeric reactant comprising at least one of a recurring unit of formula (VII) and/or a recurring unit of formula (VIII) in a solvent to form a dissolved or partially dissolved polymeric reactant;



(VII)



(VIII)

wherein:

z is independently 1 or 2;

A<sup>7</sup> and each A<sup>8</sup> are oxygen; and

R<sup>10</sup> and each R<sup>11</sup> are independently selected from the group consisting of hydrogen, ammonium, and an alkali metal; and

reacting the dissolved or partially dissolved polymeric reactant with a second reactant and a third reactant, wherein the second reactant comprises the first drug and the third reactant comprises the second drug.

45. The method of Claim 44, wherein the second reactant comprises a substituent selected from the group consisting of hydroxy and amine.

46. The method of Claim 44 or 45, wherein the third reactant comprises a substituent selected from the group consisting of hydroxy and amine.

47. The method of any one of Claims 44 to 46, comprising reacting the dissolved or partially dissolved polymer reactant with at least a portion of the second reactant before reacting with the third reactant.

48. The method of any one of Claims 44 to 46, comprising reacting the dissolved or partially dissolved polymer reactant with at least a portion of the second reactant at about the same time as reacting with the third reactant.

49. The method of any one of Claims 44 to 46, comprising reacting the dissolved or partially dissolved polymer reactant with at least a portion of the second reactant after reacting with the third reactant.

50. The method of any one of Claims 44 to 49, wherein the method further comprises reacting the dissolved or partially dissolved polymeric reactant with a fourth reactant, wherein the fourth reactant comprises at least one selected from the group consisting of a polydentate ligand, a polydentate ligand precursor with protected oxygen atoms, a group that comprises a third drug, a group that comprises a targeting agent, a group that comprises an optical imaging agent, a group that comprises a magnetic resonance imaging agent, and a group that comprises a stabilizing agent.

51. The method of Claim 50, wherein the fourth reactant comprises a substituent selected from the group consisting of hydroxy and amine.

52. The method of Claim 50 or 51, wherein the third drug is an anticancer drug.

53. The method of Claim 52, wherein the anticancer drug is selected from the group consisting of a taxane, a camptotheca, and an anthracycline.

54. The method of Claim 53, wherein the taxane is selected from the group consisting of paclitaxel and docetaxel.

55. The method of Claim 54, wherein paclitaxel is conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C2'-carbon.

56. The method of Claim 54, wherein paclitaxel is conjugated to the recurring unit of formulae (I), (II), (III), (IV), (V), and/or (VI) at the oxygen atom attached to the C7-carbon.

57. The method of Claim 53, wherein the camptotheca is camptothecin.

58. The method of Claim 53, wherein the anthracycline is doxorubicin.

59. The method of any one of Claims 50 to 58, wherein the third drug is different from the first drug and the second drug.

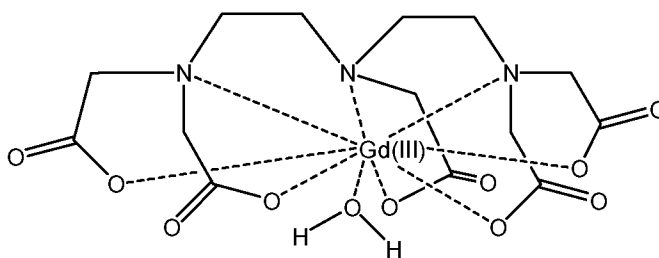
60. The method of any one of Claims 50 to 59, wherein the targeting agent is selected from the group consisting of an arginine-glycine-aspartate (RGD) peptide, fibronectin, folate, galactose, an apolipoprotein, insulin, transferrin, a fibroblast growth factor (FGF), an epidermal growth factor (EGF), and an antibody.

61. The method of any one of Claims 50 to 59, wherein the targeting agent interacts with a receptor selected from the group consisting of  $\alpha_v\beta_3$ -integrin, folate, asialoglycoprotein, a low-density lipoprotein (LDL), an insulin receptor, a transferrin receptor, a fibroblast growth factor (FGF) receptor, an epidermal growth factor (EGF) receptor, and an antibody receptor.

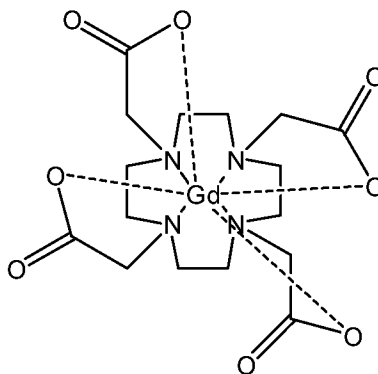
62. The method of any one of Claims 50 to 61, wherein the optical imaging agent is selected from the group consisting of an acridine dye, a coumarine dye, a rhodamine dye, a xanthene dye, a cyanine dye, and a pyrene dye.

63. The method of any one of Claims 50 to 62, wherein the magnetic resonance imaging agent comprises a Gd(III) compound.

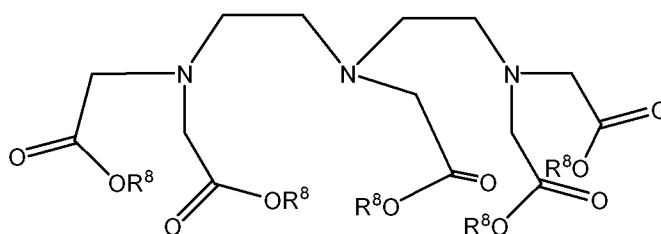
64. The method of Claim 63, wherein the Gd(III) compound comprises:



65. The method of Claim 63, wherein the Gd(III) compound comprises:

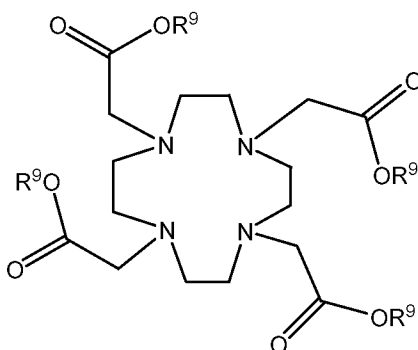


66. The method of any one of Claims 50 to 65, wherein the polydentate ligand comprises:



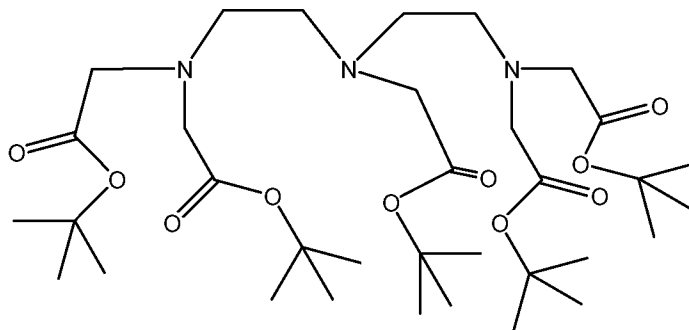
wherein each  $R^8$  is independently hydrogen, ammonium, or an alkali metal.

67. The method of any one of Claims 50 to 65, wherein the polydentate ligand comprises:



wherein each  $R^9$  is independently hydrogen, ammonium, or an alkali metal.

68. The method of any one of Claims 50 to 67, wherein the polydentate ligand precursor with protected oxygen atoms comprises:



69. The method of any one of Claims 50 to 68, wherein the stabilizing agent is polyethylene glycol.

70. The method of any one of Claims 44 to 69, further comprising reacting the dissolved or partially dissolved polymeric reactant in the presence of a coupling agent.

71. The method of Claim 70, wherein the coupling agent is selected from the group consisting of 1-ethyl-3-(3-dimethylaminopropyl)-carbodiimide (EDC), 1,3-dicyclohexyl carbodiimide (DCC), 1,1'-carbonyl-diimidazole (CDI), N,N'-disuccinimidyl

carbonate (DSC), N-[(dimethylamino)-1H-1,2,3-triazolo-[4,5-b]pyridine-1-yl-methylene]-N-methylmethanaminium hexafluorophosphate N-oxide (HATU), 2-[(1H-benzotriazol-1-yl)-1,1,3,3-tetramethylaminium hexafluorophosphate (HBTU), 2-[(6-chloro-1H-benzotriazol-1-yl)-1,1,3,3-tetramethylaminium hexafluorophosphate (HCTU), benzotriazole-1-yl-oxy-tris-pyrrolidino-phosphonium hexafluorophosphate, bromo-tris-pyrrolidino-phosphonium hexafluorophosphate, 2-[(1H-benzotriazol-1-yl)-1,1,3,3-tetramethylaminium tetrafluoroborate (TBTU), and benzotriazol-1-yl-oxy-tris-(dimethylamino)phosphonium hexafluorophosphate (BOP).

72. The method of any one of Claims 44 to 71, wherein the solvent is a polar aprotic solvent.

73. The method of Claim 72, wherein the solvent is selected from the group consisting of N,N-dimethylformamide (DMF), dimethyl sulfoxide (DMSO), N-methyl-2-pyrindone (NMP), and N,N-dimethylacetamide (DMAc).

74. The method of any one of Claims 44 to 73, further comprising reacting the dissolved or partially dissolved polymeric reactant in the presence of a catalyst.

75. The method of Claim 74, wherein the catalyst is 4-dimethylaminopyridine (DMAP).

76. A method of treating or ameliorating a disease or condition comprising administering an effective amount of the polymer conjugate of any one of Claims 1 to 42 or the composition of Claim 43 to a mammal in need thereof.

77. The method of Claim 76, wherein the disease or condition is selected from the group consisting of lung tumor, breast tumor, colon tumor, ovarian tumor, prostate tumor, and melanoma tumor.

78. The method of Claim 76, wherein the disease or condition is selected from the group consisting of lung cancer, breast cancer, colon cancer, ovarian cancer, prostate cancer, and melanoma.

79. A method of diagnosing a disease or condition comprising administering an effective amount of the polymer conjugate of any one of Claims 1 to 42 or the composition of Claim 43 to a mammal in need thereof.



80. The method of Claim 79, wherein the disease or condition is selected from the group consisting of lung tumor, breast tumor, colon tumor, ovarian tumor, prostate tumor, and melanoma tumor.

81. The method of Claim 79, wherein the disease or condition is selected from the group consisting of lung cancer, breast cancer, colon cancer, ovarian cancer, prostate cancer, and melanoma.

82. A method of imaging a portion of tissue comprising contacting a portion of tissue with an effective amount of the polymer conjugate of any one of Claims 1 to 42 or the composition of Claim 43.

83. The method of Claim 82, wherein the tissue is from a tumor selected from the group consisting of lung tumor, breast tumor, colon tumor, and ovarian tumor.

84. The method of any one of Claims 76-83, wherein the polymer conjugate is administered intravenously.

85. Use of the polymer conjugate of any one of Claims 1 to 42 or the composition of Claim 43 for the preparation of a medicament for treating or ameliorating a disease condition.

86. The use of Claim 85, wherein the disease or condition is selected from the group consisting of lung tumor, breast tumor, colon tumor, ovarian tumor, prostate tumor, and melanoma tumor.

87. The use of Claim 85, wherein the disease or condition is selected from the group consisting of lung cancer, breast cancer, colon cancer, ovarian cancer, prostate cancer, and melanoma.

88. Use of the polymer conjugate of any one of Claims 1 to 42 or the composition of Claim 43 for the preparation of a medicament for diagnosing a disease or condition.

89. The use of Claim 88, wherein the disease or condition is selected from lung tumor, breast tumor, colon tumor, ovarian tumor, prostate tumor, and melanoma tumor.

90. The use of Claim 88, wherein the disease or condition is selected from the group consisting of lung cancer, breast cancer, colon cancer, ovarian cancer, prostate cancer, and melanoma.

91. Use of the polymer conjugate of any one of Claims 1 to 42 or the composition of Claim 43 for the preparation of a medicament for imaging a portion of tissue.

92. The use of Claim 91, wherein the tissue is from a tumor selected from the group consisting of lung tumor, breast tumor, colon tumor, and ovarian tumor.

93. The use of any one of Claims 85 to 92, wherein the polymer conjugate is administered intravenously.

94. A compound of any one of Claims 1 to 42 or the composition of Claim 43 for treating or ameliorating a disease condition.

95. The compound of Claim 94, wherein the disease or condition is selected from the group consisting of lung tumor, breast tumor, colon tumor, ovarian tumor, prostate tumor, and melanoma tumor.

96. The compound of Claim 94, wherein the disease or condition is selected from the group consisting of lung cancer, breast cancer, colon cancer, ovarian cancer, prostate cancer, and melanoma.

97. A compound of any one of Claims 1 to 42 or the composition of Claim 43 for diagnosing a disease or condition.

98. The compound of Claim 97, wherein the disease or condition is selected from lung tumor, breast tumor, colon tumor, ovarian tumor, prostate tumor, and melanoma tumor.

99. The compound of Claim 97, wherein the disease or condition is selected from the group consisting of lung cancer, breast cancer, colon cancer, ovarian cancer, prostate cancer, and melanoma.

100. A compound of any one of Claims 1 to 42 or the composition of Claim 43 for imaging a portion of tissue.

101. The compound of Claim 100, wherein the tissue is from a tumor selected from the group consisting of lung tumor, breast tumor, colon tumor, and ovarian tumor.

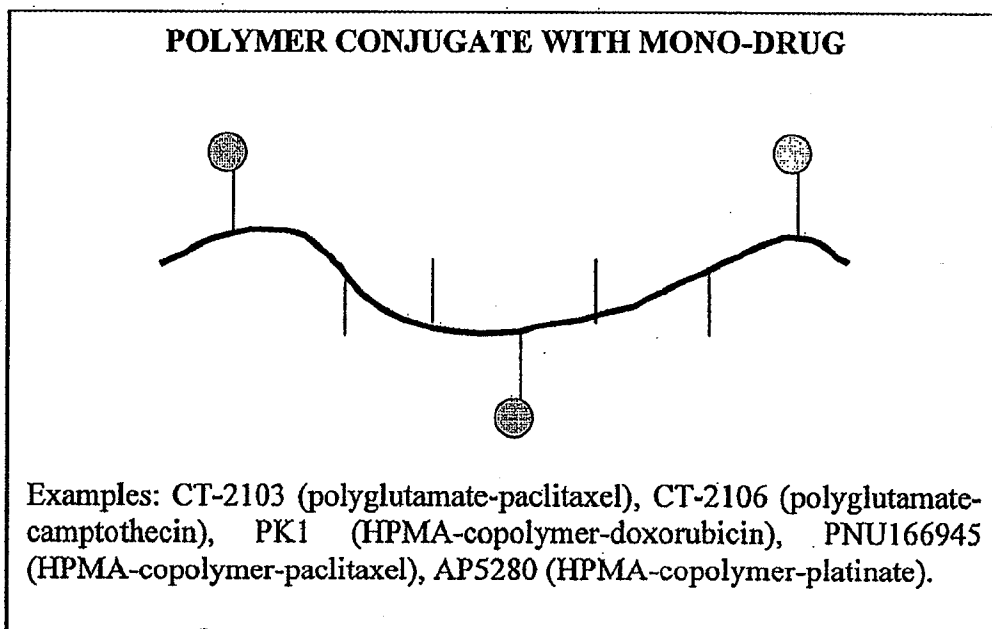
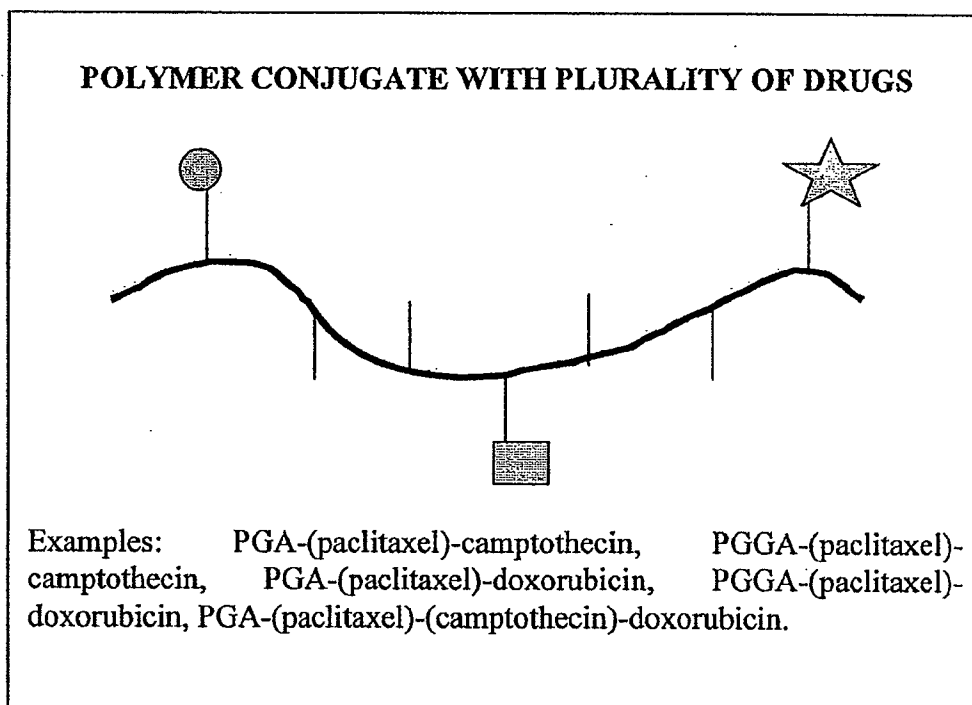
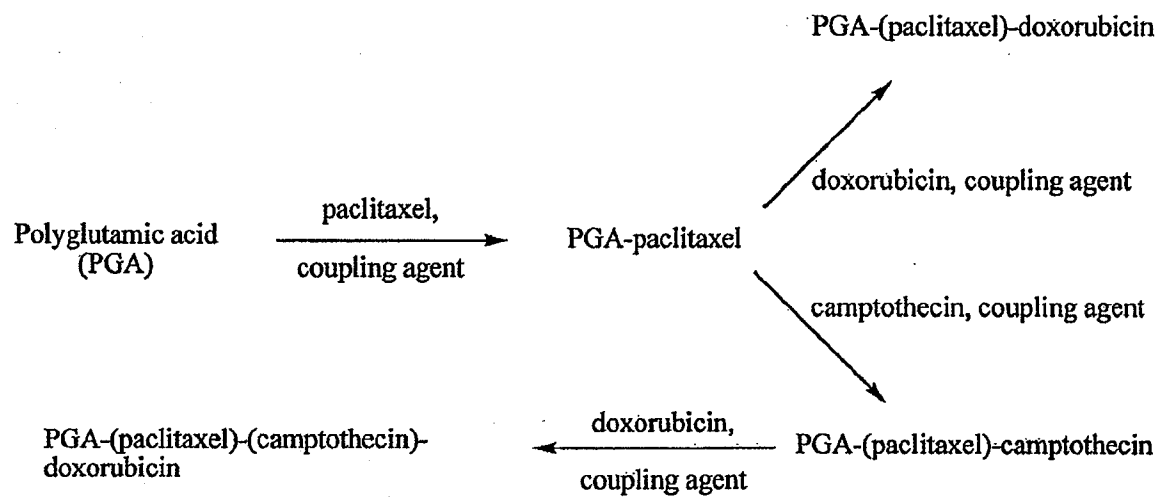
**FIGURE 1****FIGURE 2**

FIGURE 3



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FIGURE 4

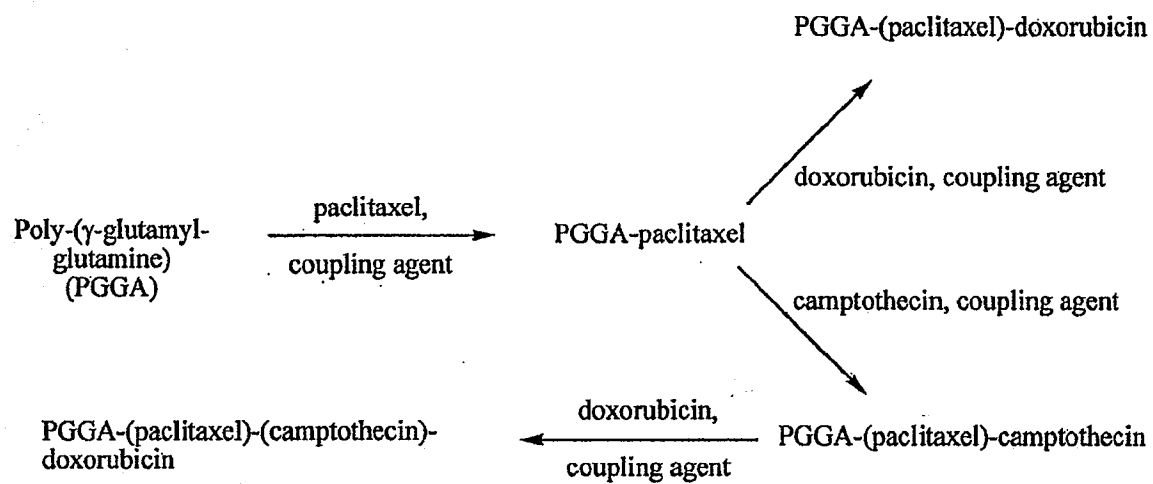
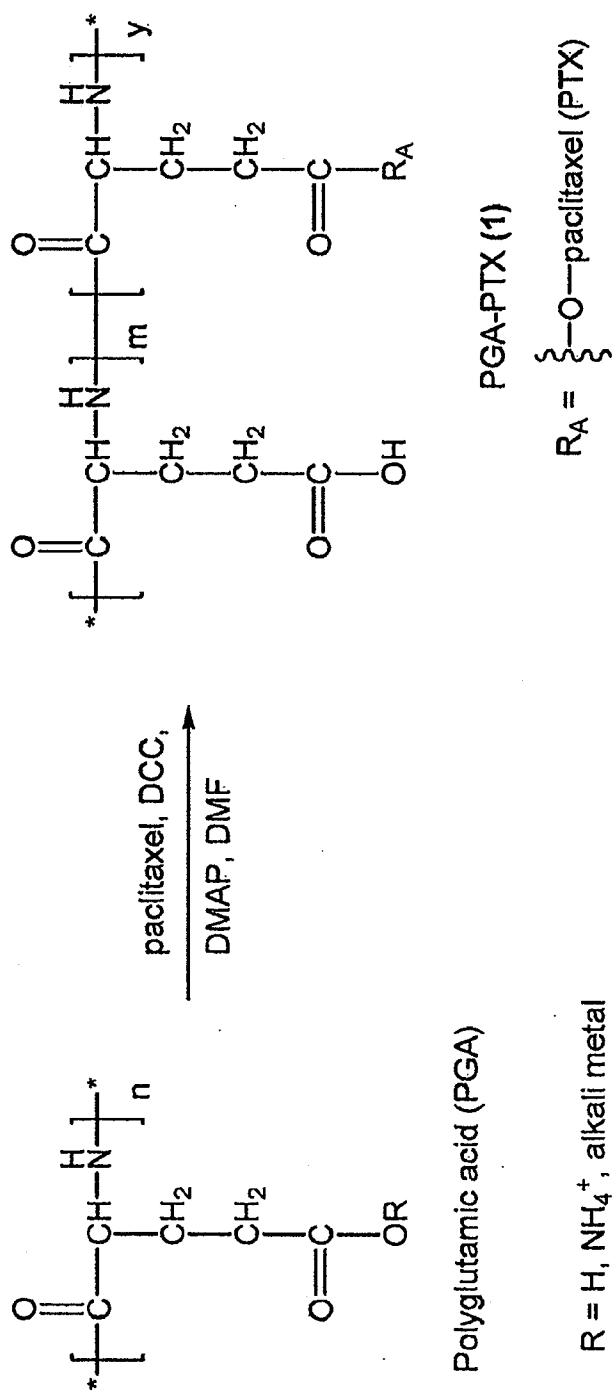
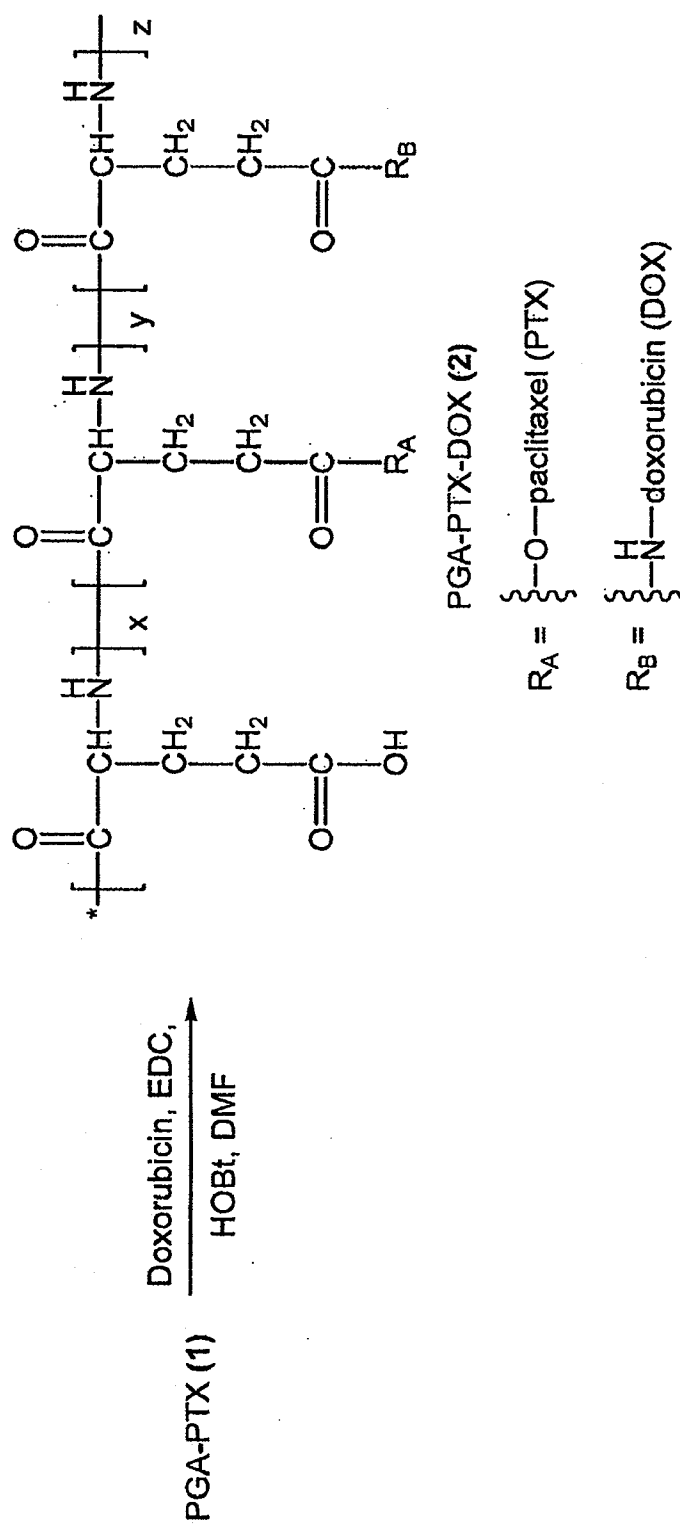


FIGURE 5

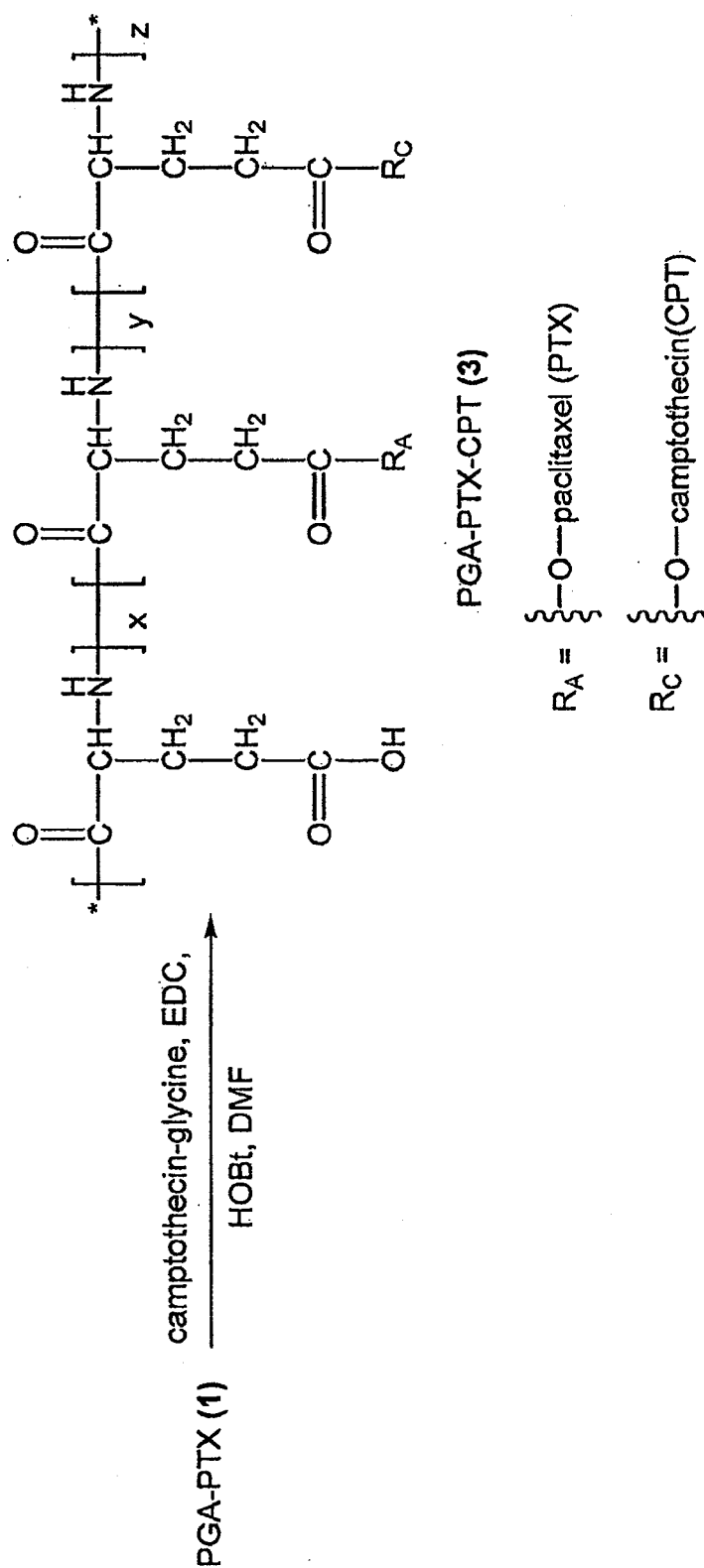


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FIGURE 6

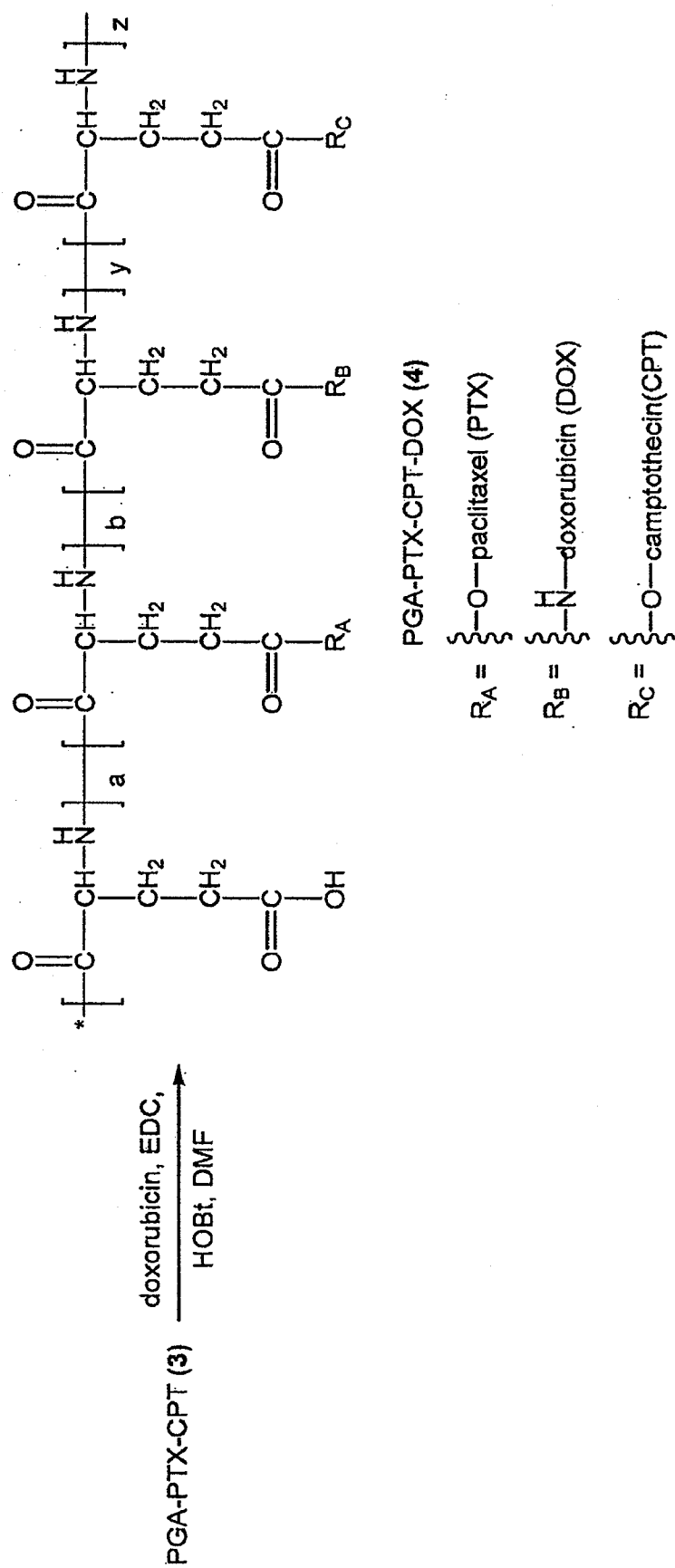


**FIGURE 7**



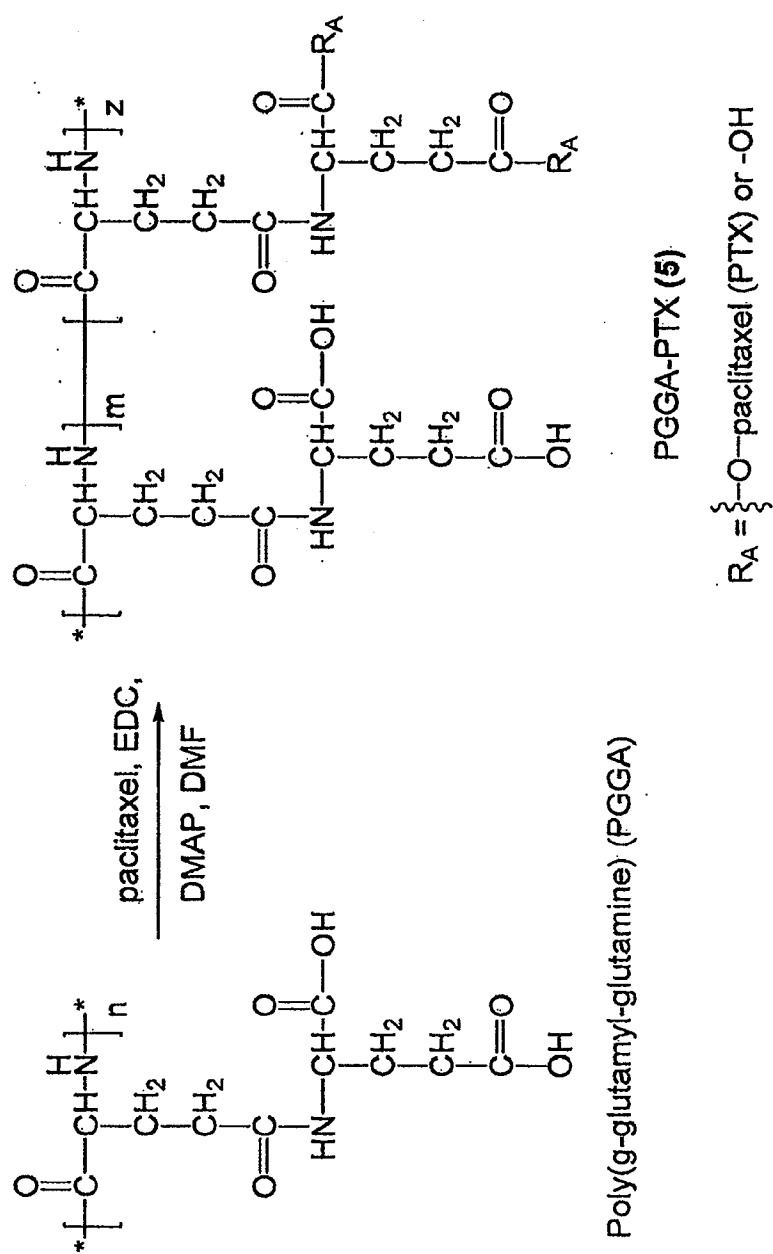


## FIGURE 8



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FIGURE 9



**FIGURE 10**

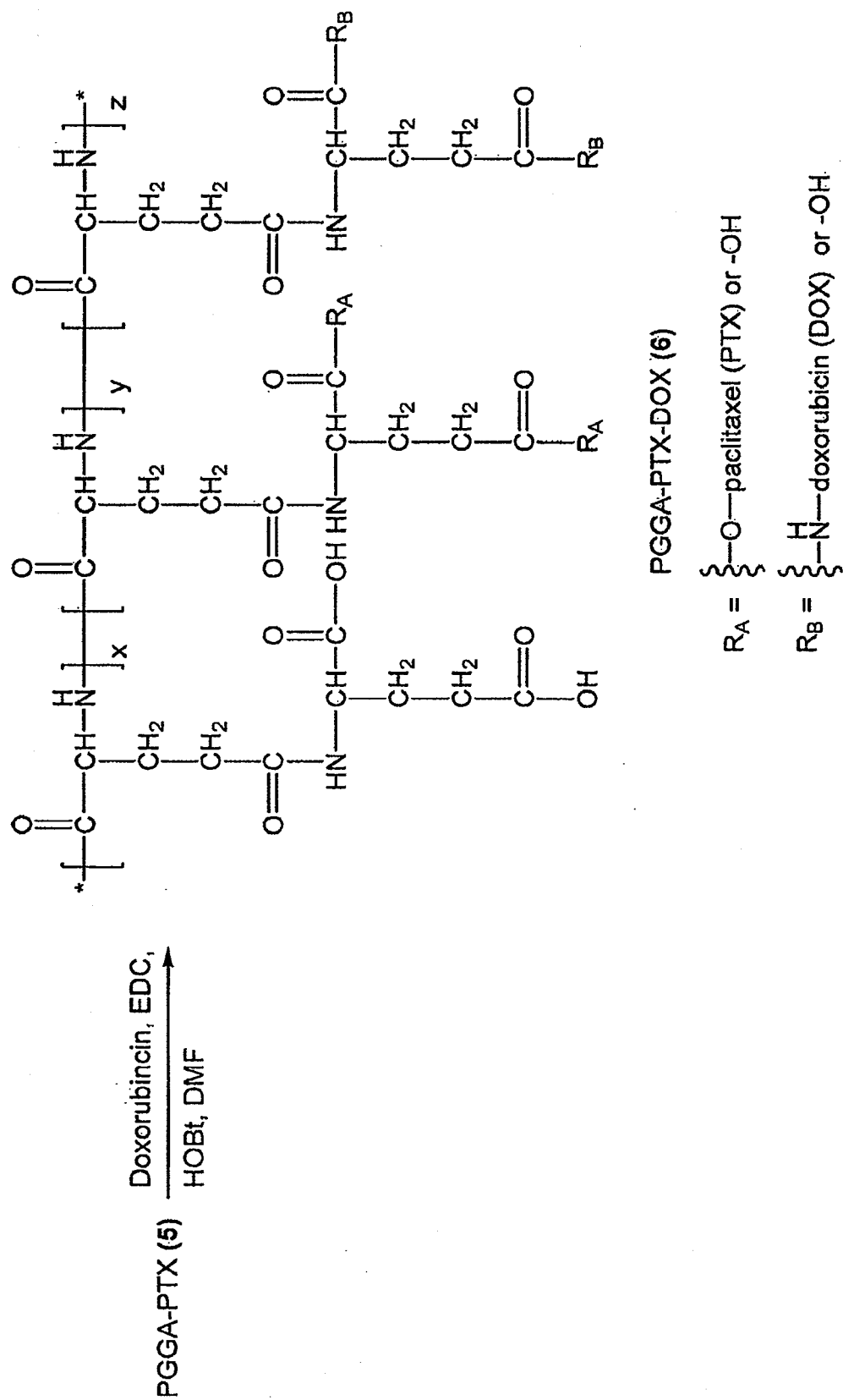




FIGURE 12

