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(54) CLATHRIN REPLACEMENT THERAPEUTICS

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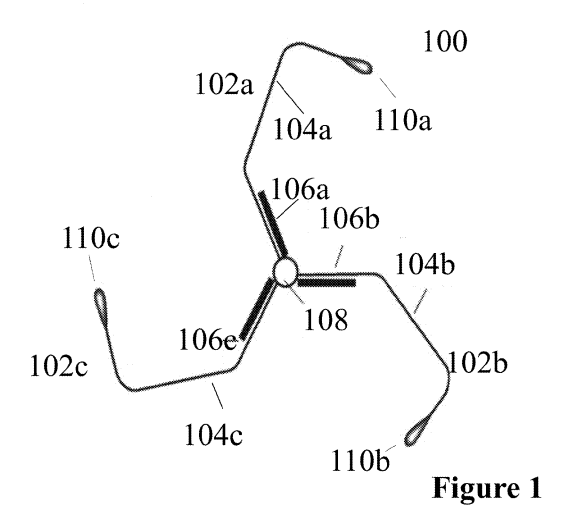
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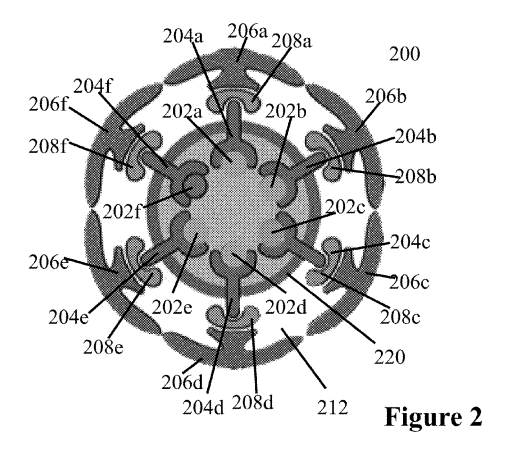
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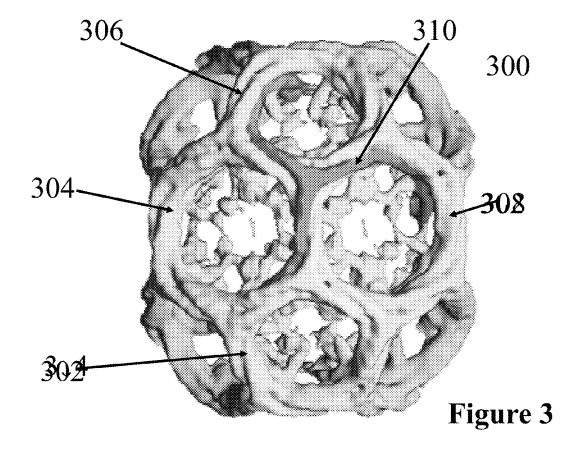
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(57)ABSTRACT

The invention in suitable embodiments is directed to replacement therapeutics. In one aspect, a medicament is comprised in whole or in part of one or more clathrin heavy chain protein that is formed from a plurality of isolated, synthetic or recombinant clathrin protein molecules. In one embodiment, a man-made clathrin heavy chain protein composition replaces and/or modifies cell elements or processes, in vivo or in vitro, thereby treating a disease, condition, or disorder comprising at least one of cell.







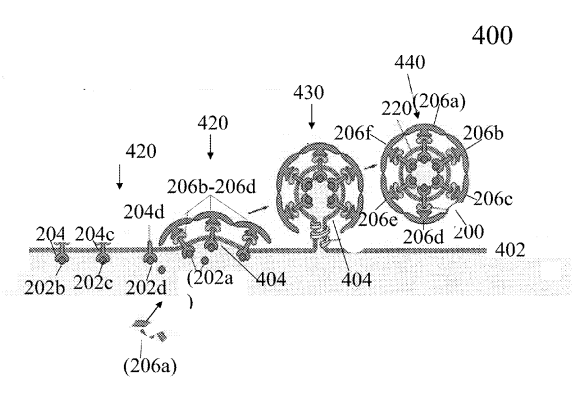


Figure 4

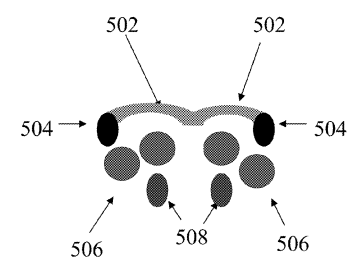
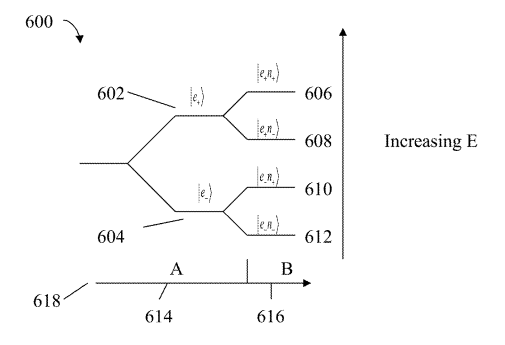


Figure 5



Increasing Magnetic Field

Figure 6

CLATHRIN REPLACEMENT THERAPEUTICS

FIELD OF THE INVENTION

[0001] This is a division of pending USPTO Utility Application No. 12399906, with the title, "DYNAMIC BIO-NANOPARTICLE ELEMENTS", originally filed on Mar. 6, 2009, and claims priority to that date. The invention relates generally to the field of nanoparticles, and more specifically, in one embodiment, to bio-nanoparticle elements formed from materials comprised of self-assembling clathrin protein molecules. In one aspect one or more man-made clathrin heavy chain protein is a medicament that replaces and/or modifies cell elements or processes, in vivo or in vitro. In another invention embodiment, the invention relates to a multifunction nanoscale bio-nanoparticle platform, such as a biomedical platform, bio-molecular platform, electronics platform, information processing platform, and the like, and using such bio-nanoparticle platforms for treating disorders of cells, in vivo or in vitro.

BACKGROUND OF THE INVENTION

[0002] Structures at the nanoscale are sometimes referred to as nanoparticles. Some nanoparticles comprise cage elements that form cavities and or comprise vesicle elements; examples of which in the prior art teach elements such as nano-carbon endohedral cages (Fullerenes); capsids, the protein shell of a virus; liposomes; lipids; heat shock proteins; ferritins; vault ribonucleoprotein particles; Clathrin protein cages; and Coatomer I/II protein cages, among other various cage- or vesicle-forming elements. Additionally, prior art teaches that protein cage elements can coat vesicle elements; for example, Clathrin and Coatomer coated vesicles (CCV's). Additionally, prior art teaches that one or more types of cargo elements can be located internally with respect to a cage and vesicle element.

[0003] A cavity forming protein cage and a cage coated vesicle implementation is taught in issued U.S. Pat. No. 7,393,924 (Jul. 1, 2008, Vitaliano et al.) The cage and cage coated vesicle elements are formed in vitro from a plurality of isolated Clathrin/Coatomer protein subunits. As taught in U.S. Pat. No. 7,393,924, the enhanced functionalization capabilities of the isolated Clathrin and Coatomer I/II protein molecules enable a number of properties and features that make them superior to other cage and cage coated vesicle elements in the prior art.

[0004] But the instant invention teaches nanoscale element fabrication, assembly, operation, behavior and properties that are unique from prior protein art that encompasses various types of cavity-forming cage structures formed in vitro from a plurality of self-assembling subunits. For example, a fully formed Clathrin cage element as taught in U.S. Pat. No. 7,393,924, and generally speaking taught in other Clathrin art, is comprised of a plurality of 3-legged triskelia, each triskelion having 6 protein subunits; 3 Clathrin heavy chain and 3 Clathrin light chain subunits.

[0005] In marked contrast, the instant invention teaches that complete cages comprised of a plurality of 3-legged triskelia are not required to comprise one or more types of efficacious elements. Instead, in its most essential embodiment the instant invention teaches one or more nanoscale elements of one or more types formed from isolated, synthetic and or recombinant amino acid residues comprising in

whole or in part one or more types of Clathrin and or Coatomer I/II proteins of one or more isoforms, including cloned isoforms. These isoforms with their differing amino acid sequences comprise (in this example, humans) the various types of Clathrin heavy chains, the various types of Clathrin light chains, encompass the distinct heavy chain and light chain segments and domains, and in the case of Coatomer, comprise and encompass its domains and subunits, with different combinations of the latter known to exist within Coatomer complexes. Examples of amino acid sequences comprising Clathrin and Coatomer proteins, and their respective isoforms are listed in SEQ ID NO:1 to SEQ ID NO:30. Accordingly, one or more instant invention embodiments may also comprise minimalist, non-cage elements of one or more types. The minimalist element structure afforded by the instant invention affords a much broader and richer variety of element configurations and embodiments than those taught in prior Clathrin or other protein

[0006] For example, freed of the constraints of only forming cavity-forming protein cages in vitro, one or more non-cage invention elements may also form one or more other types of nanoscale elements and structures, enabling new classes and types of applications. Example non-cage embodiments include, but are not limited to, functionalized nano-tubule structures; protein-based nano-dendrimers suitable for biomedical and bio-molecular applications; and self-assembling, stable, bioactive, protein-based, hydrogel nanoparticles (nanogels). In other embodiments, one or more nanoscale elements and structures may be additionally formed and comprised of one or more non-invention elements of one or more types. Such structural plasticity and flexible element functionality are not taught in prior protein cage art.

[0007] Prior art often teaches one or more types of protein cages that carry one or more types of additional elements, e.g., cargo, to enable overall functionality and produce efficacious results. However, unlike prior art, the instant invention teaches, in one embodiment, one or more noncage or cage elements may carry no additional elements like cargo, yet still can comprise inherently efficacious elements of one or more types, like drug elements, but not limited to. In one embodiment, one or more invention elements operating alone and without any additional elements such as cargo and the like comprise unique new types of inherently efficacious agents and elements that are distinctly different in behavior and functionality from prior art, and their unique features correspondingly enable new types of applications. [0008] In another embodiment, one or more elements and or their additional elements in whole or in part may require only minimal functionalization to be efficacious; e.g., they may not require PEGylation or other types of functionalization to operate effectively.

[0009] In another embodiment, one or more elements carry one or more types of cargo and the cargo acts as the efficacious element. In another embodiment, one or more elements together with cargo elements act in efficacious concert.

[0010] In another embodiment, one or more elements are penetrating elements that enter one or more cells and gain access to the cytosol and intracellular elements of one or more types, including one or more cell organelles. Such elements may, in one embodiment, require minimal functionalization. In another embodiment, one or more elements

may comprise one or more membrane fusion elements. These various features are not taught in prior protein cage art. In one embodiment, using cell crossing techniques yield efficacious cancer treatments, gene therapy, and the like.

[0011] Further, in cage, cavity, and vesicle prior art, one or more types of additional elements, e.g., cargo, are often inserted into a complex, fully formed structure, a sometimes difficult and laborious process. But the invention, in one or more embodiments, teaches that using utilizing non-cage elements of one or more types makes the addition of one or more elements less difficult as there is no insertion process into a cage, cavity, or vesicle to contend with. In another embodiment, additional element functionalization is simplified by decorating just the external surface of a cage, a feature not taught in prior Clathrin art.

[0012] In another embodiment, one or more assay, diagnostic, therapeutic, and prosthetic applications and the like can be performed ensemble using the same bioengineered element.

[0013] These various functionalization capabilities enable a highly flexible nano-platform that features improved stability, rigidity, functionality and loading capacity relative to other nanoparticles, and being comprised of ubiquitous proteins, features low antigenicity in one or more embodiments. In one illustrative embodiment, one or more elements may be harmlessly dissolved, passed, and or excreted from the body.

[0014] In one embodiment, the current application teaches one or more elements comprising one or more types of hybrid elements and arrangements, which can produce efficacious results. In one embodiment, one or more invention elements are conjugated to natural biological/molecular elements, like cells, but not limited to, forming one or more types of hybrid elements in vitro and or in vivo. Such hybrid elements may operate alone or with additional elements, e.g., with cargo. In another embodiment, such hybrid elements may fuse in vitro and or in vivo with non-invention elements, such as those comprising natural elements in cells, but not limited to. This type of hybrid/fusion capability and flexibility is not taught in the prior art.

[0015] In another embodiment, the current application teaches one or more elements, functioning alone or with one or more additional elements, which comprise efficacious replacements for one or more elements of one or more types, including non-invention elements. In one embodiment, one or more elements may replace one or more types of naturally occurring cell elements, to efficacious effect. This replacement capability is not taught in the prior art.

[0016] In one embodiment, the instant invention teaches one or more elements, functioning alone or with one or more additional elements, which comprise one or more cellular repair elements, of one or more types; a capability not taught in the prior art. In another embodiment the elements are cellular regeneration elements.

[0017] Prior art also does not teach that cage, vesicle elements, or their various subunit elements efficaciously operate in the extra-cellular spaces, e.g., in the synaptic spaces between neurons. But the instant invention teaches one or more types of elements capable of such extracellular operation, including for the in situ remediation, removal and or sequestration of undesirable organic and or non-organic elements.

[0018] The invention further teaches a biological model that is consistent, not from the complete cage element level

up, but from the minimalist, non-cage element level up, in vitro and in vivo, making drug discovery safer, more efficacious, more time and cost effective, and overall, a much more rapid process than prior art.

[0019] In another embodiment, one or more elements may comprise one or more types of minimalist, non-cage elements than that taught in prior art for doing clinical trials of one or more types of agents, including their targeted agent delivery, including high precision dosing.

[0020] In one embodiment, the instant invention teaches one or more elements that in whole or in part execute one or more types of actions for creating, spawning, comprising, modifying, repairing, regenerating, reassembling, and or control and regulation of one or more cells, cellular elements, cell organelles, including like actions and behaviors involving cellular processes such as endocytosis, exocytosis, mitosis, trafficking and signaling, communication between cells, receptor upregulation and downregulation, other behaviors, and the like. Failures and defects in any of these cellular elements and processes can lead to diseases, for example, cancer. This type of efficacious behavior is not taught in prior art, including in protein cage art.

[0021] In one invention embodiment, one or more elements, with or without additional elements, and in some embodiments with minimal functionalization, enter the central nervous system, including passing the blood brain barrier (BBB) for efficacious effect. Although different protein cage types, e.g., viruses, have been investigated as MRI nano-probes, some types of these cages in prior art did not cross the BBB, and other types in prior art were shown to be immunogenic after crossing the BBB.

[0022] In one embodiment, the invention enables post administration delivery of one or more types of agents into the CNS in 30 minutes or less. In other embodiments, delivery of agents occurs in 30 minutes or more. In another embodiment, agents operate in the inter-neuronal spaces. Prior art does not teach such flexible CNS delivery arrangements.

[0023] The instant invention teaches self-directing, self-replicating, self-adapting, self-repairing, self-regulating, and or self-regenerating methods for one or more minimalist, non-cage elements, which can also perform on-the-fly target prioritization. Prior protein cage art does not teach such self-modifying methods at a minimalist, non-cage element level.

[0024] Prior art does not teach enabling and or utilizing quantum mechanical effects using just one or more minimalist, non-cage elements. But in one embodiment, the instant invention teaches enabling and utilizing such quantum mechanical effects.

[0025] The instant invention also teaches a plurality of elements of one or more types that can, in one illustrative embodiment, function as biomedical platform and the like, and in another example embodiment, function as a biomolecular component platform and the like, or as an information processing platform that can carry out algorithmically defined actions, and other types of platforms.

[0026] Thus, there exists a need for an improved bio-nano-structure element that overcomes the limitations in the prior art for various types of in vivo and in vitro applications.

SUMMARY OF THE INVENTION

[0027] The invention, in one aspect, remedies the deficiencies of the prior art by teaching modifiable, interactive, dynamic bio-nanoparticle elements, some of which may comprise minimalist, non-cage embodiments, with or without one or more additional elements of one or more types located on and or in one or more elements; whose applications, in one or more embodiments, focus on forming in whole or in part one or more nanoscale elements and structures of one or more types that execute one or more functions and or effect one or more ends in vivo and or in vitro.

[0028] In one illustrative embodiment, the invention is an improvement over other in vivo biodegradable polymer nanospheres, liposomes, lipids, capsids agent delivery systems, as well as endohedral Fullerenes and other bionanoparticles in the prior art because the invention enables, among other unique features:

[0029] Simplified nanoscale fabrication

[0030] Simplified cargo and other element type attachment.

[0031] Cell and organelle crossing, and or membrane fusion.

[0032] Low antigenic, "green" nanotechnology.

[0033] Interaction, control, and regulation of cellular processes, like endocytosis, exocytosis, mitosis, trafficking and signaling, communication between cells, receptor upregulation and downregulation, other cellular behaviors, and the like.

[0034] Entering the CNS, including passing the blood brain barrier, and in some cases, in less than 30 minutes post administration.

[0035] One or more elements that carry no additional elements, like cargo, and operating alone produce an efficacious effect, acting like a drug, for example.

[0036] Hybrid invention elements comprised of one or more types of non-invention elements, e.g., natural cell elements.

[0037] Self-modifying, orchestrated actions at a minimalist, non-cage level using natural control laws that govern biological elements.

[0038] Methods and behaviors defined by algorithms. [0039] In one particular embodiment, one or more of self-assembling Clathrin and or Coatomer elements are functionalized, modified and or bioengineered using commercially available biotechnology tools and other tools and techniques known in the art, which makes the invention more versatile and cost-effective than the existing art.

[0040] In another embodiment, one or more elements are also comprised of one or more non-invention elements, e.g., one or more invention elements are conjugated to natural biological/molecular elements, like cells, but not limited to, forming one or more types of hybrid elements in vitro and or in vivo.

[0041] In one illustrative embodiment, one or more elements can be of any suitable size. According to an illustrative embodiment, one or more elements are nanoscale elements.

[0042] The invention, in one embodiment, teaches one or more elements that dynamically and interactively respond to changing in vivo and or in vitro environments; e.g., change of pH, temperature, biochemical, or biological conditions, and the like.

[0043] In one embodiment, one or more elements, in one or more configurations, utilize self-directing, self-adapting, self-assembling, self-repairing, self-regenerating, self-regulating, and or self-replicating methods.

[0044] In one embodiment, one or more elements, in one or more configurations, utilize goal directed methods.

[0045] In one embodiment, one or more elements utilize, respond to, and or exhibit one or more effects, such as quantum mechanical, mechanical, photonic, acoustic, electrical, biochemical and chemical, and the like.

[0046] The invention, in one embodiment, provides one or more elements that maintain structural and or functional integrity long enough to do useful work, in vivo and or in vitro

[0047] According to one feature, one or more elements re-supply, repair, reassemble and or regenerate defective, destroyed and or inoperable elements of one or more types, including non-invention elements, in vivo and or in vitro.

[0048] In another embodiment, one or more types of elements, unlike other nanoparticles in the art; such as nano-carbon, virus capsids, as well as nano-coating elements like polysorbate; may exhibit no or limited immunogenic, toxic, and or environmental impact effects, and depending on cargo and other element type also may require little or no functionalization.

[0049] In another embodiment, elements maintain structural integrity at room temperature in vitro and vivo, which eliminates the need for elaborate structure stabilizing mechanisms, like cooling systems.

[0050] Another advantage of the invention is that its protein material does not exhibit extreme hydrophobicity.

[0051] According to another feature, one or more elements are protected from the external environment, and the invention is stable with respect to dissociation and any element toxicity is sequestered from the surrounding in vivo and or in vitro environment.

[0052] In some embodiments, bonding and or attachment methods of one or more types, e.g., covalent, non-covalent, and any other bond type that can be explained by quantum theory, are used to directly attach one or more elements, internally or externally to one or more other elements in an ordered arrangement.

[0053] In one embodiment, one or more elements each may bond with one or more other elements, of one or more types, including invention and non-invention elements.

[0054] In one embodiment, one or more elements may additionally have located on and or in them one or more cargo elements of one or more types, formed from one or more types of molecules.

[0055] In another embodiment, the invention features precise, highly ordered placement of additional elements, like cargo elements, with minimal inter-element spacings on one or more elements and structures.

[0056] In one embodiment, one or more cargo elements comprise natural, isolated, synthetic and or recombinant elements.

[0057] In one embodiment, one or more cargo carrying elements include in whole or in part one or more non-invention elements of one or more types.

[0058] In one embodiment, one or more cargo elements and or cargo carrying elements comprise hybrid elements of one or more types.

[0059] In one embodiment, one or more elements of one or more types do not carry cargo elements.

[0060] In one embodiment, nanoscale ensembles comprising one or more types of elements allow for a large variety and number of possible cargo element configurations.

[0061] In one embodiment, one or more elements may additionally have located on and or in them one or more elements such as ligand elements, receptor elements, adaptor protein elements, and the like, formed from one or more types of molecules, which may also comprise one or more hybrid elements formed from one or more non-invention elements.

[0062] In another embodiment, one or more elements may be comprised of one or more elements derived in part from one or more types of elements, for example, but not limited to, an amino acid sequence derived from a Clathrin or Coatomer protein.

[0063] In another illustrative embodiment, one or more elements, in one or more configurations, are coated in whole or in part with chemicals, metals, biomaterials, and or other substances, of one or more types.

[0064] In another illustrative embodiment, one or more elements, in one or more configurations, comprise one or more organic, inorganic, and or synthetic material elements, of one or more types, in one or more forms and or phases, in whole or in part

[0065] In one embodiment, one or more elements are radiation shielded, radio frequency (RF) shielded, thermally shielded, chemically shielded, and the like, in whole or in part, and in one or more configurations.

[0066] In various embodiments, one or more elements may be of more than one functionalization type, and or express more than one type of functionality.

[0067] In one embodiment, one or more elements in whole or in part may require minimal or no functionalization to be efficacious elements, like a drug and the like, but not limited to

[0068] In another embodiment, one or more elements in whole or in part comprise one or more structures, of one or more types.

[0069] In another embodiment, one or more elements in whole or in part comprise a shape programmable and or shaped scaffolding system via which one or more elements of one or more types form one or more structures with one or more types of shapes and or functions.

[0070] In one embodiment, one or more elements act as one or more types of efficacious replacements for one or more other elements, including non-invention elements, in vitro and or in vivo, e.g., act as replacements for one or more natural elements commonly found in cells, but not limited to. This type of replacement functionality is not taught in prior art, including protein cage art.

[0071] According to one approach, various self-assembling and self-directed methods are employed. Elements and or their platforms can be formed from the bottom-up, one element at a time. Another advantage of bottom-up fabrication is that it reduces the amount of superfluous material that surrounds each cargo element, reducing the element's exposure to contaminant background radiation and thereby improving the functional effectiveness of the element.

[0072] In one embodiment, the instant application teaches one or more nanoscale elements of one or more types formed from isolated, synthetic and or recombinant amino acid residues comprising in whole or in part one or more types of Clathrin and or Coatomer I/II proteins of one or more isoforms, including cloned isoforms. The efficacious elements may comprise minimalist, non-cage forming elements

in one or more embodiments. In other embodiments, one or more Clathrin or Coatomer cage elements comprise efficacious elements.

[0073] In one embodiment, one or more elements may additionally comprise a hybrid molecular element formed from one or more other types of molecules.

[0074] The instant invention teaches that in one or more non-cage element embodiments it features unique types of dynamic properties and capabilities not found in fully self-assembled cavity-forming cage structures as taught in the prior art.

[0075] In one embodiment, an element is comprised of one or more 3-legged triskelia, each triskelion having 6 protein subunits; 3 Clathrin heavy and 3 light chain subunits. In another example embodiment, the instant invention teaches one or more configurations as being comprised of only 3 Clathrin heavy subunits or only 3 light chain subunits. In another illustrative embodiment, configurations comprised of less than 3 Clathrin heavy or 3 light chain subunits are enabled. In another embodiment, the invention teaches elements comprising in part one or more types of Clathrin and or Coatomer I/II proteins of one or more isoforms

[0076] Likewise, the invention teaches one or more highly flexible element embodiments formed from Coatomer I/II proteins. In one embodiment, one or more nanoscale elements of one or more types are formed from isolated, synthetic and or recombinant amino acid residues comprising in whole or in part one or more types of Coatomer I/II proteins of one or more isoforms, including cloned isoforms. Components of both COP1 and Clathrin-adaptor coats share the same structure and the same motif-based cargo recognition and accessory factor recruitment mechanisms, which leads to insights on conserved aspects of coat recruitment, polymerization and membrane deformation. These themes point to the way in which evolutionarily conserved features underpin these diverse cell pathways.

[0077] In one example embodiment, one or more elements comprised of Coatomer (COPI and COPII) proteins, which can efficaciously act alone or with additional elements, are used instead of Clathrin proteins, preferably in those applications where Coatomer characteristics would be more desirable than those of Clathrin. Coatomer I/II protein elements may, in one or more embodiments, be comprised of one or more alpha, beta, beta', gamma, delta, epsilon and or zeta subunits. Different combinations of these subunits are known to exist within Coatomer complexes. According to an illustrative embodiment, a Coatomer subunit is a nanoscale element. In one invention embodiment, Clathrin and Coatomer elements and one or more methods may be used together in one or more configurations, taking advantage of their respective capabilities.

[0078] Freed from the constraints of only assembling into cavity forming cages in vitro, one or more non-cage elements of one or more types may self-assemble into one or more other types of complex elements and or material forms, enabling new classes of applications. For example, but not limited to, using techniques known in the art, bioengineered strands of Clathrin and or Coatomer proteins form functionalized nano-tubules (Zhang, et al. 2007) for biomedical applications and bio-molecular components. In another bioengineered embodiment, invention elements comprise repeatedly branched, highly symmetrical structures, forming protein-based nano-dendrimers suitable for biomedical and bio-molecular applications. In another embodiment, self-

assembling, stable, bioactive, protein-based, hydrogel nanoparticles (i.e., nanogels), some with tunable structural properties, are enabled. Generally, hydrogels are of interest to the biomedical field, e.g., for treating trauma, because the hydrated networks can provide a physiological environment where biological species can survive or grow. In other embodiments, one or more other types of non-cage forming structures, elements, and forms of materials comprised of invention elements are formed using techniques known in the art.

[0079] Unlike cage, cavity, and vesicle systems in the prior art where one or more additional elements, e.g., cargo, are inserted into a complex, fully formed structure; a sometimes difficult and laborious process; the invention, in one embodiment, teaches that it can be functionalized with one or more additional elements at a much more fundamental nano-element level, e.g., by using non-cage elements of one or more types formed from amino acid residues of Clathrin or Coatomer proteins. Such functionalized, minimalist elements may further self-assemble in vitro into one or more nanoscale structure elements, including cages. This makes the addition of one or more elements easier and simpler as there is no insertion process into a completely formed cage, cavity, or vesicle. In another embodiment, additional element functionalization is simplified by decorating just the external surface of a cage.

[0080] According to one illustrative configuration, one or more types of elements, such as cargo elements, may interfere with the invention's overall operation if carried in the same element as other element types. Instead, the problematic elements are carried in a separate element that exclusively carries non-interfering elements, thereby inhibiting disruptive interference of invention operations. Such non-interfering elements may be functionally and or physically linked with other elements carrying other element types.

[0081] In one embodiment, one or more elements efficaciously operate alone and carry no additional elements, e.g., cargo. In one embodiment, such solo element functionality produces a unique new type of efficacious element, and its unique features correspondingly enable new types of applications.

[0082] Some embodiments include a molecule having an unpaired electron, a transition metal ion, which can be found in the active centers of many proteins (metalloproteins), or a material having any defect that produces an unpaired electron

[0083] According to one in vivo application for enhanced medical imaging, paramagnetic lanthanide, transition metal ion complexes, and the like are cargo elements that modify the NMR relaxation times of nearby proton nuclei of H2O molecules, leading to brighter images and enhanced contrast between areas comprising the contrast agent and the surrounding tissues.

[0084] In another illustrative embodiment, one or more elements accept free radical molecules such as nitroxide molecule spin labels for electron paramagnetic resonance (EPR) based invention applications.

[0085] In another illustrative embodiment, one or more elements accept and or comprise one or more types of labels and assay strategies, and instruments for detection of one or more such labeled and or assay elements may include, but are not limited to: fluorescence and confocal microscopy, flow cytometry, laser scanning cytometry, fluorescence microplate analysis and biochips, immunoassay systems,

nucleic acid-based diagnostics, and the like. In various embodiments, one or more elements meet and or surpass the requirements for label and assay sensitivity, accuracy and convenience.

[0086] In another embodiment, one or more types of elements such as comprising in whole or in part one or more large molecule elements, small molecule elements, cargo elements, agent elements, device elements, drug elements, and the like, enter the CNS, including passing the blood brain barrier, in 30 minutes or less and or in 30 minutes or more, post administration, and, depending on cargo and other element type, may require minimal functionalization for such element passage.

[0087] In some configurations, one or more elements comprise a cargo element, while in other configurations they comprise multiple elements, of one or more types. In some configurations, one or more or each of the elements and or cargo elements is a metal, and or may include one or more metals. Alternatively, each of the elements and or cargo elements is or includes non-metal elements. In other embodiments, elements and or cargo elements are exclusively non-metal elements that may include gases, as well as other elements like biological elements, drugs, optics, polymers, etc. In another embodiment, one or more elements and or additional elements comprise one or more types of material forms, including a solid, gas, vapor, crystal, and the like. In another embodiment one or more invention and or non-invention elements, in one or more combinations, comprise one or more types of isolated, synthetic and or recombinant elements.

[0088] An invention element, in one functionalized configuration, includes receptor molecules; natural, isolated, synthetic and or recombinant, for capturing and ordering the placement of one or more elements, like cargo elements, on one or more elements.

[0089] An invention element, in another functionalized configuration, includes adapter molecules; natural, isolated, synthetic and or recombinant, disposed between the receptor molecules and one or more elements to couple the receptor molecules to another element, like to a cargo element.

[0090] An invention element, in one functionalized configuration features ligands, natural, isolated, synthetic and or recombinant, including drugs, of one more types attached to receptors and or adapter protein elements.

[0091] In one configuration, one or more elements, of one or more types, are attached to one or more types of amino acids on one or more elements.

[0092] In another configuration, biotin-avidin is used as a coupler of one or more elements, of one or more types, to one or more elements of one or more types.

[0093] In another configurations, PEGylation, a cross-linker, molecular bridge, molecular tether, and the like are used to attach one or more elements, of one or more types, to one or more elements of one or more types.

[0094] In one example, molecules of one or more types are attached to a short molecular tether to one or more elements via site directed substitution mutagenesis, followed by reaction of a unique amino acid group with a specific molecular label.

[0095] In another embodiment, free radicals, toxic elements, other types of undesirable elements and the like circulating within an in vivo environment are scavenged via molecular tethers, via other elements of one or more types

attached to one or more invention elements, and or via direct binding to one or more elements.

[0096] In another embodiment, the invention takes full advantage of protein flexibility and plasticity to create elements of one or more types that are bonded, fastened, fused, and or affixed to one or more other elements, of one or more types.

[0097] In one illustrative embodiment, one or more elements and or bonded elements are coated in whole or in part with other elements, such as chemical, biological and or metallic materials, and the like. The coating elements may be or include organic, inorganic, and or synthetic materials, or a combination thereof.

[0098] In another invention embodiment, site directed mutagenesis is used to incorporate one or more elements, of one or more types, into one or more other elements, of one or more types.

[0099] In one embodiment site-directed mutagenesis using one or more types of primer; including its reverse complement; are used to insert one or more DNA sequences of one or more types into one or more coding regions of one or more elements.

[0100] In another embodiment, cloning is done of one or more genes encoding one or more elements. In another embodiment, one or more amino acids and or their encoder gene are controlled, regulated, modified, and the like, by one or more methods known in the art to produce an efficacious effect, in vivo and or in vitro.

[0101] In one embodiment, one or more elements of one or more types comprise targeted and or non-targeted drug elements, biological elements, other forms of healthcare elements, including cosmetic elements, in one or more configurations or combinations, for diagnosing, remedying, inhibiting, mitigating, curing, and or preventing one or more types of diseases, infections, physical or mental trauma, other forms of physical and mental afflictions, and the like, of one or more types, including types featuring minimal immunogenic and or toxic effects.

[0102] In one embodiment, one or more elements are used as a means for evaluating drug advancement and efficacy.

[0103] The invention teaches a biological model and or method that is consistent from a minimalist component level up, e.g., amino acid residues comprising in part one or more Clathrin and or Coatomer I/II proteins of one or more isoforms, making drug discovery safer, more efficacious, more time and cost effective, and overall, a much more rapid process.

[0104] In one personalized medicine embodiment, the invention reduces drug side effect profiles and or produces greater agent efficacy, as well as excludes agents that may have no efficacy in a particular individual. The invention, in one embodiment, provides for individual patient factors such as genotype, phenotype, age, gender, ethnicity etc., to be taken into account by one or more elements and factored into dosing and administration consideration.

[0105] In one embodiment, one or more elements comprise one or more types of pluripotent stem cells and or comprise one or more stem cell delivery methods.

[0106] According to one feature, one or more elements may be or include one or more research, therapeutic, diagnostic, vaccine, assay, and or prosthetic agents, in one or more configurations, and thereby constitute one or more types of biomedical elements. Such biomedical elements may be, for example, nano-structured and/or include chemi-

cal, biological and/or metallic materials. The biomedical elements may be or include organic, inorganic, and or synthetic materials, or a combination thereof.

[0107] Medical, biomedical, bioengineered, and or biological applications and platforms of the instant invention may include, but are not limited to, imaging; sensor; genetic and protein assay; diagnostic; drugs and drug delivery; prosthetic; inter- and extra-cellular tissue; whole organ; circulatory system; medical device; implantable defibrillator; pacemaker; coronary stents; angioplasty device; and other like applications.

[0108] In one embodiment, one or more elements comprise one or more applications that perform analysis, of one or more types, of disorders of complex inheritance.

[0109] In one embodiment, one or more elements comprise one or more applications that perform analysis, of one or more types, of pharmacologic therapy.

[0110] In one embodiment, one or more elements comprise one or more types of prognosis and therapy selection—"theradiagnostics".

[0111] In one embodiment, one or more elements comprise one or more genomic applications of one or more types.

[0112] In one embodiment, one or more elements comprise one or more oncology applications of one or more types.

[0113] In one or more embodiments, one or more elements may use routes of administration comprising one or methods of one or more types, such as those defined by CDER Data Element Number C-DRG-00301 in the US FDA Data Standards manual. Routes of in vitro administration of one or more elements may also comprise one or more forms.

[0114] In one or more embodiments, one or more pharmaceutical and drug formulations of one or more types are used, in whole or in part, such as tablet, capsule, soft galantine capsule, topical, injections, eye drops, syrups and liquids, soap and cosmetics, birth control device, and the like, but not limited to, as well as one or more types of biologics, chemical compounds, water soluble compositions, and the like, but not limited to. In vitro formulations may also comprise one or more formulations of one or more types in one or more embodiments.

[0115] According to one feature, one or more elements respond to one or more external and/or internal stimuli, which can be, for example, mechanical, chemical, biological, metabolic, covalent, non-covalent, photonic, sonic, acoustical, thermal, fluidic, electromagnetic, magnetic, radioactive, quantum mechanical, or electrical in nature. Examples of such a stimulus response is altering a cargo element carried by an element; the altering of the element itself; causing changes in cellular process like endocytosis, exocytosis, mitosis, trafficking and signaling, and the like, including other conformational changes.

[0116] In another embodiment, photonic energy impacting one or more elements produces electrical current, and or photonic energy, e.g., a laser.

[0117] In general, in another embodiment, one or more element and or platform are physically and/or functionally cooperative with other suitable types or forms of elements, agents, organisms, materials, substances, components, devices, and or systems, including non-invention elements, in vitro and/or in vivo.

[0118] The invention, in one embodiment, provides for a plurality of elements comprising aggregated, complex self-

assembled nanoscale structures that dynamically bind together one or more types of endogenous, exogenous, homogeneous, and or heterogeneous elements into one or more complex elements, which also may have one or more payload types.

[0119] The invention, in one embodiment, provides a capability for in vivo and in vitro integration of one or more types of elements into other elements, devices and mechanisms, some of which may also be non-invention elements, that also may be linked together functionally or logically, including with other devices and or operators, locally or at a distance, significantly enhancing the overall capabilities of the invention.

[0120] In one embodiment, the invention provides for the ability of one or more elements to track, recognize, attack and or destroy multiple targets on the fly, in vivo and in vitro, using dynamic target prioritization for a single element type and or multiple element types.

[0121] In one application, one or more elements, including cargo elements, comprise one or more types of targeted agent delivery systems and or agents in vivo or in vitro, including high precision dosing, using, as appropriate, ligands, targeting moieties, and or other vectors. In one application, one or more targeted elements comprise one or more research, remedial, inhibitory, mitigation, preventive, prosthetic, assay, and or other type of bio-molecular agent or device, in one or more combinations, and may altogether comprise a unified element and or platform.

[0122] The invention, in one embodiment, provides for a method for targeted delivery systems that leverage and utilize biological control laws and that may act as self-directed systems.

[0123] According to another invention embodiment, one or more targeted elements may use molecular-imprint technology, which is used for the production of molecule-specific cavities that mimic the behavior of receptor binding sites, without the temperature sensitivity of natural systems.

[0124] According to another feature, biodegradable films may also be used as a pliable template for one or more targeted elements, which are pressed into a biodegradable film and then removed, leaving a physical mold of the element's shape. The film can then be hardened and used by an element to detect a particular element, which may be, but is not limited to, a particular receptor, protein, or cell, since its complex imprint shape on the film will bind only to that particular biological element.

[0125] In one embodiment, the invention provides for a targeting system using biodegradable nanocapsules for delivery of one or more elements in vivo or in vitro.

[0126] In another application, a nanoscale platform comprised of a plurality of elements performs molecular-level and or cellular-level target site loitering, monitoring, repair, construction and or dynamic, interactive control and regulation of biological systems, in vitro and in vivo.

[0127] In another embodiment, one or more elements, including in whole or in part one or more non-invention elements, operating alone or with one or more additional elements, comprise one or more types of membrane fusion elements. In one embodiment, the resulting biological processes and interactions from such fusion may lead to a series of controlled, regulated, extended, modulated, purposefully, and or self-directed methods and or behaviors of elements.

[0128] In one example embodiment, one or more elements

in whole or in part execute one or more types of actions

involving conformational changes, bonding, attachment, and or the fusion of one or more elements to a cell membrane, one or more of which actions may lead to changes in cellular processes, such as endocytosis, exocytosis mitosis, trafficking and signaling, and the like, and or enable the precise dispatch and sequenced delivery of selected agents from an element to a target cell. Alternatively, a series of interlocking steps between a part of a cell membrane, and all, or a subset of the materials comprising an element may cause the cessation of one or more element's delivery to a target cell, and or enable delivery from other sources.

[0129] In another configuration, one or more elements dynamically respond to natural environmental conditions and manifest special functions. The various control laws that regulate biochemical reactions and physiological processes often display features that allow biomolecules or biological structures to perform more tasks than are reasonably expected from a simple mechanical device. In one embodiment, the invention takes deliberate advantage of these biological control laws. Via the use of bio- and genetic engineering methods known in the art, the invention makes use of these control laws to dynamically regulate complex in vivo and in vitro biochemical reactions and physiological processes. An example of biological control laws at work is the automatic self-directed, self-assembly of in vitro and in vivo Clathrin and Coatomer proteins.

[0130] In one embodiment, intramolecular dynamics of biomolecules and the concerted and interlocking steps of conformational changes lead to deliberately purposeful actions. For example, one or more elements may fit spatially and each step in a process fits temporally (kinetically) with an element of anticipation of the purposeful outcome.

[0131] In another example case, the spatially and temporally defined events between the cell and one or more elements may cause the invention to release diagnostic and monitoring agents to determine the most appropriate course of therapeutic action. The calculated utilization of biological control laws by one or more elements may, for example, provide for a sophisticated drug delivery system that provides optimal dosing by altering its drug delivery behavior, as well as producing minimal side effect profiles.

[0132] A further advantage of the invention is that it provides elements that can be bio-engineered to prevent in vivo uptake by one or more types of organs, tissue, cells, and bone. In the converse, another advantage is that one or more elements can be bio-engineered for highly selective uptake by one or more types of targeted cells, tissue, organs, bone, as well as by other organic and inorganic matter. In another embodiment, one or more elements comprise a non-selective uptake, non-targeted drug delivery system.

[0133] In another embodiment, the invention provides for the ability of one or more elements to intelligently monitor, control and regulate, react, and further adjust biological processes after delivery of the payload, enabling high precision dosing.

[0134] Another advantage of the invention is that Clathrin can cross cell membranes including the blood brain barrier (Gragera et al 1993) and can move through the synaptic clefts (Granseth et al 2007). In one embodiment, bioengineered Clathrin actively transports substances in and out of cells including neurons and blood brain barrier cells.

[0135] In another embodiment, one or more elements, operating alone or with one or more additional elements, comprise one or more types of cell membrane crossing

elements and gain access to the cytosol and intracellular elements of one or more types, including one or more cell organelles. Such elements may, in one embodiment, require minimal functionalization to cross the cell membrane and or enter a cell organelle.

[0136] In one embodiment, one or more elements, in whole or in part, in one or more combinations, take one or more actions to create, spawn, comprise, modify, regenerate, reassemble, and or control and regulate one or more cells, cellular elements and or cellular processes of one or more types.

[0137] In one embodiment, one or more elements, in whole or in part, in one or more combinations, take one or more actions to rectify and or repair failures and defects in cellular processes, such as, endocytosis, exocytosis, mitosis, trafficking and signaling, and the like. Such failures and defects can lead to diseases, for example, cancer.

[0138] In one embodiment, one or more elements comprise in situ in vivo elements for remediation, removal and or sequestration of one or more types of contaminants, toxins, undesired organic or inorganic elements, and the like. [0139] In one embodiment, one or more elements comprise in situ environmental elements for remediation, removal and or sequestration of one or more types of in vitro environmental contaminants and or toxins; for example, chlorinated solvents TCE, PCE, PCBs, c-DCE, DNAPL, heavy metals (chromium), biofilm, synthetic chemicals, and the like.

[0140] In one embodiment, some or all elements may also operate under the control and influence of other in vitro and or in vivo elements, including non-invention elements, and altogether may comprise a scalable, nanoscale platform.

[0141] In general, in another aspect, the invention is directed to a method of forming one or more types of scalable platforms, including the steps of providing one or more embodiments of the elements to deliberately carry out a series of tasks of one or more types, which tasks and or methods may be externally directed or internally self-directed, or a combination thereof. In other embodiments, one or more nanoscale platforms may be additionally comprised of one or more non-invention elements and platforms of one or more types.

[0142] One or more elements, in one platform embodiment, may also modify, process, manipulate, encode and decode, input, output, transmit, communicate, store and or read information using techniques and methods known in the art, in vivo and in vitro.

[0143] In one embodiment, scalable information processing platforms use some or all elements as bits that are programmable into a plurality of logical states. In another configuration, the invention features a scalable information-processing platform that may include one or more elements.

[0144] As a general characteristic, one or more elements may take any suitable form, and multiple embodiments may be used as elements, and or further combined in any suitable manner to create one or more cargo carrying and or non-cargo carrying nanoscale elements ("elements"), and or multifunction nanoscale platforms ("platforms") of one or more types, operating in vitro and or in vivo, such as: multiple polypeptide elements and platforms; biological elements and platforms; large molecule elements and platforms; small molecule elements and platforms; biomedical elements and platforms; medical elements and platforms; diagnosis, cure, mitigation, treatment, prevention of disease

or other type of drug elements and platforms; targeted and or non-targeted delivery elements and platforms; cell, cell organelles, or cell material crossing elements and platforms; personal medicine elements and platforms; elements and platforms that, post administration, in whole or in part enter the central nervous system, including passing the blood brain barrier in 30 minutes or less and or in 30 minutes or more; healthcare elements and platforms; reproductive health elements and platforms; substance abuse disorder treatment elements and platform; bioengineered elements and platforms; cosmetic elements and platforms; agricultural elements and platforms; sensor elements and platforms; research and development elements and platforms; scientific elements and platforms; crystal elements and platforms; electronic elements and platforms; photonic energy elements and platforms; information processing or storage elements and platforms; energy storage elements and platforms; in situ elements and platforms for remediation, removal and or sequestration of undesirable elements and platforms of one or more types; quantum mechanical elements and platforms; telecommunication elements and platforms; and the like; one or more of which nanoscale elements and platforms may be additionally comprised of one or more non-invention elements and platforms of one or more types, and with or without one or more types of cargo elements located on and or in one or all or a subset of

[0145] In general, in a further aspect, the invention is directed to a method of forming one or more formations of nanoscale elements formed in vitro from one or more elements of one or more types formed from isolated, synthetic and or recombinant amino acid residues comprising in whole or in part one or more types of Clathrin and or Coatomer I/II proteins of one or more isoforms, including cloned isoforms; with or without one or more additional elements of one or more types located on and or in one or more elements; forming in whole or in part one or more types of element carrying and or non-element carrying nanoscale elements and structures; one or more of which elements may also comprise one or more non-invention elements of one or more types, forming hybrid elements; wherein one or more elements, using one or more types of methods, executes one or more functions and or effects one or more ends in vivo and or in vitro.

BRIEF DESCRIPTION OF THE DRAWINGS

[0146] The foregoing and other aspects of the invention may be more fully understood from the following description, when read together with the accompanying drawings in which like reference numbers indicate like parts.

[0147] FIG. 1 is a conceptual diagram depicting a Clathrin triskelion comprised of one or more elements of one or more types employed in an illustrative embodiment of the invention.

[0148] FIG. 2 is a conceptual cross-sectional view of one or more Clathrin protein, receptor, adaptor protein, and cargo elements in an illustrative embodiment.

[0149] FIG. 3 is a computer generated frontal view of an actual Clathrin cage comprised of a plurality of Clathrin triskelia, and, in an illustrative embodiment, comprising one or more invention elements.

[0150] FIG. 4 is a flow diagram depicting conceptually the formation of individual Clathrin elements during endocyto-

sis, which also serves to illustrate how the instant invention operates in one or more embodiments.

[0151] FIG. 5 is a conceptual diagram depicting Coatomer I/II protein comprised of one or more subunit and domain elements of the type employed in an illustrative embodiment of the invention.

[0152] FIG. 6 is an exemplary energy level diagram 600 illustrating the energy levels associated with a hyperfine interaction between electron and nuclear spin in the presence of magnetic fields.

DESCRIPTION OF THE ILLUSTRATIVE EMBODIMENTS

[0153] The instant invention is comprised of one or more formations of nanoscale elements formed in vitro from one or more elements of one or more types formed from isolated, synthetic and or recombinant amino acid residues comprising in whole or in part one or more types of Clathrin and or Coatomer I/II proteins of one or more isoforms, including cloned isoforms, and which operate in vitro and or in vivo. In one embodiment, one or more elements form one or more configurations of one or more types, described below.

[0154] FIG. 1 is a conceptual diagram illustrating the basic unit of Clathrin, a three-leg pinwheel protein structure, and each complete leg is typically called a 'monomer'. The arrangement of the monomers in the three-dimensional protein is the quaternary structure. Each Clathrin leg monomer is further comprised of two subunits, one 190 kDa subunit ("heavy chain") and one 24-27 kDa subunit ("light chain"). Three, two-subunit Clathrin monomers self-assemble and combine to create triskelion element 100. It is this triskelion morphology that allows Clathrin to form its unique polyhedral network.

[0155] In FIG. 1, the assembled triskelion element 100 is comprised of three monomer leg elements 102a-102c. The three leg elements 102a-102c extend radially from a hub section 108. The filamentous portion of Clathrin triskelion legs 102a-102c is formed by a continuous superhelix. A naturally occurring Clathrin leg is about 47.5 nm (475 Å) long. In the instant invention, Clathrin leg length and or molecular weights can be modified and or adjusted by using bioengineering techniques known in the art.

[0156] In the case of humans, there are two isoforms each of Clathrin heavy chain (CHC17 and CHC22) and light chain (LCa and LCb) subunits, all encoded by separate genes. CHC17 forms the ubiquitous Clathrin-coated vesicles that mediate membrane traffic. CHC22 is implicated in specialized membrane organization in skeletal muscle. CHC17 is bound and regulated by LCa and LCb, whereas CHC22 does not functionally interact with either light chain. [0157] In one embodiment, a Clathrin triskelion is composed of a trimer of heavy chains 104a-104c each bound to a single light chain 106a-106c, respectively. In the case of one isoform embodiment, CHC17 (SEQ ID NO:1), a Clathrin heavy chain element is comprised of a 1675 amino acid residue protein, which is encoded by a gene consisting of 32 exons. In the case of another isoform embodiment, CHC22, a Clathrin heavy chain element is comprised of a 1640 amino acid residue protein (SEQ ID NO:2).

[0158] In one or more invention embodiments, efficacious elements formed in part from Clathrin amino acid residues include, but are not limited to, a N-terminal globular domain 110*a*-110*c* (residues 1-494) that interacts with adaptor proteins (e.g., AP-1, AP-2, b-arrestin), a light chain-binding

region (residues 1074-1552), and a trimerization domain (residues 1550-1600) near the C-terminus.

[0159] One or more of the Clathrin heavy chain amino acid sequences as described in SEQ ID NO:1 and SEQ ID NO:2, but not limited to, and in whole or in part may be modified, altered, adapted or functionalized in one or more ways in one or more embodiments of the invention.

[0160] In the illustration, the three Clathrin monomer elements 102a-102c are comprised of six subunit elements, three of which subunits are the heavy chain subunit elements 104a-104c. The three heavy chain subunits are comprised of several distinct domains and segments, one or more of which may comprise one or more invention elements in one or more embodiments, and may be functionalized via one or more techniques known in the art.

[0161] In general, each heavy chain comprises eight repeated motifs (CHCR 0-7), which make up the proximal, knee, distal and ankle segments of a Clathrin leg. The heavy-chain amino terminus folds into the terminal domain (TD) and is attached to CHCRO by a helical linker. (Brodsky, 2004). The three Clathrin heavy chains are joined at their C-termini (located within hub element 108), extending into proximal and distal leg domains ending in globular N-terminal domain elements 110a-110c, and which are responsible for peptide binding. The Clathrin heavy chain terminal domains provide multiple interaction sites for a variety of adaptor proteins (AP) that can bind multiple receptors occupied by ligands. These sites prevent chemical interactions between cargo elements. The heavy chain N-terminal domain elements 110a-110c are each comprised of a seven-bladed beta-propeller connected to a flexible linker region, respectively. This propeller domain interacts with a host of accessory proteins participating in receptor-mediated endocytosis such as adaptor proteins, non-visual arrestins and the uncoating ATPase, hsc70. The propeller domain is followed by a long filamentous segment, which is interrupted by a bent region between the distal and proximal domains, and ends in the trimerization domain at the C-terminus.

[0162] Besides harboring determinants important for driving the association of individual Clathrin molecules during lattice formation, each of the three heavy chain 104a-104c proximal domains also include binding sites for attaching the three light chain subunit elements 106a-106c, respectively, forming three complete Clathrin monomers. The three light chain subunits are also comprised of several distinct domains and segments, one or more of which may comprise one or more invention elements in one or more embodiments, and may be functionalized via one or more techniques known in the art.

[0163] Among other roles, Clathrin light chains prevent Clathrin heavy chains from interacting with each other. On the other hand, assembly proteins bind to light chains and cause a change in them such that they no longer prevent heavy chains from interacting. Clathrin light chains consist of what has been described as a linear array of domains: regions of protein discernable from the primary sequence or with distinct biochemical properties. These are an N-terminal segment, a region that is 100% conserved between light chains, a portion to which Hsc70 binds, a calcium binding domain, a region which binds the heavy chain, a site for neuronal-specific splice inserts and then finally a calmodulin-binding domain at the C-terminus domain (Royle, 2006).

The light chain C-terminal residues are also important for enhancing the in vitro assembly of hub 108 at low pH.

[0164] One or more of the Clathrin light chain amino acid sequences as described in SEQ ID NO:12 and SEQ ID NO:13 but not limited to, and in whole or in part may be modified, altered, adapted or functionalized in one or more ways in one or more embodiments of the invention.

[0165] In one embodiment, each of the 3 heavy chain subunits 104a-104c may each have 3 light chains subunits 106a-106c attached, respectively, forming the typical, three-monomer Clathrin triskelion structure. But in another embodiment, each leg 102a-102c may include only the 3 Clathrin heavy chain subunits 104a-104c, respectively, which is distinctly unique from the classic Clathrin monomer configuration. In yet another unique embodiment, only 3, non-attached light chain subunits 106a-106c are used.

[0166] In one distinctive embodiment of the invention, a 3-legged pinwheel configuration 100 is not enabled, and only partial pinwheel structures are used. In one embodiment, a partial pinwheel configuration of one or two legs (one or two Clathrin monomers) is comprised of one or two Clathrin heavy chains and one or two corresponding light chain subunits. In another embodiment, one or two elements comprised of only one or two Clathrin heavy chain subunits are used; e.g., subunits 102a, or 102a-102b. In one embodiment, only one or two unattached light chain subunits are used.

[0167] In another distinctive embodiment of the invention, one or more elements of one or more types are formed from isolated, synthetic and or recombinant amino acid residues comprising in part one or more types of Clathrin heavy chain and or light chain proteins of one or more isoforms as described in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:12 and SEQ ID NO:13, respectively.

[0168] In one embodiment, one or more N-terminal domain elements, e.g., 110a, 110b and or 110c are bioengineered to facilitate, modify, regulate or control peptide binding of one or more types, as well as interaction sites for one or more types of adaptor proteins.

[0169] In one embodiment, one or more domain elements of heavy chain subunits and or light chain subunits are bioengineered to facilitate, modify, regulate or control one or more Clathrin protein characteristics and or behaviors in vivo and or in vitro.

[0170] FIG. 2 is a conceptual cross-sectional view of a biological endohedral consisting of Clathrin protein elements. In this illustrative embodiment, one or more elements 102a-102c, 106a-106c, 104a-104c, 110a-110c, element 108, and or one or more types of elements formed from isolated, synthetic and or recombinant amino acid residues comprising in whole or in part one or more Clathrin proteins of one or more isoforms, and with or without one or more additional elements of one or more types, may comprise one or more multiple polypeptide elements of one more types. The latter are labeled in FIG. 2 as elements 206a, 204a, 202a, and 208a, which are formed in vitro, and also may operate in vitro and or in vivo. One or more of elements 206a, 204a, 202a, and or 208a may comprise one or more types of functionalization, include invention and non-invention elements, express one or more types of functionality, and or form one or more types of structures.

[0171] In one illustrative embodiment, but not limited to, one or more elements 206a may comprise one or more elements 102a-102c, 106a-106c, 104a-104c, 110a-110c,

element 108, and or one or more types of elements formed from isolated, synthetic and or recombinant amino acid residues comprising in whole or in part one or more Clathrin proteins of one or more isoforms, and express one or more types of functionality in one or more embodiments.

[0172] In another embodiment, one or more elements 206a may be comprised of, and or help comprise one or more types of non-invention elements, such as a natural cell element in one embodiment, comprising one or more types of hybrid elements in one or more embodiments.

[0173] In another embodiment, one or more elements 206a may be comprised of, and or help comprise one or more types of isolated, synthetic, recombinant and or natural molecules in one or more embodiments.

[0174] In one illustrative embodiment, but not limited to, one or more elements 202a may comprise cargo elements of one or more types, including natural, isolated, synthetic and or recombinant, including natural and or synthetic ligands and or drugs, and may express more than one type of functionality. In one embodiment, one or more other elements, of one or more types, including invention and non-invention elements each may bond with one or more respective cargo elements 202a.

[0175] In one embodiment, one or more cargo elements 202a are cavity forming and are non-permeable, semi-permeable, and or permeable, and or can change from one permeable state to another. In one embodiment, the cavity forming elements comprise one or more types of elements and or agents, including gas, vapor or fluid, with or without dopants. In one embodiment, one or more cargo cavities elements comprise one or more types of elements and or agents, including one or more types of metals.

[0176] In another illustrative embodiment, one or more efficacious cargo elements 202a carried on one or more elements may comprise the total functionality. In another embodiment, one or more other elements, of one or more types, including invention and non-invention elements may act in concert with one or more cargo elements 202a to achieve ensemble efficacy.

[0177] In one embodiment, but not limited to, one or more elements 204a may comprise attachment and or receptor elements for one or more elements 202a of one or more type. and or express more than one type of functionality. In one embodiment, one or more other elements, of one or more types, including invention and non-invention elements each may bond with one or more respective elements 204a. In another embodiment, receptor molecules 204a can be bioengineered to recognize and associate with specific molecules, which may also be synthetic and or natural ligands and or drugs. In another embodiment, receptor molecules **204***a* can be natural, isolated, synthetic and or recombinant. [0178] In one embodiment, but not limited to, one or more elements 208a of the instant invention may comprise the major types of adaptor elements, like the heterotetrameric adaptor protein (AP) elements, and the monomeric GGA (Golgi-localizing, Gamma-adaptin ear domain homology, ARF-binding proteins) adaptors. In one illustrative embodiment, elements 208a comprise one or more small sigma subunits of various adaptins from different AP adaptor elements. The AP complex family has six members in mammals: AP-1A, AP-2, AP-3A and AP-4 are ubiquitously expressed. The other two members, AP-5 and AP-6, are cell-type specific isoforms of AP-1A and AP-3A: the epithelium-specific AP-1B and the neuron-restricted AP-3B.

(Ohno, 2006). In another embodiment, AP180, like AP-2 and AP-3, binds to N-terminal domains 110a-110c of Clathrin. In one embodiment, one or more AP elements may be functionalized at one or more heavy chain terminal domain elements 110a-110c. In one embodiment, one or more other elements, of one or more types, including invention and non-invention elements each may bond with one or more respective elements 208a. In another embodiment, adapter molecules 208a are bioengineered to recognize specific receptor molecules and to couple the receptor molecules to Clathrin and or Coatomer protein elements. In another embodiment, adapter molecules 208a can be natural, isolated, synthetic and or recombinant.

[0179] In one embodiment, one or more elements 206a, 204a, and or 208a operate alone without cargo element 202a, and comprise one or more types of inherently efficacious solo acting elements.

[0180] In one embodiment, unlike prior Clathrin art, a plurality of elements 206a, 204a, and or 208a operate without cargo elements 202a, and comprise an inherently efficacious cage element 212 of one or more types, like a drug element, for example, which is unlike prior Clathrin art. [0181] In one embodiment, also unlike prior Clathrin art, a plurality of elements 206a, with or without one or more additional other elements comprise cage element 212, and element 212 has one or more elements, of one or more types and affixed via one or methods, located on the outside part of cage element 212; that is, located outside the cavity formed by cage 212. In another embodiment, further unlike prior Clathrin art, a plurality of elements 206a, with or without one or more additional other elements, comprise cage element 212, and element 212 has one or more elements, of one or more types and affixed via one or methods, located on both the outside, and inside parts (i.e., located within the cage cavity), of cage element 212.

[0182] According to one invention feature, cargo attachment element 204a and or element 208a shields cargo element 202a in the same element 206a from interacting. According to another feature, the shielding properties of element 206a shields and inhibits chemical and molecular interactions between it and the external environment. According to a further feature, element 206a protectively sequesters cargo elements 202a from the external environment.

[0183] In another embodiment, one or more non-invention, "natural" Clathrin elements 206b-206f (the term "natural" hereinafter generally refers to non-isolated, non-recombinant, and non-synthetic protein elements) join with one or more isolated, recombinant, and or synthetic elements; in this example, 206a; to form a natural/invention hybrid Clathrin cage element 212. In another embodiment, hybrid cage element 212 may also be comprised of natural cage element 220, which is a vesicle, forming a hybrid Clathrin Coated Vesicle.

[0184] FIG. 3 is a computer generated frontal view of a Clathrin cage 300 comprised of a plurality of natural Clathrin triskelia elements 302-308, respectively. In an illustrative embodiment, element 310 is an invention element, comprised of three heavy chain elements 104a-104c—which may or may not include three respective light chain elements 106a-106c—forming a hybrid or fused cage 300 comprised of natural elements and invention elements. In this role, element 310 comprises an efficacious replacement for a natural triskelia element.

[0185] FIG. 4 is a flow diagram 400 depicting, conceptually, the formation of a plurality of natural Clathrin elements 206b-2026f, and, in this example, along with invention element (206a) into cage 200, which at step 440, shows Clathrin coated vesicle 220. The process by which natural Clathrin molecules 206b-206d obtain natural cargo molecules 202b, 202c, and 202d in this example is known as Clathrin mediated endocytosis (CME), a process wherein a cell takes in macromolecules by forming vesicles derived from the plasma membrane. Endocytosis is crucial to cellular function. Via CME, cells internalize cargo attachment elements, transmembrane channels, transporters and extracellular ligands such as hormones, growth factors and nutrients.

[0186] In one embodiment, one or more invention elements are biologically engineered to take or induce one or more types of actions, such as to create, spawn, comprise, modify, repair, regenerate, reassemble, and or control and regulate CME, as well as exocytosis, mitosis, trafficking, signaling processes, other behaviors, and the like. Defects and disorders in any of these critical cellular processes can lead to disease, and one or more types of these processes may be modified in one or more embodiments of the instant invention, for example, to achieve therapeutic effect.

[0187] In one embodiment, the instant invention takes or induces one or more efficacious actions involving receptor-mediated endocytosis that encompass nutrient uptake (LDL, transferrin, etc.), membrane recycling, membrane protein recycling, antigen uptake, synaptic vesicle recycling, and signaling receptor down-regulation.

[0188] In one or more embodiments, one or more invention elements comprise counterparts to natural Clathrin proteins that may inherently behave as a drug; e.g., one or more invention elements are functionalized for in vivo delivery and carry no additional elements, such as cargo. Such solo acting element embodiments would interact in one or more ways with natural cells and their processes, and by so doing diagnose, regulate and or cure one or more diseases and disorders relating to endocytosis.

[0189] An increase of a cellular component is called upregulation. Upregulation is an increase in the number of receptors, e.g., see elements 204b, 204c, and 204d in FIG. 4, on the surface of target cells, making the cells more sensitive to a hormone or another agent. For example, there is an increase in uterine oxytocin receptors in the third trimester of pregnancy, promoting the contraction of the smooth muscle of the uterus. In one or more embodiments, one or more invention elements, either by acting alone and or in part with other elements of one or more types, including natural and or non-invention elements, efficaciously modify, control and regulate, interfere with, create, and or spawn elements, and or induce actions or behaviors that increase the upregulation of one or more types of receptors of the surfaces of target cells.

[0190] On the other hand there is downregulation, an example of which is the cellular decrease in the number of receptors to a molecule, such as a hormone or neurotransmitter, which reduces the cell's sensitivity to the molecule. In the literature, downregulation is the process by which a cell decreases the quantity of a cellular component, such as RNA or protein, in response to an external variable. In one or more embodiments, one or more invention elements, either by acting alone and or in part with other elements of one or more types, including natural and or non-invention

elements, efficaciously modify, control and regulate, interfere with, create, and or spawn elements, and or induce actions or behaviors that increase the downregulation of one or more types of receptors.

[0191] Exocytosis is the reverse process of endocytosis, whereby a cell directs secretory vesicles out of the cell membrane. These membrane-bound vesicles contain soluble proteins to be secreted to the extracellular environment as well as membrane proteins and lipids that are sent to become components of the cell membrane. Exocytotic vesicles are usually not Clathrin-coated; most of them have no coat at all. However, two observations suggest that Clathrin effectively 'tracks' vesicle proteins leaving a synapse. In one study (Granseth, et al, 2008) the amount of a Clathrin light chain (LC) tagged with the element mRFP leaving the synapse was proportional to the number of vesicles released by the stimulus, as assessed by the amplitude of a sypHy signal (sypHy is an improved fluorescent reporter of exocytosis). Second, in the same study the movement of LC-mRFP began without a significant delay and peaked with the sypHy signal. The movement of Clathrin out of the synapse together with synaptophysin and synaptobrevin is most easily explained as representing CME (Clathrin mediated endocytosis) of vesicles at sites removed from the active zone. This interpretation is consistent with studies showing that the machinery for CME is not at the active zone, but in the surrounding regions of membrane (Heuser & Reese, 1973; Ringstad et al. 1999; Qualmann et al. 2000; Teng & Wilkinson, 2000). Thus, Clathrin is naturally found in the extracellular space and may play a role in regulating exocytosis and or endocytosis. In one or more illustrative embodiment, one or more elements of one or more types may efficaciously operate in inter- and or extra-cellular spaces of one or more types; for example, perform remediation, sequestration, or removal of one or more types of undesirable elements.

[0192] Membrane trafficking only occurs during interphase. As the cell enters mitosis, Clathrin-mediated membrane traffic is rapidly shut down and only resumes in late telophase. Clathrin may therefore have a separate function that is distinct from membrane trafficking, which operates during mitosis. Clathrin is thus a multifunction protein: during interphase its function is in membrane trafficking and during mitosis it has a role in stabilizing spindle fibers (Royle, 2006). In one invention embodiment, mitosis may be efficaciously controlled and regulated, modified, and or induced via one or more methods and instances of the instant invention.

[0193] In another embodiment, one or more elements are comprised of, but not limited to, one or more isolated, synthetic, and or recombinant adaptor protein molecules, tubulin protein molecules, dynamin protein molecules, epsin protein molecules, endophilin protein molecules, synaptotagmin protein molecules, and or other types of protein molecules associated with Clathrin and Coatomer proteins and processes, for efficacious effect.

[0194] In another embodiment, one or more natural adaptor protein molecules, tubulin protein molecules, dynamin protein molecules, epsin protein molecules, endophilin protein molecules, synaptotagmin protein molecules, and or other types of protein molecules involved with associated with Clathrin and Coatomer proteins and processes form efficacious hybrid elements when also comprised of one or more types of invention elements.

[0195] The CME process involves a dynamic interaction between Clathrin and a wide range of other protein molecules, and altering the compositions and behaviors of the various molecular parties involved. For example, the cell uses endocytosis to control and regulate the density of receptors on the cell surface and to acquire nutrients. Endocytosis of ligand-activated cargo attachment elements is essential for the proper attenuation of a variety of signal transduction processes, as well as for co-localization of activated cargo attachment elements with downstream signaling molecules. Endocytosis also counterbalances secretion, preventing continuous expansion of the plasma membrane. Endocytosis thus internalizes macromolecules and fluid, and after sorting, directs the internalized molecules for degradation or recycling.

[0196] The endocytosis process begins when proteins bound to cargo attachment elements accumulate in coated pits 404, which are specialized regions of the cell membrane 402 where it is indented and coated on its cytoplasmic side with a bristle-like coat composed of two natural proteins: Clathrin and protein adapters. Most, if not all, intracellular transport vesicles are encased in a proteinaceous coat, one class of which is Clathrin-coated vesicles (CCVs). CCVs also mediate the transport of lysosomal hydrolases from the trans-Golgi network, as well as the efficient internalization of extracellular solutes such as nutrients, hormones, growth factors, and immunoglobulins at the plasma membrane.

[0197] Clathrin also transports proteins from the Golgi to other organelles. In neurons, endocytosis is critical to allow rapid synaptic vesicle regeneration. Besides Clathrin, there are other coat-forming proteins, such as COP I and COP II, which mediate intracellular traffic and there are Clathrin-independent endocytic pathways which mediate internalisation of a variety of cargo (Royle, 2006).

[0198] In one invention embodiment, the natural endocytosis process is transformed into a versatile therapeutic method to regulate the intensity, localization, half-life and function of signaling elements (signalosomes) that form in cells upon, for example, binding of growth factors, cytokines and morphogens to their cognate receptors. In one example embodiment, the invention rectifies breakdowns in the function of endocytic adaptors that might facilitate impairment of tissue homeostasis and consequent tumor development. In another illustrative embodiment, one or more invention elements, acting alone or not, interact with natural adaptor proteins required for appropriate receptor downregulation and which play distinct roles in oncogenesis. (Crosetto, et al. 2005) In another embodiment, CME elements might also comprise one or more invention cargo elements (202a in FIG. 4), which can be drugs, other ligands, and the like.

[0199] In one embodiment, referring to FIG. 4, a natural Clathrin coated vesicle 220 is desired to form to endocytose over-expressed natural receptor elements 204b and 204c that are initially located outside cell membrane 402. The appearance of one or more types of invention elements, such as element (206a) in the illustrative example, outside cell membrane 402 and or by crossing 402, dynamically begin to create, induce, spawn, mediate, control and regulate, regenerate, and or interact with one or more natural endocytosis processes and behaviors. With the prompting of one or more types of invention Clathrin elements, one or more biological processes acting on cell membrane 402 induce a Clathrin bud 404 to form at 420.

[0200] As shown at 430 and 440, after forming completely around bud 404, natural Clathrin elements 206b-206d pinch off (scission) from membrane 402 with the desired over expressed receptors 204b and 204c held inside vesicle 220. After excision, bud 404 has evolved into a plurality of natural Clathrin elements 206b-206f, some of which are attached to one or more types of over expressed receptor elements 204b and 204c, as well as attached to other receptor elements; which in this example are the normally expressed natural elements 204d.

[0201] In one illustrative embodiment, the otherwise allnatural plurality of Clathrin elements in FIG. 4 includes one or more non-cargo carrying; solo acting invention elements (206a), forming a "hybrid" CCV 440 with the desired efficacious properties and behavior. This hybrid CCV then follows normal pathways within the cell, causing downregulation of the desired over-expressed receptor elements, which may be associated with one or more types of neurotransmitters, viruses, cholesterol, as well as with other cargo types, restoring a cell to its normal, healthy state.

[0202] In another illustrative embodiment, natural Clathrin coated vesicle structure 440 in FIG. 4 is additionally comprised of one or more non-cargo carrying invention receptor element 204a and or adaptor element 208a (as illustrated in FIG. 2), forming a hybrid or fused Clathrin coated vesicle 440 in FIG. 4, with the desired efficacious properties and behavior. In another embodiment, one or more hybridized and or invention elements may enter the cell nucleus and or other organelles and cell elements.

[0203] The fusion and or participatory actions of one or more non-additional element carrying, solo acting invention elements 206a, 204a, and or 208a in FIG. 2 may yield a therapeutic effect, and are an example embodiment of inherently efficacious invention elements in action. In another embodiment, natural or hybrid CCV 440 in FIG. 4 also includes one or more invention cargo molecules (202a) that may have been transported into the cell via their attachment to one or more natural and or invention receptor elements.

[0204] Referring again to FIG. 4, in another example embodiment, a therapeutic effect is accomplished via one or more invention elements by regulating EGFR (epidermal growth factor receptor), which exists on the cell surface and is activated by binding of its specific ligands including epidermal growth factor and transforming growth factor α (TGF α).

[0205] When these natural cargo attachment elements are activated, cells rapidly clear them from the surface and destroy them. Control of EGF receptor signaling is performed by Clathrin-mediated endocytosis. Natural Clathrin coats also exist on endosomes and are involved in endosomal sorting of the EGFR. A defect in this overall process will likely lead to uninhibited growth of cells and tumors. EGFR expression, over-expression, or mutation is associated with cancer progression, advanced disease, drug resistance, aggressive disease, poor prognosis, and reduced survival. EGFR is considered one of the main proteins elevated in breast, lung, and prostrate cancers, among others. Brain cancer is also implicated with over-expressed EGFR. Other work has shown that using monoclonal antibodies for EG1-R, or anti-EGFR, has proven an effective strategy for getting nanoparticles to specifically attach themselves to cancer cells. Additional work has shown effectiveness of EGFR as the cancer-targeting pathway. In one embodiment, CME, cell fusion, cell penetrating, and or one or more types of other participatory actions of one or more solo operating, efficacious invention elements 206a, 204a, and or 208a in FIG. 2 may yield a therapeutic effect in controlling, regulating, or mediating EGFR activity. In another example embodiment of modulating EGFR activity, cargo elements (202a) in FIG. 4 may comprise one or one or more types of cancer drugs or biologicals delivered directly into cells and organelles that are transported into the cell via their attachment to one or more natural and or invention receptor elements during CME, by cell fusion, by directly penetrating cell membrane 402, and or by one or more types of other participatory actions. In another embodiment, invention cargo elements (202a) may comprise one or more diagnostic agents, or combine one or more diagnostic agents and therapeutic agents in the same payload. In one or more embodiments, one or more invention elements of one or more types may thus comprise an efficacious method for the diagnosis, treatment, remedying, curing, and or prevention of one or more types of cancers, including those cancer types that fall outside the scope of EGFR-related activity.

[0206] FIG. 5 is a conceptual diagram illustrating the basic units of Coatomer I and II proteins. COPII and Clathrin cages are both constructed from ∂-solenoid and β-propeller building blocks (Fotin et al., 2004b; ter Haar et al., 1998; Ybe et al., 1999). In various embodiments of the invention, one or more elements of one or more types are formed from isolated, synthetic and or recombinant amino acid residues comprising in whole or in part one or more types of Coatomer proteins of one or more isoforms, including cloned isoforms. Examples of various Coatomer subunit amino sequences are listed in SEQ ID NO:15, SEQ ID NO:18, SEQ ID NO:21, SEQ ID NO:26, SEQ ID NO:27, SEQ ID NO:29, and SEQ ID NO:30. In another embodiment, one or more Coatomer subunit amino acid sequences may be modified, altered, adapted or functionalized in one or more ways in one or more embodiments of the invention.

[0207] In one embodiment, Coatomer is comprised of seven distinct subunits: alpha, beta, beta', gamma, delta, epsilon and zeta subunits, respectively.

[0208] In Clathrin, a triskelion assembly unit lies at each vertex, and the $\[\partial \]$ -solenoid legs of neighboring triskelia interdigitate extensively as they extend toward the adjacent vertices; the $\[\beta \]$ -propeller is not part of the architectural core and instead projects in toward the membrane to interact with adaptor molecules (Fotin et al., 2004; Kirchhausen, 2000). In contrast, the COPII assembly unit is a rod that constitutes the edge of a cuboctahedron, and four rods converge to form the vertex with no interdigitation of assembly units. $\[\partial \]$ -solenoid domains form the core of the edge, but, unlike Clathrin, the COPII vertices are formed from $\[\beta \]$ -propellers. In summary, the COPII and Clathrin lattices seem not to share common construction principles other than the use of $\[\partial \]$ -solenoid and $\[\beta \]$ -propeller folds.

[0209] Crystallographic analysis of the Coatomer II assembly unit reveals a 28 nm long rod, element 502, comprising a central solenoid dimer capped by two β propeller domains, elements 504, at each end. GTPase, elements 508, bind to adaptor elements 506, which bind to elements 502. In the illustration, element 502a is an invention element that acts as an efficacious replacement element for one or more natural element 502, forming a hybrid Coatomer element. The structural geometry and properties of COPI coats remain to be determined. However, by analogy to the COPII and Clathrin structural units, they

probably involve a preassembled cage protein (CP) scaffold that is generated by the β-propeller-containing and ∂-sole-noid-containing subunits and an adaptor protein (AP) sub-complex. Together these could form an AP-CP heptaheteromeric functional unit in the cytosol. (Gurka, et al. 2006)

[0210] COPI and COPII play a major role in exocytosis, as also can their invention element counterparts. Clathrin can also play a role in exocytosis, but to a lesser extent than Coatomer. The exocytosis process refers to the fusion of intracellular vesicles with the plasma membrane. It occurs via two major processes, a constitutive pathway and a regulated pathway. These are the major ways that the cell secretes materials, wherein a cell secretes macromolecules (large molecules) by fusion of vesicles with the plasma membrane. Coatomer-coated vesicles, which are typically less than fifty nanometers in size, are also involved in vesicular transport between the Golgi apparatus, endoplasmic reticulum and plasma membrane. Coatomer I vesicles shuttle elements from the Golgi to the endoplasmic reticulum (ER). Coatomer II vesicles shuttle elements from the ER to the Golgi. Coat-protein I/II subunits (COPs) require ATP to assemble into a coat and unlike Clathrin coats, the Coatomer coat remains on the vesicle until docking occurs. In some instances, Coatomer proteins are also involved in endocytosis, but are unrelated to Clathrin. Thus, while Clathrin also mediates endocytic protein transport from the ER to the Golgi, Coatomers (COPI, COPII) primarily mediate intra-Golgi transport, as well as the reverse Golgi to ER transport of dilysine-tagged proteins. Coatomers reversibly associate with Golgi (non-Clathrin-coated) vesicles to mediate protein transport and for budding from Golgi membranes. In one or more embodiments, one or more COPI/ COPII invention elements and or Clathrin invention elements, either by acting alone and or in part with other elements of one or more types, including natural and or non-invention elements, efficaciously modify, control and regulate, interfere with, create, and or spawn elements and or induce actions or behaviors involving exocytosis.

[0211] Cells of the mammalian immune system undergo selective changes in protein glycosylation during differentiation, immune activation, and autoimmune disease. In many, if not most of these types of diseases endocytosis and cellular trafficking and signaling plays a role. Referring again to FIGS. 1, 2, 3, 4, (and 5, in some embodiments), but not limited to, in one embodiment, one or more invention elements of one or more types, in whole or in part selectively interfere with, fuse with, control and regulate, induce, and otherwise modify endocytosis, receptor-specific processing, trafficking and signaling, and other behaviors for efficacious effect in one or more types of autoimmune diseases, including, but not limited to, one or more types of diabetes, CNS autoimmune diseases, and other types of autoimmune diseases that effect the body.

[0212] Referring again to FIGS. 1, 2, 3, 4, (and 5 in some embodiments), but not limited to, in one embodiment, one or more invention elements of one or more types selectively interfere with, control and regulate, and or modify secretory products that participate in inflammation and immunoregulation; and also in other embodiments, whereby endocytosis mediated by specific receptors for immunoglobulin or by other opsonins is important in removal of damaged self or foreign particles. In another embodiment, defects in membrane receptor function, whether inherited or acquired, and

the pathogenesis of immune diseases may be remedied, inhibited, mitigated, and or prevented.

[0213] Referring again to FIGS. 1, 2, 3, 4, and 5, in one embodiment, but not limited to, one or more invention elements of one or more types efficaciously fuse with and or functionally replace one or more natural elements commonly found in endocytosis, exocytosis, mitosis, trafficking and signaling, and the like, either by acting alone and or in part with other elements of one or more types, including natural and or non-invention elements.

[0214] Referring again to FIGS. 1, 2, 3, 4, and 5, but not limited to, in another embodiment, one or more invention elements of one or more types efficaciously cross over into a cell, its elements, and or its organelles, such as its nucleus, either by acting alone and or in part with other elements of one or more types, including natural and or non-invention elements.

[0215] Referring again to FIGS. 1, 2, 3, 4, and 5, in another embodiment, but not limited to, one or more invention elements efficaciously create, spawn, comprise, modify, repair, regenerate, reassemble, and or control and regulate one or more natural elements commonly found in endocytosis, exocytosis, mitosis, trafficking and signaling, other cellular behaviors, and the like, either by acting alone and or in part with other elements of one or more types, including natural and or non-invention elements.

[0216] Referring again to FIGS. 1, 2, 3, 4, and 5, in another embodiment, but not limited to, one or more invention elements efficaciously utilize natural and or genetically engineered elements to encode components of the intracellular sorting machinery that mediate the selective trafficking of lipids and proteins in the secretory and endocytic pathways, to efficacious effect.

[0217] Referring again to FIGS. 1, 2, 3, 4, and 5, in another embodiment, but not limited to, one or more invention elements efficaciously utilize genetic agents and elements, including, but not limited to, proteins; peptides; DNA and DNA variants; RNA and RNA variants such as mRNA, iRNA and siRNA; RNA-induced silencing complex (RISC), other genetic-modifying agents and methods, and the like.

[0218] In another embodiment, but not limited to, one or more invention elements efficaciously utilize one or more oligonucleotides in antisense therapy. These antisense DNA drugs work by binding to messenger RNAs from disease genes, so that the genetic code in the RNA cannot be read, stopping the production of the disease-causing protein.

[0219] In another illustrative embodiment, one or more elements may comprise one or more RNAi (RNA interference) elements and or RNAi variants such as small interfering RNA molecules (siRNA), but not limited to, that may collaborate with proteins in the cell and also may form a nanoscale element called a RISC (RNA-Induced Silencing Complex). RNAi and or RISCs may be used to head off a genetic disease before the first symptom appears, based on an analysis of an individual's predisposition to certain diseases. This methodology is a way of silencing a specific gene, for example, genes that direct cancer cells to proliferate or that create overproduction of proteins that cause rheumatoid arthritis. Basically, RNAi works by scanning RNA templates that may cause a disease and cleaving that RNA template, and enzymes then destroying the template before it can complete its actions on the offending DNA. One of the key barriers to successful RNAi therapy is their finding their way to a specific site in the body and then the RNAi not degrading rapidly before it can do useful work. In one illustrative embodiment, RNAi, siRNA, RISC elements and or other suitable methods may be targeted by an invention element such that one or more such RNA elements seek out and destroy potentially harmful genetic elements and or other genetic processes.

[0220] As noted in the literature, Clathrin heavy chain is known to be a cytosolic protein that functions as a vesicle transporter. However, the Clathrin heavy chain exists not only in cytosol but also in cell nuclei. The p53 gene, in which mutations have been found in >50% of human cancers, encodes a protein that plays an important role in preventing tumorigenesis. Clathrin heavy chain expression enhances p53-dependent transactivation, whereas the reduction of Clathrin heavy chain expression by RNA interference (RNAi) attenuates its transcriptional activity. Moreover, Clathrin heavy chain binds to the p53-responsive promoter in vivo and stabilizes p53-p300 interaction to promote p53-mediated transcription. Thus, nuclear Clathrin heavy chain is required for the transactivation of p53 target genes and plays a distinct role from Clathrin-mediated endocytosis (Enari, et al 2006). In one embodiment, p53 and or one or more other types of genes, their diseases and disorders, and or RNAi related activities may be efficaciously controlled and regulated, mitigated, prevented, and or modified via one or more embodiments of the instant invention.

[0221] Referring again to FIGS. 1, 2, 3, 4, and 5, in another embodiment, but not limited to, one or more elements, acting alone or not, would achieve therapeutic effect by deliberately controlling and regulating, or modifying faulty exocytosis and or endocytosis processes that produce disorders and diseases. This is a health critical situation, as the role of dopamine receptors and transporters; the excitability of dopaminergic neurons; and the regulation of extracellular dopamine levels in the brain, especially in relation to the diseased state, has proven to be imperative for a further understanding of dopaminergic neurotransmission as a whole. For example, dopaminergic neurotransmission critically depends on exocytotic release and neuronal uptake of dopamine, as well as on diffusion away from the release site. Once target cells are reached, dopamine can bind to and activate dopamine receptors. The subsequent cellular response depends on the type of dopamine receptor that is activated and the signal transduction mechanisms that are coupled to these receptors. Disturbances in one or more of the above-mentioned aspects of dopaminergic transmission could lead to severe neurological and neuropsychiatric disorders such as Parkinson's disease, depression, addiction, schizophrenia, attention deficit hyperactivity disorder, restless legs syndrome, Tourette syndrome, and the like, and in or more invention embodiments, one or more such disorders may be efficaciously treated.

[0222] Referring again to FIGS. 1, 2, 3, 4, and 5, in another embodiment, but not limited to, one or more elements, during some operations may interact with, for example, an externally applied magnetic field, like during NMR. However, since invention protein elements are electrically neutral, only minimal (e.g., no) structural distortion of the elements occurs in the presence of the magnetic field. Therefore, using invention elements to capture other types of elements, which may be, for example, one or more NMR contrast agents for developmental imaging and diagnostic

studies, and which contrast agents may also be capable of crossing cellular membranes, protects and extends the utility of the invention.

[0223] Referring again to FIGS. 1, 2, 3, 4, and 5, in another embodiment, but not limited to, one or more elements may comprise, for example, one or more metal ions including, but not limited to, the gadolinium (III) chelate compounds of DTPA, DO3A, DOTA and other variations of these linear and macrocyclic ligands that act as targeted and or non-targeted contrast agents.

[0224] Direct Gd3+-OH2 chemical bonds, which exchange rapidly with other bulk H2O molecules, produce the mechanism whereby unpaired electrons on Gd3+ relax the proton nuclei of many nearby H2O molecules. Accordingly, the behavior of T1 contrast agents, such as those based on gadolinium requires good direct contact with tissue water molecules (spin-lattice relaxation mechanism) to be efficient. Thus, it is often preferable to bind them to the external surface of the carrier. (Hooker, et al. 2007) In one embodiment, one or more elements facilitate better contact to tissue water because one or more contrast agents of one or more types are not located in the interior part of a cage (in its cavity), but rather, located on much more exposed non-cage elements of one or more types. In one embodiment, one or more cage element 212 has one or more contrast agents of one or more types located on the outside part of cage element 212; or on both the inside and outside parts of element 212.

[0225] In another illustrative embodiment, one or more imaging or study elements comprise one or more treated manganese minerals, such as oxides, silicates, and carbonates for imaging and study enhancement.

[0226] Besides Gd3 complexes, there is another important class of contrast agents for MRI that is based on polysaccharide coated iron oxide particles. Their peculiarity stems from the fact that their blood half-life and distribution to different organs of the reticuloendothelial system (RES) depend upon the particle size (Aime, et al 1998). In one embodiment, one or more elements comprise one or more of a wide range of lanthano-invention labeled derivatives for custom-designed contrast agents.

[0227] In another embodiment, one or more elements comprise one or more therapeutic agents in addition to one or more imaging contrast and diagnostic agents.

[0228] In another illustrative embodiment, targeted and or non-targeted in vivo delivery of one or more elements are internally and or externally monitored, directed, activated, deactivated and or regulated, locally and or at a remote distance by, for example, but not limited to, NMR, ESR, ultrasound, radio transmissions, and or biochemical reactions.

[0229] Additionally, in other embodiments, NMR is combined with other techniques, such as ENDOR, which combines the best aspects of ESR and NMR, to yield high sensitivity and nuclear selectivity, respectively, for in vivo and in vitro studies.

[0230] In one embodiment, one or more different sized, paramagnetic coated, quantum dots, and or photonic dots are used as one or more contrast markers in magnetic resonance imaging (Mulder, et al., 2009). In other embodiments, one or more different sized quantum dots, and or photonic dots may be used in positron emission tomography (PET) for in-vivo molecular imaging, or as fluorescent tracers in optical microscopy.

[0231] In another configuration, one or more types of elements comprise one or more radiodiagnostic agents for nuclear medicine.

[0232] Referring again to FIG. 2, in further illustrative embodiments, free-floating cargo may be carried in cavity forming cargo elements 202a that comprise a fluid, gas, or vapor; which free-floating cargo, for example, may be one or more molecular ensembles for enhanced medical imaging, and which cargo may also be carrying one or more therapeutic agents.

[0233] Referring again to FIGS. 1, 2, 3, 4, and 5, in another embodiment, but not limited to, one or more invention elements comprise one or more types of elements in whole or in part, such as one or more drug and pharmacological elements; biological elements; biomedical or medical elements; and the like, including healthcare elements; bioengineered elements; cosmetic elements; and the like.

[0234] Referring again to FIGS. 1, 2, 3, 4, and 5, but not limited to, in one embodiment, one or more elements of one or more types comprise targeted and or non-targeted drug delivery elements, including their high precision dosing, or other forms of healthcare elements for diagnosing, remedying, inhibiting, mitigating, curing, and or preventing one or more types of diseases, infections, physical or mental trauma, or other forms of physical and mental afflictions.

[0235] Referring again to FIGS. 1, 2, 3, 4, and 5, but not limited to, in one embodiment, one or more elements comprise an in vitro and or in vivo model and or system for research study, including a model, method, and or system for the research and development of new drugs, therapies, prosthetics, and drug delivery systems, including an accelerated drug discovery process.

[0236] Referring again to FIGS. 1, 2, 3, 4, and 5, in another embodiment, but not limited to, one or more elements, acting alone or not, are utilized for studying, discovering, preventing, curing, mitigating, and or healing one or more types of animal, tree, plant, grain, grass, agricultural, vegetable, and or fungal diseases, disorders, infestations, and or blights.

[0237] Referring again to FIGS. 1, 2, 3, 4, and 5, in another embodiment, but not limited to, one or more elements are used for studying, discovering, designing, and or enabling of genetically engineered elements, for example, one or more types of genes, cells, and other biological elements and products in animals, trees, plants, grains, grasses, agriculture, vegetables and fungi.

[0238] In another illustrative embodiment, one or more elements comprise one or more methods for nourishing and or promoting healthy growth in one or more types of animals, trees, plants, grains, grasses, agriculture, vegetables and or fungi.

[0239] Referring again to FIGS. 2 and 4, in another embodiment, but not limited to, the heat shock cognate protein, hsc70, and its molecular co-chaperone auxilin, help to regulate the natural endocytosis aftermath of natural CCV uncoating and disassembly. Hsc70 also promotes uncoating and disassembly of Coatomer I and II vesicles. In cells over-expressing ATPase-deficient hsc70 mutants, uncoating of CCVs is inhibited in vivo. In one embodiment, bioengineered elements may be used to regulate under or over expression of hsc70 and or auxilin. In one example embodiment, using a monoclonal antibody or other agent type as cargo against hsc70 blocks the hsc70-mediated release of invention and or non-invention Clathrin from coated

vesicles. In another example embodiment, or more auxilin elements comprise invention elements.

[0240] In one illustrative embodiment, one or more elements are stable with respect to dissociation, including one or more associated non-invention elements.

[0241] In another illustrative embodiment, disassembly and dissolution of one or more elements are deliberately inhibited and control and regulated, including one or more associated non-invention elements.

[0242] In one illustrative embodiment, one or more elements remain stable for a time certain or estimated time before the onset of dissociation, including one or more associated non-invention elements.

[0243] In one illustrative embodiment, dissociation of one or more elements may occur in whole or in part, including one or more associated non-invention elements.

[0244] In one illustrative embodiment, one or more cargo elements may comprise one or more uncoating and dissociation agents and or use one or more methods for controlled and regulated release of agents or cargo from one or more elements, including one or more associated non-invention elements.

[0245] In another embodiment, disassembly and dissolution of one or more elements, including one or more associated non-invention elements are inhibited, controlled and regulated, and or promoted by using one or more specific agents, stimuli, and or other methods.

[0246] In one embodiment, but not limited to, one or more invention elements of one or more types are formed in vitro via the following protocols, which may be modified and or substituted by one or more other types of protocols in one or more invention embodiments: (Adapted from Campbell, C et al., Biochemistry 23, 4420-4426 (1984), Pearse & Robinson, EMBO J. 9:1951-7 (1984), and Zhu, et. al., Methods in Enzymology, 328, 2001, Kedersh N, et al., J. Cell Biology 103, 1986.)

[0247] (Adapted from Campbell, C et al., Biochemistry 23, 4420-4426 (1984), Pearse & Robinson, EMBO J. 9:1951-7 (1984), and Zhu, et. al., Methods in Enzymology, 328, 2001, Kedersh N, et al., J. Cell Biology 103, 1986.)

[0248] Part I. Method of Differential Centrifugation.

[0249] 1. Make up 1 L of a buffer (buffer A) that comprises: 50 mM Mes pH 6.5, 100 mM NaCl, 1 mM EGTA, 0.5 mM MgCl $_2$, 0.02% NaN $_3$, 1 mM DTT a day prior to experiment and storage at 4° C.

[0250] 2. Add 1:100 PMSF proteases inhibitor to buffer A (200 ul/20 ml).

[0251] 3. Collect and wash 14 rat brains (\sim 2.0 g) and livers (\sim 20.0 g). Wash and place the brains in ice-cold buffer A. Perfuse the livers with ice-cold PBS and collect them in ice-cold buffer A.

[0252] 4. Mince and homogenize the brains in a Potter-Elvehjem grinder with 2 volume of ice-cold buffer A per total brain wet weight (~90 ml). Do the same with the livers (~400 ml).

[0253] 5. Centrifuge the homogenate at 23,000 g (11,900 rpm) in a Sorvall GSA or at 13,000 rpm in a Sorvall SS34 rotor for 45 min at 4° C.

[0254] 6. Collect the supernatant and centrifuge at 43,000 g (18,000 rpm) in a Sorvall SS34 rotor or at 20,000 rpm in a ti 45 Beckman rotor for 1 h at 4° C.

[0255] 7. Resuspend the pellet in 10 ml of ice-cold buffer A, use a loose-fitting Teflon-glass Dounce homogenizer.

[0256] 8. Collect homogenate in a 50 ml conical tube. Wash pestle and glass homogenizer with 5 ml of buffer A, and add this to homogenate until total volume is 15 ml. Add 1:100 PMSF

[0257] 9. Dilute the homogenate 1:1 with 15 ml of 12.5% Ficoll/12.5% sucrose (both in ice-cold buffer A), and mix by inversion to ensure homogeneity.

[0258] 10. Centrifuge at 43,000 g (18,000 rpm) in a Sorvall SS34 rotor or at 20,000 rpm in a ti 45 Beckman rotor for 30 min at 4° C.

[0259] 11. Collect the supernatant in a graduate cylinder and dilute it 1:5 in ice-cold buffer A. Add 1:100 PMSF

[0260] 12. Centrifuge the supernatant at 100,000 g (33,000 rpm) in a Beckman 70.1Ti rotor or at 31,100 rpm in a ti 45 Beckman rotor for 1 h at 4 $^{\circ}$ C.

[0261] 13. Collect pellet and resuspend in 5-10 ml of ice-cold buffer A by using a loose-fitting Teflon-glass Dounce homogenizer. Add 1:100 PMSF

[0262] 14. Leave the homogenate on ice for about 30 min, and take an aliquot of 10 ul for EM, and dilute 1:10 for brain, 1:100 for liver.

[0263] Part II. Purification of CCVs Using Density Gradients (Zhu's CCVs and Clathrin Coat Preparation). Submit the Crude Clathrin-Coated Vesicles from Fresh Rat Brain to Discontinuous Sucrose Gradient for Remove Contaminating Vaults

[0264] 1. CCVs resuspended in (5-10 ml) buffer A

[0265] 2. Preparer a discontinuous sucrose gradient in SW28 tubes by carefully layering 5 ml of 40%, 5 ml of 30%, 6 ml of 20%, 8.5 ml of 10%, and 8.5 of 5% sucrose solutions in buffer A from bottom to top.

[0266] 3. CCVs (5-10 ml) is laid on top of the gradient and centrifuged at 100,000 g (25,000 rpm) in a SW28 rotor for 1 hr at 4° C.

[0267] 4. Collect twenty-six 1.5 ml factions from the top. [0268] 5. Small aliquots from every other faction are analyzed for CCVs using 10% SDS-PAGE. [Fractions comprising the CCVs (typically fractions 12-21 as numbered from the top of the gradient) are combined, diluted with 3 volumes of buffer A, and centrifuge at 112,000 g (31,100 rpm) in a ti 45 Beckman rotor for 1 h at 4° C. or at 33,000 rpm in a Beckman 70.1Ti rotor for 1 h at 4° C. Add 1:100 PMSF]

[0269] 6. Resuspend the pellet in ice-cold buffer A, do a protein assay to yield an approximate concentration. Usually add 1 to 2 ml of buffer A.

[0270] 7. Aliquot the homogenate in aliquots of 200 ul and store at -80° C. Take an aliquot of 10 ul each for EM and SDS-gel PAGE.

[0271] Part III. Isolation of Triskelia and APs from CCVs Using Keen's Method.

[0272] 1. Dialyze CCVs against 0.01M Tris buffer, Ph 8.5, 3 mM azide for 5 hours.

[0273] 2. Centrifuge at 240,000 g (51,200 rpm) for 20 min at 4° C. Because you are using low amount of sample; (IF we have less than 2 mL, Do not use the lid or close the centrifuge tubes of the 70.1 Ti rotor.) The soluble coat proteins comprising triskelial and APs are separated from the residual Clathrin-coat vesicle membranes.

[0274] 3. Collect the soluble fraction and do protein assay.[0275] 4. Take an aliquot of 10 ul for EM and 50 ul for SDS-gel PAGE.

[0276] Part IV. Separation by FPLC of AP-1 from AP-2 with Hydroxyapatite Column

[0277] Solutions:

Stocks:	1M NaH ₂ PO ₄ ; pH 7.1 5M NaCl 10% NaN ₃	(30 g/250 ml)
Low PO ₄ buffer (500 ml):	10 mM NaH ₂ PO ₄ ; pH 7.1 100 mM NaCl 0.02% NaN ₃ 0.1% beta-Mercaptoethanol	(5 ml of stock) (10 ml of stock) (1 ml of stock) (0.5 ml) (RT)
High PO ₄ buffer (200 ml):	500 mM NaH ₂ PO ₄ ; pH 7.1 100 mM NaCl 0.02% NaN ₃ 0.1% beta-Mercaptoethanol	(RT) (100 ml of stock) (4 ml of stock) (0.4 ml of stock) (0.2 ml) (RT)

[0278] Both buffers need to be filtered and degassed prior to use.

[**0279**] AP Buffer:

100 mM MES, pH 7.0	39 g/2 l
150 mM NaCl	17.5 g/2 l
1 mM EDTA	4 ml of 500 mM solution/2 1
0.02% NaN ₃	4 ml of 10% solution/2 l
0.5 mM DTT	-> add just before use
	(4° C.)

[0280] Hydroxyapatite Column:

[0281] 5 ml Econo-Pac CHT-II from BioRad; the column is stored at 4° C. in low PO₄ buffer

[0282] Procedure:

[0283] Connect the hydroxyapatite column to the FPLC system via the BioRad adaptors. Put a 0.2μ syringe filter at the inlet of the column.

[0284] Use the following FPLC settings:

[0285] Sensitivity: 1

[0286] Flow: 1 ml/min

[0287] Chart Recorder speed: 0.5 cm/min

[0288] Make sure the fraction collector is set at "ml" and a volume of "1"

[0289] Pump A is used for the low PO_4 buffer; Pump B for the high PO4 buffer. Wash the pumps with Valve 1 in position "3".

[0290] Once the FPLC system is set up, start washing the column with 20 ml of high PO_4 buffer (=20 min). Be sure to switch on UV-Lamp.

[0291] This is followed by equilibration of the column with low PO_4 buffer; i.e. until the baseline is stable. The backpressure of the system should be approx. 0.1 MPa and must not exceed 0.35 Mpa.

[0292] During the equilibration phase (Valve 1 in position "1"="Load"), the 50 ml superloop is loaded with the AP sample (Pump C; 5 ml/min).

[0293] With the column equilibrated and the superloop loaded, switch Valve 1 into position "2"="Inject". The APs are injected over the column at a flow rate of 1 ml/min.

[0294] After the injection is completed, continue running low PO_4 buffer over the column until the baseline is stable. Don't forget to prepare 1.5 ml tubes for the fraction collector

 $\boldsymbol{[0295]}$ AP-1 and AP-2 are then eluted from the column using Method 6:

0.0	CONC % B	0.0
0.0	VALVE.POS	1.1
0.0	CM/ML	0.50
0.0	PORT.SET	6.1
40.0	CONC % B	0.0
40.0	ML/MIN	1.00
50.0	CONC % B	100

[0296] The elution profiles for AP-1 and AP-2 tend to vary considerably from one purification to another; AP-1 is eluted first

[0297] AP-1 tends to be eluted from the column in three to four 1 ml fractions, usually starting at around #13. AP-2 is usually eluted in up to 15 fractions, starting at around #25. The fractions comprising the APs need to be verified by SDS-PAGE (two gels of 10% or 12%)

[0298] Wash column with low PO_4 buffer; store at 4° C. [0299] Pooled AP-1 fractions and pooled AP-2 fractions are dialyzed against 1 liter of AP buffer overnight, and for a few more hours after exchanging the buffer (4° C.). The samples are then stored at 4° C.

[0300] Typically, the concentration for Clathrin (peak fractions) is approx. 0.5 mg/ml, for AP-1 and AP-2 between 0.3-0.5 mg/ml.

[0301] According to one illustrative embodiment, but is not limited to, recombinant Clathrin formation may be achieved in the following exemplar manner. Stoichiometric quantities of adaptor elements 208a comprising AP-1 and AP-2 are required for Clathrin self-assembly at physiological pH. However, in vitro Clathrin self-assembly occurs spontaneously below about pH 6.5. Recombinant terminal and distal domain fragments are produced and combined with recombinant-produced hub fragments in assembly buffer as described below in order to induce formation of one or more Clathrin elements, such as those comprising elements 206a, for use in the invention.

[0302] In one illustrative technique, bovine Clathrin heavy chain cDNA encoding heavy chain amino acids 1-1074 (SEQ ID NO: 1) is cloned into the pET23d vector (Novagen) between the NcoI (234) and XhoI (158) sites. Expression of the cloned sequence results in a terminal and distal domain fragments having a C-terminal polyhistidine tag. Hub fragments corresponding to amino acids 1074-1675 (SEQ ID NO: 1) are cloned into vector pET15b (Novagen) between the BamHI (319) and XhoI (324) sites. Expression of the hub fragments produces the proximal leg domain and central trimerization domain of the Clathrin hub with an N-terminal polyhistidine tag. Vectors comprising the heavy chain and hub domains are expressed in E. coli by induction with 0.8 mM isopropyl-B-D-thiogalactopyranoside for 3 hours at 30 degrees Celsius. Expressed proteins are isolated, recombinant, and or synthetic from bacterial lysate in binding buffer (50 mM Tris-HCl (pH7.9), 0.5M NaCl, 5 mM imidazole) in a nickel affinity resin using the polyhistidine tag. Proteins are eluted with 206a mM EDTA and dialyzed against 50 mM Tris-HCl (pH7.9). Hub fragments are further isolated, recombinant, and or synthetic using size exclusion chromatography on a Superose 6 column (Pharmacia).

[0303] In another exemplar technique, Clathrin assembly reactions are performed using expressed heavy chain and hub fragments by overnight dialysis at 4 degrees Celsius in

assembly buffer (100 mM 2-(N-morpholino) ethanesulfonic acid, pH 6.7, 0.5 mM MgCl2, 1 mM EGTA, 1 MM Tris (2-carboxyethyl)-phosphine hydrochloride, 3 mM CaCl2. Assembly reactions are centrifuged for 5 minutes at 12,000 rpm. The supernatant is then centrifuged for 45 minutes at 45,000 rpm (100,000×g). The pellets are resuspended in assembly buffer, and protein composition is determined on SDS-PAGE. The efficiency of element **206***a* formation can be determined by electron microscopy by diluting assembly reactions 1:5 in 10 mM Tris pH7.9, and placing aliquots on a glow-discharged carbon-coated grid, using 1% uranyl acetate as the stain.

[0304] According to another illustrative embodiment, but is not limited to, recombinant Clathrin formation may be achieved in the following exemplar manner, as described by Rapoport, et al. (MBC 2008): A cDNA encoding rat Clathrin heavy chain (Kirchhausen et al., 1987a) is used as a template to generate full-length (1675 HC), nested C-terminal truncations (1661 HC, 1643 HC, 1637 HC, 1630 HC, and 1596 HC), internal deletions (1675 PIVYGQ HC, 1643 PIVYGQ HC, and 1675 QLMLTA HC), and mutations (1643LML-AAA HC) of the heavy chain; each is then subcloned into the insect cell expression vector pFastBac1 (Invitrogen, Carlsbad, Calif.). A cDNA encoding rat liver Clathrin light chain LCa (Kirchhausen et al., 1987b) is used as the template to subclone the region encoding the full light chain (residues 1-256) into the insect cell expression vector pFastBacHTb. The final construct (rLCali) comprises at its N terminus a 6x-His-tag followed by a linker of 20 residues. Baculoviruses suitable for infection and expression are generated with the Bac-to-Bac system (BD Biosciences, San Jose, Calif.). Virus stocks are obtained after four rounds of amplification, and they are kept in the dark at 4° C. The open reading frame of rat brain Clathrin light chain LCa1 is also used as a template to subclone it into the bacterial expression vector pET28b (Novagen, Madison, Wis.) between the NcoI and EcoR1 restriction sites so as to generate a native, nontagged light chain. All constructs are verified by DNA sequencing. Clathrin heavy chains together with light chain are expressed in Hi5 insect cells (1L, 1-1.5 206a cells/ml) grown for 2-3 d in spinner flasks at 27° C. in Excell 420 medium after coinfection with the appropriate viruses. Alternatively, Clathrin heavy chain only is expressed in a similar way. The cells are centrifuged at 1000 rpm for 10 min at room temperature by using an H6000A rotor (Sorvall, Newton, Conn.), and the pellets are resuspended in 20 ml lysis buffer (50 mM Tris, pH 8.0, 300 mM NaCl, 1 mM EDTA, 3 mM mercaptoethanol, and half of a tablet of Complete Protease Inhibitor Cocktail [Roche Applied Science, Indianapolis, Ind.]). The resuspended pellets are sonicated for 1 min on ice (Flat tip at 20% power, Ultrasonic processor XL; Heat Systems, Farmingdale, N.Y.), cell debris is removed by centrifugation at 90,000 rpm for 20 min at 4° C. by using a TLA 100.4 rotor (Beckman Coulter, Fullerton, Calif.), and the supernatant (20 ml) is dialyzed at 4° C. for 12 h against 2×2 l of cage buffer (20 mM [2-(N-morpholino) ethanesulfonic acid] MES, pH 6.2, 2 mM CaCl2, 0.02% NaN3, and 0.5 m Mdithiothreitol [DTT]). The sample is then centrifuged at 4° C., first at low speed (1000 rpm for 10 min) to remove large aggregates and then at high speed (54,000 rpm for 1 h) by using a Ti rotor (Beckman Coulter). The pellet, primarily comprising Clathrin (presumably assembled as cages) is resuspended in 6 ml of 100 mM MES, pH 6.5, 3 mM EDTA, 0.5 mM MgC12, 0.02% NaN3, 0.5 mM DTT,

and 0.5 mM phenylmethylsulfonyl fluoride) followed by addition of 3 ml of 2.4MTris, pH 7.4, 1 mM DTT, and incubation for 20 min at room temperature, a condition used to dissociate native Clathrin assemblies. The sample is centrifuged at 90,000 rpm for 20 min at 4° C. by using a TLA 100.4 rotor, and most of the Clathrin is recovered in the supernatant. The resulting sample is subjected to gel filtration chromatography (90 cmר=3 cm column comprising Sephacryl-S 500 [GE Healthcare, Little Chalfont, Buckinghamshire, United Kingdom] in 0.5 M Tris, pH 7.4, 0.04% NaN3, and 0.5 mM DTT) at room temperature and with a flow of 2 ml/min. Fractions of 5.5 ml comprising the Clathrin peak (100 ml) are pooled and then subjected to adsorption chromatography (5 ml, hydroxyapatite, Econo-Pac CHT-II; Bio-Rad, Hercules, Calif.); the column is pre-equilibrated with low phosphate buffer (10 mM NaH2PO4, pH 7.1, 100 mM NaCl, 0.02% NaN3, and 0.5 mM DTT) and eluted with a linear gradient from low to high phosphate concentration (500 mM NaH2PO4, pH 7.1, 100 mM NaCl, 0.02% NaN3, and 0.5 mM DTT) at room temperature with a flow of 1 ml/min. Fractions (1 ml) are collected into microcentrifuge tubes comprising 21 of 0.5 M EDTA. Typical Clathrin yields are in the range of 3-40 mg per 1 1 of cell culture. Western blot analysis is used to confirm the expression of Clathrin heavy and light chains. The rat Clathrin light chain rLCa1b is expressed in Escherichia coli strain BL21(DF3). The bacteria are grown in Luria-Bertani (LB) medium comprising 30 mg/l kanamycin at 37° C. with shaking (250 rpm) to an optical density of 0.5. Expression is induced by addition of isopropyl-d-thiogalactoside (IPTG) (final concentration, 0.6 mM). After 3 h, the cell are harvested by centrifugation at 5000 rpm for 10 min at 4° C. by using an H6000A rotor (Sorvall) and resuspended in ice-cold lysis buffer (20 mM Bis-Tris adjusted to pH 6.0 at room temperature, 0.5 mM dithiothreitol, 1 mM EDTA, and Complete Protease Inhibitor Cocktail) by using 20 ml of lysis buffer per 3.5 g of wet cell weight. The suspension is placed into a glass vessel, and the vessel is immersed in boiling water for 4 min and then chilled on ice. The boiled suspension is centrifuged at 54,000 rpm for 30 min at 4° C. by using a 60Ti rotor (Beckman Coulter) to remove the precipitated material. rLCa1b is purified from the filtered supernatant (0.2-msyringe filter) by anion exchange chromatography at 4° C. on a HiTrap MonoQ column equilibrated with buffer A (20 mM Bis-Tris, adjusted to pH 6.0 at room temperature, and 0.5 mM dithiothreitol) and eluted using a linear gradient from 0 to 32% buffer B (20 mMBis-Tris, adjusted to pH 6.0 at room temperature, 0.5 mM dithiothreitol, and 1 M NaCl). For the in vitro reconstitution of Clathrin, recombinant heavy chain (expressed in insect cells without light chain) is mixed with excess rLCa1b (expressed in bacteria) by using a weight ratio of 3:1 (equivalent to a molar ratio HC:LC of 1:2.4) just before cage or coat assembly for 40 min at room temperature.

[0305] Part V. Clathrin Coat Formation

[0306] Reagents

[0307] 1. Coat Formation Buffer

80 mM Mes hydrate pH 6.5 31.23 g/2 L 20 mM NaCl 2 mM EDTA 8 mL of 500 mM stock solution/2 L 0.4 mM DTT 1.6 mL of 500 mM stock solution/2 L

[0308] 2. Clathrin [0309] 3. AP-2 [0310]Procedure [0311] (1) Place a solution of clathrin and AP-2 into a dialysis chamber [$\dot{0}312$] clathrin: AP-2 =3:1 to 4:1 (w/w) [0313] (2) Dialyze over night against coat formation buffer; replace buffer and dialyze for an additional 3-4 h. [0314] (3) Transfer to a centrifuge tube, centrifuge to remove larger aggregates [0315] rotor: TLA-100.4, 12000 rpm, 4° C., 10 min [0316] (4) Transfer supernatant to fresh centrifuge tube, centrifuge to collect coats [0317] rotor: TLA-100.4, 65000 rpm, 4° C., 12 min [0318] (5) Immediately withdraw supernatant with a 1 mL pipette. [0319] (6) Wash carefully with buffer around the pellet. [0320] (7) Resuspend the pellet by adding buffer, allowing to stand at room temperature for 10-15 min, then slowly wash buffer over the pellet to resuspend using a micropipettor (avoid foaming) [0321] volume: 120-150 μ L for a pellet of ~3 mm diameter [0322] Part VI. Clathrin Cage Formation

[0323] Reagents

[0324] 1. Cage Formation Buffer:

[0325] 20 mM Mes, pH 6.2 (3.9 g/l) (7.8 g/2 l)

[0326] 2 mM CaCl2 (2 ml of 1 M/l) (4 ml of 1 M/2 l)

[0327] 0.02% NaN3 (2 ml of 10%/l) (4 ml of 10%/2 l)

[0328] 0.5 mM DTT (1 ml of 500 mM/l) (2 ml of 500 mM/21

[0329] 2. Clathrin

[0330] Procedure

[0331] (1) Place a solution of Clathrin (0.5-1 mg/mL) into a dialysis chamber

[0332] (2) Dialyze over night against cage formation buffer; replace buffer and dialyze for an additional 3-4 h.

[0333] (3) Transfer to a centrifuge tube, centrifuge to remove larger aggregates

[0334] rotor: TLA-100.4, 12000 rpm, 4° C., 10 min [0335] (4) Transfer supernatant to fresh centrifuge tube, centrifuge to collect coats

[0336] rotor: TLA-100.4, 65000 rpm, 4° C., 12 min [0337] (5) Immediately withdraw supernatant with a 1 mL pipette.

[0338] (6) Wash carefully with buffer around the pellet.

[0339] (7) Resuspend the pellet by adding buffer, allowing to stand at room temperature for 10-15 min, then slowly wash buffer over the pellet to resuspend using a micropipettor (avoid foaming)

[0340] Production of Recombinant Auxilin

[0341] A protein chimera of glutathione transferase (GST) with bovine auxilin (spanning residues 547-910) is generated by fusion in the vector pGEX4T-1 and then used for expression in E. coli BL21 (Fotin et al., 2004a). The bacteria are grown in LB medium supplemented with ampicillin to an OD600 0.5-0.6 at 37° C. Protein expression is induced by addition of 1 mM IPTG (final concentration) and the cells grown for another 4 h at 25° C. The cells (from 1 1 of culture) are centrifuged at 5000 rpm for 15 min at 4° C., and the pellet is kept frozen overnight. The pellet is resuspended in 25 ml of pGEX lysis buffer (20 mM HEPES, pH 7.6, 100 mM KCl, 0.2 mM EDTA, 20% glycerol, 1 mM DTT, and half a tablet of Complete Protease Inhibitor Cocktail) and sonicated on ice using three consecutive sonication cycles of

60, 30, and 30 s (standard microtip, 20% power). The sample is centrifuged at 45,000 rpm for 1 h at 4° C. by using a 60Ti rotor, and the supernatant mixed with 0.5 ml of a 50% (vol/vol) slurry of glutathione-Sepharose 4 beads (GE Healthcare). After 2 h of end-over-end rotation at 4° C., the beads are poured into a propylene Econo-Column (Bio-Rad), washed with 15 ml of pGEX lysis buffer, and then washed with 15 ml of 25 mM HEPES, pH 7.0, 100 mM NaCl, and 0.1 mM EGTA. Elution of GST-auxilin (in 2 ml) is achieved by supplementing the solution with 50 mM glutathione, adjusted to pH 8. These steps are carried out at 4° C. Release of the GST portion is achieved by incubation of 1 mg of GST-auxilin with 1 U of thrombin at room temperature for 6 h. Proteolysis is ended by addition of 1 mg of Pefabloc SC (Roche Applied Science). The 40-Da auxilin fragment is further purified using a Mono S column (Pharmacia, Peapack, N.J.). The sample is first dialyzed overnight against MES buffer A (50 mM MES, pH 6.7, 1 mM EDTA, and 3 mM-mercaptoethanol), and then it is loaded onto the column (pre-equilibrated with MES buffer A) and eluted with a linear gradient of buffer A and with MES buffer B (50 mM MES, pH 6.7, 500 mM NaCl, 1 mM EDTA, and 3 mM-mercaptoethanol) at a flow of 1 ml/min. The auxilin sample is stored at 80° C. with 20% glycerol (final concentration).

[0342] Production of Recombinant Hsc70

[0343] N-terminal 6x-His-tagged bovine Hsc70 (full length) cloned into the pET2lavector is expressed in E. coli BL21. The bacteria are grown at 37° C. in LB supplemented with 0.1 mg/ml ampicillin to an OD600 of 0.5, transferred to 28° C., and induced with 0.1 mM IPTG for 5 h. The cells are centrifuged at 5000 rpm for 15 min at 4° C., and the pellets from 11 culture resuspended in 25 ml 50 mM Tris, pH 8.0, 300 mM NaCl, 1 mM ATP, 2 mM MgCl2, 10 mM-mercaptoethanol, and half a tablet of Complete Protease Inhibitor Cocktail without EDTA. The supernatant obtained after sonication and centrifugation (as with auxilin) is mixed with 1 ml of 50% (vol/vol) slurry of nickelnitrilotriacetic acid-agarose beads (QIAGEN, Valencia, Calif.) for 4 h by endover-end rotation at 4° C. The beads are placed into an Econo Pac column and then washed with 30 ml of 50 mM Tris, pH 8.0, 300 mM NaCl, 10 mM-mercaptoethanol, 10 mM imidazole, 1 mM ATP, and 1 mM MgCl2). Hsc70 is then eluted at 4° C. with 5-6 ml of the same solution supplemented with 200 mM imidazole. Fractions of 1 ml are collected into microcentrifuge tubes comprising 40 l of 0.1 M EGTA. The samples comprising 20% glycerol (final concentration) are stored at 80° C.

[0344] According to another illustrative embodiment, Clathrin and or Coatomer I/II proteins are extracted and prepared from Clathrin and or Coatomer I/II coated vesicles obtained from non-rat, non-bovine organic tissue, including from human tissue, in whole or in part. In another embodiment, Clathrin and or Coatomer I/II coated proteins are extracted and prepared from Clathrin and or Coatomer I/II coated vesicles obtained by donor/recipient tissue matching using established techniques. In another embodiment, Clathrin and or Coatomer I/II proteins are prepared, in whole or in part, by using stem cells, cloning and or other genetic manipulation techniques known in the prior art to produce genetically matched tissue for a donor recipient.

[0345] According to one illustrative embodiment, the coat protein I (COPI) assembly process is carried out by preparing Coatomer subunits from cytosolic preparations, includ-

ing methods, but are not limited to, as essentially described in Spang, et al., Proc. Natl. Acad. Sci. USA. 1998 September 15; 95 (19): 11199-11204. Coatomer, a nanoscale element comprised of seven distinct subunits (alpha, beta, beta', gamma, delta, epsilon and zeta subunits, respectively) and ADP-ribosylation factor (ARF, an N-myristylated small GTP-binding protein) are the only cytoplasmic proteins needed.

[0346] In another illustrative embodiment, the coat protein I (COPI) assembly process is carried out by preparing Coatomer subunits from cytosolic preparations, including methods, but are not limited to, as essentially described in Sheff, et al, The Journal Of Biological Chemistry, Vol. 271, No. 12, Issue Of March 22, Pp. 7230-7236, 1996 "Purification of Rat Liver Coatomer (COPI")—Purification of rat liver Coatomer is accomplished through a substantial modification of the method of Waters and Rothman (13). Unless otherwise noted, all operations are performed at 4° C. Approximately 250 g of fresh liver from 10-15 adult Sprague-Dawley rats (Harlan Sprague-Dawley) are homogenized in 2 volumes of buffer (25 mM Tris, pH 7.5, 320 mM sucrose, 500 mM KCl, 2 mM EDTA, 1 mM dithiothreitol) comprising protease inhibitors (2 mg/ml pepstatin A, antipain, and leupeptin; 1 mM phenylmethylsulfonyl fluoride) using a polytron homogenizer with 1.5-cm cutter assembly at maximum speed for three 1-min bursts on ice with 1-min rests. The lysate is cleared by sequential centrifugation at 9000 3 g for 15 min followed by centrifugation of the supernatant at 100,000 3 g for 1 h. This material (S100) is stored at 270° C. for up to 4 months. For a typical purification, 150 ml of S100 is diluted 6-fold with cytosol buffer (25 mM Tris, pH 7.5, 1 mM dithiothreitol, 1 mM EDTA plus protease inhibitors as above). Protein concentration is 5 mg/ml. Ammonium sulfate is added to 25% of saturation and stirred for 15 min on ice, and then precipitate is removed by centrifugation, and the supernatant is brought to ammonium sulfate at 45% of saturation with stirring on ice. The precipitate is collected by centrifugation and redissolved in 150 ml of cytosol buffer. An additional 120 ml of cytosol buffer is added and then 30 ml of 60% (w/v) polyethylene glycol 3350 in distilled H2O with gentle stirring. The mixture is incubated at 4° C. for 30 min, and the precipitate is collected by centrifugation at 10,000 3 g for 15 min. The precipitate is resuspended in 20 ml of G buffer (10 mM Tris, pH 7.5, 0.2 mM ATP, 0.2 mM CaCl2), the insoluble material is removed by centrifugation, and the supernatant is passed over a 20-ml column comprising 250 mg of DNase-I (Sigma) coupled to agarose (Affi-Gel-10, Bio-Rad, prepared according to the manufacturer's directions) to remove contaminating actin and actin binding proteins. Eluent is desalted into cytosol buffer using 10DG desalting columns (Bio-Rad) and applied to a 50-ml DEAE cellulose column (DE52, Whatman) equilibrated in cytosol buffer. COPI is eluted with a 100-400 mM KCl gradient over 200 ml, with the elution of COPI followed by spot blot on nitrocellulose using EAGE antibody. In a final step, peak COPI fractions are pooled, diluted 1:1 with cytosol buffer, and applied to a 1-ml Mono-Q column (Pharmacia) equilibrated in cytosol buffer and mounted on a fast protein liquid chromatography apparatus (Pharmacia). The column is swished with 300 mM NaCl and then eluted with a 350-400 mM NaCl gradient over 20 ml. COPI, as assayed by the presence of b-COP on a spot blot using EAGE antibody, eluted as a single peak. The presence and purity of COPI is confirmed by SDS-

PAGE. An alternative final step is employed in preparing samples for two-dimensional dimensional gels. Here, DEAE eluent is concentrated in a Centricon-30 microconcentration (Amicon) to 400 ml and applied to a 24-ml Superose-6 (Pharmacia) column equilibrated in cytosol buffer with 50 mM KCl. As with Mono-Q, COPI eluted in a single peak. This final step produces a somewhat lower yield and comprises some contaminants between 30 and 100 KD by SDS-PAGE. For copurification of labeled CHO cytosol and rat liver COPI, all quantities are divided by 3, 1 ml of labeled cytosol is added to 50 ml of rat liver S100, and the Mono-Q column is used as the final step.

[0347] The increasing interest in the targeting of foreign moieties at sites in the body where their activity is required is addressed by the invention in one more embodiments. It is important that agents, like drugs, particularly those having undesirable side effects, are delivered to the site where they are supposed to act. Many molecular species require that they be delivered in a site specific manner, often to particular cells, for example, polynucleotides (anti-sense or ribozymes), metabolic co-factors or imaging agents. One such system has been described by Wu et al., J. Biol. Chem., 263, 14621-14624 and WO-A-9206180, in which a nucleic acid useful for gene therapy is conjugated with polylysine linked to galactose which is recognized by the asialoglycoprotein cargo attachment elements on the surface of cells to be targeted. However, there are many occasions, such as in the delivery of a cytotoxic drug, when it would not be satisfactory to use a delivery system in which the targeting and or masking moiety and or vector to be delivered is so exposed. This need is addressed by various delivery system embodiments of the invention that possess the flexibility to target a wide range of biologically active foreign moieties.

[0348] In one embodiment, the invention includes one or more elements having one or more suitable sites for subsequent attachment of a targeting and or masking moiety and or vector, and one or more elements having one or more surfaces and or protein coats to which one or more targeting and or masking moieties and or vectors have already been attached.

[0349] In one embodiment, one or more masking moieties are attached to the surface of one or more invention elements. These masking moieties prevent the recognition by a specific cell surface and instead allows for intravenous administration applications. For example, the surface masking characteristics may be provided by poly (ethylene glycol) (PEG) by using various PEG-PLA and PLGA mixtures. PEG conjugation masks the protein's surface, reduces its renal filtration, prevents the approach of antibodies or antigen processing cells and reduces its degradation by proteolytic enzymes. In one embodiment, PEGylated elements significantly improve element stability and prevent leakage of agents from elements. Studies have shown that proteinbased nanoparticles and liposomes without PEGs have a short circulation time due to rapid uptake by macrophages of the reticulo-endothelial system (RES), primarily in the liver and spleen. Finally, PEG conveys to molecules its physicochemical properties and therefore modifies biodistribution and solubility of peptide and non-peptide nanoparticles. Thus, recent studies have used mostly nanoparticles with PEGs. The PEG coating is highly hydrated and this layer protects against interactions with molecular and biological components in the blood stream, as well as nonspecific binding to tissue. In one embodiment, one or more elements, in one or more configurations, are internally and or externally attached, coated, and treated, in whole or in part by using steric stabilizers including, but not limited to, steric stabilizers selected among dipalmitoyl phosphatidyl ethanolamine-PEG, PEG-stearate, the esters of the fatty acids from the myristic acid to the docosanoic acid with methyl ether PEG, the diacylphosphatidyl ethanolamines esterified with methyl ether PEG and the polylactates and the polygly-colactates esterified with methyl ether PEG. In one embodiment, one or more elements are not required to be PEGy-lated to efficaciously operate.

[0350] In another embodiment, one or more elements, and in one or more configurations are internally and or externally coated or treated in whole or in part with surfactants, including, but not limited to, surfactant agents selected among soy-bean phosphatidylcholine, dioleyl phosphatidylcholine, dipalmitoyl phosphatidylcholine, hydrogenated soy-bean phosphatidylcholine, phosphatidylethanolamine and phosphatidylserine), and or with cosurfactants, including, but not limited to cosurfactant agents selected among ethanol, propanol, isopropanol, butanol, sodium taurocholate, sodium glycocholate, propylene glycol, butyric acid and benzoic acid.

[0351] In one or more embodiments, ligands can be of one or more efficacious types, such as drugs, and may be bioengineered, and or comprise isolated, recombinant, synthetic, and or cloned elements.

[0352] In one embodiment, one or more types of ligands may be functionalized and or attached in one or more ways to one or more elements.

[0353] In one embodiment, ligands are natural ligands of one or more types. In another embodiment, one or more types of natural ligands are modified and or functionalized. In another embodiment, invention element ligands and natural element ligands are combined to comprise one or more types of hybrid ligand elements.

[0354] In another embodiment, the course of a natural ligand and or invention ligand element during cellular signaling, trafficking, downregulation, upregulation, endocytosis, exocytosis, and other cellular entry or exit, cellular inter- and or intra-actions, and the like, may be efficaciously controlled, regulated, and or modified by one or more elements to yield one or more diagnosis, cure, mitigation, treatment, prevention of disease, or other types of efficacious effects, and the like.

[0355] Examples of some natural ligands, but not limited to, that may be subject to efficacious control, modification, and or regulation in one or more invention embodiments are listed below:

[0356] Toxins and Lectins, e.g.,

[0357] Diptheria Toxin

[0358] Pseudomonas toxin

[0359] Cholera toxin

[0360] Ricin

[0361] Concanavalin A

[0362] Viruses, e.g.,

[0363] Rous sarcoma virus

[0364] Semliki forest virus

[0365] Vesicular stomatitis virus

[0366] Adenovirus

[0367] Influenza

[0368] West Nile

[0369] Serum Transport Proteins and Antibodies, e.g.,

[0370] Transferrin

[0371] Low density lipoprotein

[0372] Transcobalamin

[0373] Yolk proteins

[0374] IgE

[0375] Polymeric Ig

[0376] Maternal Ig

[0377] IgG, via Fc receptors

[0378] Hormones and Growth Factors, e.g.,

[0379] Insulin

[0380] Epidermal Growth Factor

[0381] Growth Hormone

[0382] Thyroid stimulating hormone

[0383] Nerve Growth Factor

[0384] Calcitonin

[0385] Glucagon

[0386] Prolactin

[0387] Luteinizing Hormone

[0388] Thyroid hormone

[0389] Platelet Derived Growth Factor

[0390] Interferon

[0391] Catecholamines

[0392] LDL

[0393] Neurotransmitters

[0394] Substance P

[0395] A Neurotransmitter Known to Stimulate Pain Receptors

[0396] In one or more embodiments, one or more elements are conjugated (bonded) with one or more other elements (e.g., ligands), agents, materials, and or substances of one or more types, including those developed by 3rd parties, which may be used singly or mixed together in one or more configurations for medical and biological research, diagnosis, therapy, or prosthetic purposes. One or more biomedical elements such as ligands and other types of biomedical functionalization elements may be directly and or indirectly attached, bonded, fastened, cross-linked, and or affixed to and or incorporated into one or more invention elements, as well as one or more non-invention and or natural elements. In one embodiment, attachment is achieved via molecular tethers. In another embodiment, no molecular tether is involved. In one configuration, a free radical molecule may be attached directly to one or more invention elements. In another embodiment, one or more elements may be bonded, fastened, and or affixed to one or more elements by being included in a modified protein sequence of one or more elements or bonded elements; by using a spacer; by covalent bonding; by site directed mutagenesis; by genetically engineered mutation and or modification; by peptides; by proteins; by DNA; by antibodies; by monoclonal antibodies; by recombinant elements; and via other bioengineering techniques and methods known in the art.

[0397] According to one embodiment, the protein amino acid sequence of one or more elements are modified to provide a site suitable for attachment thereto of an in vivo or in vitro targeting and or masking moiety. In one illustrative embodiment, one or more target-specific ligands and or targeting moieties are directly attached to one or more elements via one or more amino acid groups, and or attached via one or more short molecular tethers.

[0398] In another embodiment, one or more functionalization elements, of one or more types, comprise highly specific targeting agents, such as, but not limited to, antibodies, peptides or small molecules, large molecules, and other functional ligands, such as fluorophores and permeation enhancers, and the so functionalized nanoparticles may target receptors, transporter, enzymes and or intracellular processes in vivo with high affinity and specificity.

[0399] In one illustrative embodiment, one or more elements such as diagnostic, therapeutic, prosthetic, and or assay agents, but not limited to, are delivered to a target in vivo or in vitro using a variety of guidance techniques, including for example, optical (photonic), acoustic, electric, biological, chemical, mechanical reactions and forces, but not limited to, and one or more elements may be delivered singly and or in one or more configurations to one or more targets.

[0400] In another illustrative embodiment, one or more elements comprise one or more diagnostic agents like imaging contrast or radioactive agents to perform site designation, site specificity, and site retention for targeted in vivo delivery of therapeutics; the latter may also comprise part of the same diagnostic payload.

[0401] In one illustrative embodiment, the invention enables targeted agent delivery systems that retain their structural integrity and that may also loiter for a calculated period of time at the targeted area of concern after delivery of agent payload.

[0402] In one illustrative embodiment, one or more elements comprise molecules arranged in specific patterns. The pattern of elements precisely mirrors or mimics a spatial or physical pattern a target cell in a human or animal body expects to see and will recognize, and one or more elements are accepted by the target cell, which can be a cancer cell or HIV infected cell, for example.

[0403] In one embodiment, gold metal nanoparticle probes with sensor ligands and using electrical charges are bonded to one or more elements, and or attached to ligands, targeting moieties, and or vectors. The gold particles carry short strands of artificial DNA (oligonucleotides) tailored to match known segments of biological DNA that are implicated in, or linked to, disease.

[0404] Target-specific ligand binding and any subsequent changes within or to one or more elements may be a result of either covalent or non-covalent interactions—the latter including hydrogen bonding, ionic interactions, Van der Waals interactions, and hydrophobic bonds—depending on the application, system design, receptor design, cargo type and or the interaction/application environment.

[0405] In another illustrative embodiment, one or more elements, ligands, targeting moieties, vectors, and the like utilize the method of chirality.

[0406] In another illustrative embodiment, reactions and forces arise from one or more ligands and or targeting moieties binding to targets, including covalent and noncovalent interactions, which ligands are tethered and or directly attached to one or more invention elements. Ligand binding to one or more specific targets may produce one or more conformational changes sufficient to deform and or rupture one or one or more elements in whole or in part, thereby causing one or more elements to be released. The targeting moieties can be selected by one of ordinary skill in the art keeping in mind the specific cell surface to be targeted. For example, if one wishes to target the asialoglycoprotein receptor on the hepatocytes in the liver, an appropriate targeting moiety would be clustered trigalactosamine. Once a specific targeting moiety has been selected for a particular cell to target, the different targeting moieties can be attached either by covalent linkage directly onto the surface of one or more invention elements, or by indirect linkage via, for example, a biotin-avidin bridge. In another embodiment, depolymerization (e.g., by cytosolic Hsc 70) of the Clathrin and or Coatomer element exposes one or more transmembrane proteins (V-SNARE) that direct one or more elements to their destinations by binding to a specific T-SNARE protein on the target organelle. The fusion protein SNAP25 causes the one or more elements to fuse with the target membrane

[0407] In one embodiment, avidin is attached covalently to the surface of one or more elements and a biotinylated ligand attaches non-covalently to the avidin. In another embodiment, biotin is covalently attached to the surface of one or more invention elements, and then avidin is used as a bridge between the biotinylated polymer and the biotinylated ligand. Targeting agents may also include one or more biocompounds, or portions thereof, that interact specifically with individual cells, small groups of cells, or large categories of cells. Examples of useful targeting agents include, but are not limited to, low-density lipoproteins (LDS's), transferrin, asiaglycoproteins, gp120 envelope protein of the human immunodeficiency virus (HIV), and diphtheria toxin, antibodies, and carbohydrates. A variety of agents that direct compositions to particular cells are known in the prior art (see, for example, Cotten et al., Methods Enzym, 1993, 217, 618).

[0408] In another illustrative embodiment, one or more classical structural activity relationships (SARs) based drug discovery approaches are combined with one or more other techniques to form a specific case of targeted drug delivery, for example, but not limited to, one or more structural metabolism relationships (SMRs) that in combination with SARs are sometimes termed as retrometabolic drug design approaches. These active drugs are designed to undergo singular metabolic deactivation after they achieve their therapeutic roles, and may produce specific action at the site of application without affecting the rest of the body.

[0409] In another illustrative embodiment, one or more elements comprise one or more agent functionalities and or methods that produce targeting by changing molecular properties of an overall target molecule, as a result of enzymatic conversion, but also, for example, may involve one or more pharmacophores. These elements, sometimes referred to as the targetor (Tor) moiety, are converted by site-specific enzymes to active functions. In addition to the Tor moiety, one or more other functions may be introduced into elements for in vivo use, which can be named as "protector functions" that serve as lipophilicity modifiers or protectors of certain functional groups in therapeutic agent molecules.

[0410] In other illustrative embodiments, one or more other types of targeting delivery systems and methods can be used, for example, but not limited to, in whole or in part in one or more configurations: surfactants (surface-active substances) and or cosurfactants; enzymatic physical-chemical-based targeting; site-specific enzyme-activated targeting; vectors, such as ligand-based, non-viral-based, and Protein/DNA polyplex vector targeting; receptor-based chemical targeting; organic and or inorganic synthetic elements; transmembrane proteins (V-SNARE); peptides, including peptides that cross cell membranes and home specifically to certain diseases; nanostructured dendrimers and hyperbranched polymers; molecular Trojan horses; adenovirus, herpes simplex virus, adeno-associated virus or other virus vectors for targeted delivery that do not cause toxicity;

antibodies, including monoclonal antibodies; nanoparticles, including polymer nanoparticles like polymer, polybutylcyanoacrylate, and ethyl alcohol nanoparticles; immunotoxins; hormonal therapy; tissue-specific gene expression; gene therapy; pegylated immunoliposomes; anti-sense therapy; biological elements and or agents, including biological elements and agents conjugated with other agents, such as transferrin, but not limited to such; chemical elements and agents; devices, systems, and or mechanisms; liposomes, including liposomes conjugated with transferrin, but not limited to such; conformationally-constrained peptide drugs targeted at the blood-brain barrier; endogenous blood brain barrier and or blood tumor capillary transporters; inhibiting and or modulating blood brain barrier active efflux transporters; air and or other gas bubbles; blood brain barrier breaking and or disrupting elements and agents; blood brain barrier tight junction separating and or endocytoses elements and agents; vector-mediated delivery of opioid peptides to the brain; brain drug delivery of peptides and protein drugs via vector-mediated transport at the blood brain barrier, neurotrophic, neuroprotective, and various peptides and drugs, and the like.

[0411] In another illustrative embodiment, one or more elements cross various in vivo biological barriers, such as the transmucosal passage, and may also cross the bloodbrain barrier (BBB) and the blood-cerebrospinal fluid (CSF) barrier for targeted and or non-targeted in vivo delivery of CNS agents and elements. In one embodiment, one or more BBB-passing elements comprise small and or large molecule drugs.

[0412] Natural Clathrin, and in particular its ability to 'track' vesicle proteins leaving a synapse into the extracellular space (Granseth, et al 2007) indicates that the protein is not immediately scavenged by phages and other "house-cleaning" elements in the brain, and further, may move freely about CNS spaces. In one embodiment, one or more elements efficaciously move through the CNS spaces and comprise in situ elements for remediation, removal, and or sequestration of one or more types of contaminants, toxic elements, undesirable organic or inorganic elements, and the like.

[0413] In another embodiment, extensive modification and functionalization of agents and elements may not be required for CNS entrance and or BBB passage. Only minimal functionalization may be required, depending on cargo and element type.

[0414] In another embodiment, one or more CNS-entering and or BBB-passing elements of one or more types may behave as a drug by themselves—i.e., they efficaciously operate alone without carrying additional elements, e.g., cargo elements. In another embodiment, one or more elements of one or more types carry one or more additional elements of one more types past the BBB.

[0415] In another illustrative embodiment, one or more elements enter the CNS and or cross the blood brain barrier for targeted delivery of agents and elements, including, but not limited to, small and or large molecules, non-lipid-soluble micromolecules, macromolecules, light sources, hydrophilic and or hydrophobic agents, such as therapeutic, diagnostic, and prosthetic agents, and other structured cargo to specific cells and areas within the brain, and such agents and or cargo may comprise one or more sensor agents, assay agents, diagnostic agents, prosthetic agents, and also may

comprise agents like central nervous system drugs, antibiotics, and antineoplastic agents of one or more types, but are not limited to such.

[0416] In another embodiment, one or more elements are capable of circumventing the fluid-brain barriers by intracellular routes related to three separate and distinct endocytic processes. The three endocytic processes from the least to the most specific are fluid- or bulk-phase endocytosis, adsorptive endocytosis, and receptor-mediated endocytosis. [0417] There are several transport mechanisms and techniques known in the art to be involved in the uptake of nanoparticles by the brain across the BBB (Lockman et al. 2002, Begley, 2004, de Boer et al. 2007), one or more of which may be utilized in one or more invention embodiments. These mechanisms and techniques include: simple diffusion of lipophilic molecules, the BBB-specific influx transporters, including organic anion and cation transporters and transcytosis or endocytosis. In one embodiment, one or more elements are internalized at the BBB by one or two different endocytosis mechanisms: receptor-mediated endocytosis (RME) and adsorptive-mediated endocytosis (AME). AME is triggered by an electrostatic interaction between the positively charged moiety of the peptide and the negatively charged region of the plasma membrane. In contrast, RME is specific to certain peptides such as insulin and transferrin.

[0418] In one embodiment, delivery through the blood-brain barrier of one or more types of small or large molecule cargo elements, and or molecules with polar functional groups is accomplished via chimeric peptides. The latter are formed when a transportable vector, such as cationized albumin, lectins, or a receptor-specific monoclonal antibody, is conjugated to a therapeutic compound that is normally not transported through the BBB. In one embodiment, conjugation of drugs to transport vectors is facilitated by, but not limited to, the use of avidin-biotin technology. In another embodiment, chimeric peptides are not required to pass through the blood-brain barrier, depending on cargo and element types.

[0419] In another illustrative embodiment, one or more elements may be coated with one or more surfactants and or cosurfactants, including, but not limited to, polysorbate 20, 40, 60 and 80, and or with one or more other materials and substances to cross various biological barriers, such as the transmucosal passage, and also to overcome the blood-brain barrier (BBB), the transmucosal passage, and the blood-cerebrospinal fluid barrier (CSG) for targeted delivery of agents and elements nanoparticles. In another embodiment, surfactants and or cosurfactants are not required to achieve such BBB-passing functionality, depending on cargo and element type. E.g., in the prior art, it has been shown that using such surfactants and co-surfactants can cause an immunogenic response.

[0420] In another illustrative embodiment, one or more elements may be cationized to facilitate blood brain barrier passage. In another embodiment, cationization is not required to achieve such functionality, depending on cargo and element type.

[0421] In another illustrative embodiment, one or more elements cross the blood brain barrier due to disruption of the barrier by acoustic techniques, such as by using ultrasound

[0422] In another embodiment, zonula occludens toxin and its eukaryotic analogue, zonulin, (zot) are protein

ligands attached to one or more invention elements. Zonulin, the natural ligand of the Zot target receptor, interacts with these cargo attachment elements at the blood brain barrier, unlocking the tight junctions (TJ) in the brain that regulate the blood-brain barrier at that receptor. TJ-unlocking allows passage of one or more elements through the BBB, and thereby enables delivery of small and large molecules, non-lipid-soluble micromolecules, macromolecules, light sources, and other structured cargo elements to the brain. In another embodiment, Zonulin is not required to pass through the blood-brain barrier, depending on cargo and element types.

[0423] Extracellular pathways circumventing the fluid-brain barriers in humans are comparable in the CNS of rodents and a subhuman primate. The most highly documented extracellular route is through the circumventricular organs (e.g., median eminence, organum vasculosum of the lamina terminalis, subfornical organ, and area postrema), all of which comprise fenestrated capillaries and, therefore, lie outside the BBB. In one embodiment, blood-borne macromolecules; specifically fluid-phase molecules released by the invention; escaping fenestrated vessels supplying the circumventricular organs move extracellularly into adjacent brain areas located behind the BBB.

[0424] The potential intracellular and extracellular pathways that blood-borne substances carried within one or more elements may follow in various embodiments for circumventing the fluid-brain barriers and entry to the CNS are therefore numerous, and various invention embodiments are used as appropriate. One invention embodiment, for example, uses the nasal cavity as a route for delivery of one or more types of drugs and other agents, especially for systemically acting drugs that are difficult to deliver via routes other than injection. Embodiments for the use of the nasal cavity for drug delivery also extend to circumventing the blood brain barrier. Drugs have been shown to reach the CNS from the nasal cavity by a direct transport across the olfactory region situated at the loft of the nasal cavity. It is the only site in the human body where the nervous system is in direct contact with the surrounding environment. In one embodiment, the nasal route would be important for rapid uptake of one or more types of drugs used in crisis treatments and management, such as for acute pain, epilepsy, psychic agitation, and for one or more other types of centrally acting drugs where the pathway from nose to brain provides a faster and more specific therapeutic effect. Furthermore, in another embodiment, the trigeminal nerve and, in animals, the vomeronasal organ also connects the nasal cavity with the brain tissue. One or more methods of nasal delivery to the CNS, which may also be used by the instant invention, but not limited to, are described in Dhuria, et al, 2008; Ma et al, 2007; and Thorne et al. 1995.

[0425] The nasal cavity has a relatively large absorptive surface area and the high vascularity of the nasal mucosa ensures that absorbed compounds are rapidly removed (Mainardes, et al 2006). In one embodiment, two routes, singly or in combination, are used via which one or more types of molecules are transported from the olfactory epithelium into the CNS and/or CSF. The first is the epithelial pathway, where one or more types of compounds pass paracellularly across the olfactory epithelium into the perineural spaces, crossing the cribriform plate and entering the subarachnoid space filled with CSF. From here the molecules can diffuse into the brain tissue or will be cleared

by the CSF flow into the lymphatic vessels and subsequently into the systemic circulation. The second embodiment utilizes the olfactory nerve pathway, where compounds may be internalized into the olfactory neurones and pass inside the neuron through the cribriform plate into the olfactory bulb. In another embodiment, it is possible that further transport into the brain can occur by bridging the synapses between the neurons. After reaching the brain tissue, the drugs are cleared either via the CSF flow or via efflux pumps such as p-glycoprotein at the BBB into the systemic circulation. Despite the potential of the nasal route, there are some factors that limit the intranasal absorption of drugs. These barriers include the physical removal from the site of deposition in the nasal cavity by the mucociliary clearance mechanisms, enzymatic degradation in the mucus layer and nasal epithelium and the low permeability of the nasal epithelium removed (Mainardes, et al 2006). Colloidal carriers systems, such as nanoparticles and liposomes have demonstrated great efficacy in increasing drug bioavailability via the nasal route (Illum, 2002) In one invention embodiment, one or more elements comprise a colloidal carrier for enhanced nasal delivery of one or more elements, of one or more types.

[0426] Further, in one embodiment, it is possible to greatly improve the nasal absorption of one or more types of drugs and other elements by administering them in combination with an absorption enhancer that promotes the transport of the drug across the nasal membrane. Another invention embodiment comprises a nasal drug-delivery system that combines an absorption enhancing activity with a bioadhesive effect, which increases the residence time of the formulation in the nasal cavity. In one embodiment, this method can be even more effective for improving the nasal absorption of polar drugs. In one or more embodiments, a wide range of absorption enhancer systems can be utilized. In another embodiment, depending on cargo and element types, minimal functionalization may be required to take advantage of nasal absorption for efficacious passage to brain cells.

[0427] In another illustrative embodiment, one or more elements and in one or more configurations comprise in vivo and or in vitro sensor systems, assay systems, therapeutic drugs and other suitable methods to do genetic-based (trait-based) and or phenotype (state-based) drug dosing. In one embodiment, drugs are delivered at optimally effective and safe doses per each individual.

[0428] The invention, in one embodiment, provides for individual patient factors such as genotype, phenotype, age, gender, ethnicity etc., to be taken into account by one or more elements and factored into dosing and administration consideration. It has been demonstrated that inter-individual response variability can be 40-fold or more with practically all classes of psychotropic drugs. This makes it difficult to formulate rational guidelines for dosing and interpretation of biological parameters (such as plasma or serum drug concentrations) that might be associated with a therapeutic response. Although much remains unknown, a number of factors have been characterized as important determinants of patient-to-patient variability. These encompass genetics, disease state, nutritional status, concurrent use of drugs, and other pharmacoactive substances, including demographic factors such as age, gender, and ethnicity. Therefore, there is a requirement for in vivo systems that analyze many of these factors and dynamically adjust dosing accordingly.

[0429] In one embodiment, one or more elements comprise one or more personalized medicine elements, and which elements' efficacy may be increased, because responses arising from one or more individual variability factors; such as, but not limited to, genotype, phenotype, disease state, metabolic state, nutritional status, coninstant use of drugs, and other pharmacoactive substances, and also demographic factors such as age, and ethnicity; are factored into the elements, pre-delivery and or post delivery. Side effect profiles may also be reduced via such personalized medicine embodiments.

[0430] In one embodiment, one or more elements comprise one or more patented drugs; drugs that are about to go off patent; have already gone off patent (generics); and or their active metabolites, and which drugs' efficacy may be beneficially altered and or enhanced by use of the invention. These beneficial changes in the status of an existing drug may be achieved by the invention in one or more embodiments, for example, but not limited to: the ability to target specific areas in the body; to pass the blood brain barrier; to cross over into cells and their organelles; to fuse with cell membranes; to gain access to the cytosol; to offer the benefits of low antigenicity or minimal immunogenic effects; to modify, regulate, and or control cellular processes; to more efficiently and efficaciously carry drugs; and or to dynamically and or statically adjust the drug's responses and dosages arising from inter-individual variability due to one or more factors, such as, but not limited to, genotype, phenotype, disease state, metabolic state, nutritional status, coninstant use of drugs, and other pharmacoactive substances, and also demographic factors such as age, gender, and ethnicity of the patient. New patent filings for about to go off patent drugs and drugs already off patent may be enabled by one or more invention embodiments, such as affording increased drug efficacy, and or by enabling a better safety profile for the drug in question.

[0431] In various embodiments, the instant invention can carry one or more types of biomedical or healthcare elements, for example and without limitation: one or more therapeutic elements; pharmaceutical elements; diagnostic elements; assay elements; cosmetic elements; agents for treating one or more types of autoimmune diseases; agents for treating one or more types of infectious diseases; biological elements; radioactive agents or nuclear medicine agents; contrast agents; nano-scale biosensors; restorative agents; regenerative agents; cell, tissue, organ or circulatory repair elements; drug discovery agents; drug designer agents; drug research and development agents; drug fabrication agents; drug control and regulation agents; drug modifier agents; targeted drug delivery agents; clinical drug trial agents; antibiotics; antibacterials; vaccines; antiviral and anti-parasitic drugs; cytostatics; vitamins; proteins and peptides, including enzymes; hormones or other biological elements; prosthetic elements; intelligent nano-prostheses that supplement or enhance cell, tissue, or organ functioning; surgical elements; magnetic iron oxide nanoparticles; nano-scale biosensors; assays; diagnostic systems or nanodevices for in vivo delivery of targeted therapy to combat diseases, such as cancer and HIV, and the like, including other types and forms of drug elements for the diagnosis, cure, mitigation, treatment, prevention of disease. Some or all such elements may operate under the control and influence of various other elements and or methods and comprise another type of invention platform.

[0432] In another illustrative embodiment, one or more elements in whole or in part, cure, mitigate, or treat one or more types of bodily injuries and insults, including traumatic injury, blood clots, and the like, but not limited to.

[0433] In one embodiment, nano-engineered scaffolds comprised of a plurality of elements are able to support and promote cellular differentiation and growth in injured or degenerated regions.

[0434] In one illustrative embodiment, one or more elements comprise one or more types of small and or large molecules and may utilize one or more methods to enter the CNS and or cross the blood brain barrier, in whole or in part, for delivery of one or more assay, diagnostic, therapeutic agents, and drugs, of one or more types, to cells and or targeted areas within the brain, like, for example: contrast agents; central nervous system drugs; antibiotics; antineoplastic agents, which may be used for treating malignant brain tumors (primary and or metastasized, of one or more types) or benign neoplasms; Parkinson's agents; Multiple Sclerosis agents; epilepsy agents; meningitis agents; Alzheimer's disease agents; HIV infection agents; memory agents; stroke agents; coma agents; and the like; or comprise one or more psychotropic agents or therapies of one or more types to study, diagnose, cure, mitigate, or treat of one or more types of mental health and illness, including, but not limited to, stress; anxiety; depression; mania; bipolar disorder; attention deficit (hyperactivity) disorder; panic attacks; phobias; addictions; anger; rage; suicidal thoughts and tendencies; substance abuse disorder; post traumatic stress disorder; psychoses; mental retardation; autism; delirium symptoms; schizophrenia; neuroses; and or enhancing memory; cognition; cognitive functioning; the effects of cognitive therapy, and the like; including other types and forms of drug elements for the diagnosis, cure, mitigation, treatment, or prevention of one or more types of CNS diseases. In another illustrative embodiment, one or more elements enter the CNS, including crossing the blood brain barrier, in whole or in part, to diagnose, cure, mitigate, or treat one or more types of CNS injuries and insults, including traumatic brain injury, blood clots, and the like, but not limited to.

[0435] In one embodiment, one or more elements promote neuroprotection by limiting the damaging effects of free radicals generated after head injury, a major factor contributing to neuropsychiatric degenerative disorders (e.g., Alzheimer's).

[0436] In one embodiment, nano-engineered scaffolds comprised of a plurality of elements are able to support and promote neuronal differentiation and growth in injured or degenerated brain regions.

[0437] In another illustrative embodiment, one or more elements comprise a light source, for use, for example, but not limited to, in a photodynamic therapy (PDT) system for age related macular degeneracy (AMD).

[0438] Compounds such as drugs, amino acids, carbohydrates, proteins, nucleotide bases, hormones, pesticides and co-enzymes have been successfully used in the prior art for the preparation of selective recognition matrices. A wide variety of print molecules have been used in various imprinting protocols known in the art. Of all the imprinting strategies known in the art, it has become evident that the use of non-covalent interactions between the print molecule and the functional monomers is the more versatile. The apparent weakness of these interaction types, when considered indi-

vidually, may be overcome by allowing a multitude of interaction points simultaneously. Together with the fast association and dissociation kinetics of these bond types, so that in a short time many possible combinations can be checked before the correct partners associate, this protocol has proven advantageous. Furthermore, the use of noncovalent interactions in the imprinting step closely resembles the recognition pattern observed in nature. Example invention molecular imprint embodiments in the art include, but are not limited to:

[0439] Fragmented polymer monoliths

[0440] Composite polymer beads

[0441] Polymer beads from suspension, emulsion or dispersion polymerization

[0442] In-situ polymerization

[0443] Polymer particles bound in thin layers

[0444] Polymer membranes

[0445] Surface-imprinted polymer phases

[0446] In one illustrative embodiment, the invention uses molecular-imprint technology, wherein biodegradable films are used as a pliable template for elements, which elements are pressed into a film and then removed, leaving a physical mold of the element's shape. In one embodiment, this can facilitate catalysis of certain reactions and may also be used for shape selective separations. In other embodiments, imprinted polymers may facilitate the fabrication of elements to achieve selective diffusion; as chromatographic supports for the separation of enantiomers and oligonucleotides by invention elements; to provide the recognition element for an invention chemical sensor; and for the synthesis of polymeric materials that mimic biological cargo attachment elements and are targeted by invention elements, and or play a role in the design of new drugs. In one embodiment, this invention process provides for imprinted biodegradable capsule production with target or site-specific feature sizes at the molecular level. Other invention embodiments may utilize imprinted membranes and thin films that also function as an artificial cell wall for the selective transport of targeted drugs, peptides and biologically important molecules.

[0447] Surface imprinting involves the following steps: The print molecule, usually a large one, is first allowed to form adducts with functional monomers in solution and the formed elements are subsequently allowed to bind to an activated surface such as silica wafers or glass surfaces. Thus, with this technique, a designed imprinted (imaged) surface is obtained. This approach should potentially be valuable for creating specific cell binding surfaces. When preparing molecularly imprinted polymer monoliths against large imprint species, there is a risk of permanent entrapment of the template in the polymer after polymerization. When using thin polymeric layers or imprinted surfaces this drawback may be overcome.

[0448] In one embodiment, imprinted nanocapsules using techniques known in the art and as discussed above, one or more elements utilize and or constitute a nanocapsule with manifold, multi-tiered capabilities for in vivo administration and targeted delivery. The imprinted nanocapsule is delivered in vivo to detect and target a particular in vitro imprinted biological element, which may be, but is not limited to, a particular type of receptor, protein, or cell, since its imprint shape on the nanocapsule will only bind in vivo to that particular biological element target. The molecular-level imprint process thereby provides for targeting one or

more elements using biodegradable nanocapsules for in vivo agent delivery. In addition, vectors and targeting moieties, and blood brain barrier, transmucosal, and CSF barrier breaching elements, and other elements and substances may also be attached to the surface of the molecular imprint nanocapsule or otherwise be conjugated to it.

[0449] In another illustrative embodiment, one or more elements may be used in conjunction with molecularly imprinted polymers known in the art as recognition elements in biosensor-like devices. In one embodiment, imprinted polymer embodiments may be highly resistant sensing element alternatives.

[0450] In another illustrative embodiment, one or more elements are encapsulated in whole or in part in one or more biodegradable controlled-release polymers, which polymers may also be conjugated with other elements and agents. The polymer capsule, and or one or more elements may also be coated with one or more surfactants and or cosurfactants and or with other materials and substances. One or more targeting and or masking moieties and or other targeting vectors may also be attached on the polymer surface, and or on one or more elements.

[0451] In one embodiment, one or more elements are put into one or more biodegradable controlled-release polymeric capsules, and these elements transform "dumb" polymeric delivery capsules into "smart" systems.

[0452] In the instance of polymeric nanocapsules, which may be molecular imprinted or not, illustrative controlled-release polymeric nanocapsule embodiments of the invention may include one or more of the following delivery systems, but not limited to, and in one or more configurations:

- [0453] 1. Diffusion-controlled systems
- [0454] 2. Water penetration-controlled delivery devices
- [0455] 3. Chemically controlled systems

[0456] 4. Drugs covalently attached to polymer backbone systems, which delivery systems can be further subdivided into soluble systems and insoluble systems. Insoluble systems are used as a subcutaneous or intramuscular implant for the controlled release of the chemically tethered therapeutic agent. Soluble systems are used in targeting applications.

[0457] 5. Drug release determined predominantly by erosion systems, whereby certain polymers can undergo a hydrolysis reaction at decreasing rates from the surface of a device inward, and under special circumstances the reaction can be largely confined to the outer layers of a solid device. Two such polymers are poly (ortho esters) and polyanhydrides, because the rates of hydrolysis of these polymers can be varied within very wide limits, considerable control over the rate of drug release can be achieved.

[0458] 6. Poly (ortho esters) systems, which are highly hydrophobic polymers that comprise acid-sensitive linkages in the polymer backbone.

[0459] 7. Polyanhydrides materials as bioerodible matrices for the controlled release of therapeutic agents. Aliphatic polyanhydrides hydrolyze very rapidly while aromatic polyanhydrides hydrolyze very slowly, and excellent control and regulate over the hydrolysis rate can be achieved by using copolymers of aliphatic and aromatic polyanhydrides. In this way, erosion rates over many days have been demonstrated, and erosions rates measured in years have been projected.

[0460] The form in which the foreign moiety, vector and or cargo are held within one or more elements will depend on the release properties and methods required. For release

at the targeted site, it will be important to ensure that the right conditions prevail, for example, to permit cell localization and internalization via receptor mediated endocytosis.

[0461] In one illustrative embodiment, the invention enables one or more types of delivery systems that engage in an iterative, interactive, and dynamic dialog with one or more targets; follow a sequence of actions governed by biological control laws and methods; and or use behaviors and methods as defined by graphs and or an algebra, for example, a Lie algebra. In one illustrative example, one or more elements follow an algorithm expressed by the invention, such as in this illustrative embodiment:

[0462] 1) One or more elements, that may be with or without cargo elements, docks and or loiters on or near one or more cell membranes,

[0463] 2) One or more elements enter one or more target cells, while one or more other elements continue to loiter nearby or stay docked at the cell membrane.

[0464] 3) The docked and or loitering element elements wait for a time period,

[0465] 4) The targeted cell produces one or more reactions, for example, manufactures and or secretes an agent in response to the element's docking and or delivering its cargo,

[0466] 5) The docked element and or loitering elements analyze the new cell behavior and or its secretions,

[0467] 6) The docked element or loitering elements undergo a conformational change in response to the cell's new behavior,

[0468] 7) The docked element and or loitering elements self-adapt, producing yet another conformational change in the cell, and or releases another round of one or more agents that are taken up by the targeted cell, and

[0469] 8) The foregoing process is repeated as required to achieve an efficacious effect.

[0470] In another embodiment, one or more light sources comprised of one or more elements operate in an intelligently staged sequence or orchestrated series of actions, which may be multiplexed or done in parallel by using one or more light and thermal energy emitting sources and methods. By using one or more light and or thermal energy emitting sources, optical and or thermal energies from one or more light sources operate on one or more photosensitive and or thermal sensitive elements comprising one or more elements that also comprise one or more entrapped agents. This method results in a staged series of overall actions that follow an intelligently ordered sequence of events. In an example embodiment, first a diagnostic agent from one or more elements is released by an optical and or thermal trigger, and the agent's positive finding of a disease, like cancer or HIV then causes one or more therapeutic agents to be released from the same and or other one or more other elements by one or more optical and or thermal triggers. Agent dosages are released in calculated amounts, and the dosages may be non-targeted or targeted.

[0471] In another illustrative embodiment, cavity-forming cargo elements have one or more compartments that in whole or in part are separated by one or more barriers, for example, but not limited to, one or more phospholipid membrane barriers and or one or more barriers comprised of molecular-imprinted films. The barriers may exhibit structural transitions due to internal or external stimuli. In one

embodiment, agents or cargo entrapped within one or more elements remain sequestered within their respective compartments until a change in barrier permeability state is triggered by contact, for example, by a ligand, with one or more specific targets or sites. The subsequent biochemical and or biological reactions cause the barriers to alter states into an opened state and release entrapped cargo and agents from one or more invention elements. In one example embodiment, binary mixtures of therapeutic and or diagnostic agents are mixed together as needed to dynamically and more efficaciously deal with a disease or disorder.

[0472] The invention, in one or more embodiments, comprises in whole or in part one or more elements, components, devices, systems, and the like, of one or more types, formed by using one or more engineering disciplines and related engineering technology disciplines of one or more types. Listed below are some such example invention embodiments, but are not limited to.

[0473] In one embodiment, the invention remedies the deficiencies of prior art by providing one or more elements of one or more types, a plurality of which may also comprise one or more nanoscale platforms of one or more types. A platform according to the invention may be used, for example, in biomedical, electronics, telecommunications, and information processing applications.

[0474] FIG. 6 is an exemplary energy level diagram 600 illustrating the energy levels associated with a hyperfine interaction between electron and nuclear spin in the presence of magnetic fields of the type used to do ESR spin label studies, which may be done in vivo and in vitro in one invention embodiment. The hyperfine interaction is a strictly quantum mechanical phenomenon. In an atom, the electron possesses an intrinsic quantum mechanical quantity known as spin. The nucleus of an atom also possesses spin. Intrinsic spin tends to generate a spin magnetic moment that is capable of interacting with other magnetic moments and fields. Generally, the spin magnetic moment of the nucleus does not interact with the spin magnetic moment of the electron. However, in the presence of a strong magnetic field, the spin magnetic moments of the electron and nucleus become coupled and interact.

[0475] In one illustrative embodiment, the electron is excited using pulses of electromagnetic radiation while maintaining its spin configuration. The source of the electromagnetic radiation may be, for example, an ordinary lamp, an LED, a time-varying magnetic field generator, a laser, or an electromagnetic field generator. A hyperfine interaction gives rise to electron nuclear double resonance (ENDOR) techniques. According to one illustrative embodiment of the invention, room temperature EPR and ENDOR techniques known in the art are used for performing in vivo spin probe studies.

[0476] In another embodiment, one or more elements comprise one or more diagnostic agents, and during the same NMR/MRI, or EPR, or ESR, or ESEEM, or ENDOR, or PET, or SPECT, or OCT operation, one or more elements use quantum information processing techniques known in the art can modify, process, manipulate, encode and decode, input, output, transmit, communicate, store and read information using one or more modulated signals, methodologies, or carrier signals of one or more types.

[0477] In one embodiment, one or more invention elements in one or more configurations, are bonded, tethered, or otherwise incorporated into one or more invention and or

non-invention elements, comprising functionalized nanoscale elements, components, devices, systems, and or platforms such as, but not limited to, nano-lasers, quantum dots; photonic dots; nanoscale DNA chips; protein assay chips; assay elements; environmental, protein, phenotype, DNA, and or metabolic assay and analysis elements.

[0478] In another embodiment, one or more elements may comprise a bio-lasing structure, in vivo or in vitro.

[0479] In one embodiment, one or more elements in one or more configurations comprise nano-sensor elements; including, but not limited to, radioactivity sensors; chemical sensors; biological sensors; electromagnetic sensors; acoustic sensors; visible, infrared, and or ultraviolet wavelength sensors; tactile sensors; pressure sensors; volumetric sensors; flow sensors; and temperature sensors; and one or more of which sensors may constitute a bio-molecular device.

[0480] In one embodiment, one or more elements and or platforms utilize and or employ one or more types of transmitter and or receiver elements as sensors and or for transmission of information of one or more types in vivo and in vitro.

[0481] In another embodiment, one or more elements and in one or more configurations comprise one or more nanoscale elements, components, devices, systems, and or platforms that input, read out, process, analyze, output and report on information gathered by one or more types of diagnostic, test, label, tag, reporter, sensor, and or assay elements.

[0482] In one embodiment, quantum dots and or photonic dots are released in vivo or in vitro from one or more elements, and the quantum dots and or photonic dots are coated in whole or in part in one or more surfactants, cosurfactants, and other materials or sequestering substances.

[0483] In one embodiment, quantum dots are tagged to one or more elements. The specific wavelength glow of the quantum dots enables the identification of specific pathologies, disorders, metabolic states, proteins or DNA making it possible to diagnose various diseases.

[0484] In one embodiment, one or more nanoscale quantum dot assays using tiny permutations of color tag a million or more different proteins or genetic sequences in a process called multiplexing. In one embodiment, one or more quantum dots of various sizes are excited at the same wavelength but have different emission wavelengths, and act as probes in experiments where multiple fluorescent measurements need to be made simultaneously, such as flow cytometry or confocal microscopy.

[0485] In another illustrative embodiment, one or more elements are sufficient to implement in vivo or in vitro genetic and protein nanoscale optical biological assay systems and methods. In one illustrative configuration, one or more elements comprise one or more nano-scale DNA chips known in the art, and or one or more nano-scale DNA chips known in the art to detect DNA samples formed from bonding with the target DNA on a chip, and or reference DNA nano-chips.

[0486] In another illustrative configuration, one or more elements comprise one or more protein array techniques known in the art. The array surfaces are designed to bind to one or more hydrophobic, hydrophilic (cation or anion) or specific ligands, and also include a protein array reader known in the art.

[0487] In another illustrative embodiment, one or more elements are used in a multiplexed analysis system or method that provides a nanoscale replacement for DNA-chip technology and can be used for the analysis of genetic variance, proteomics, and gene expression.

[0488] In another embodiment, one or more elements produce specific light emissions and or thermal energies caused by their coming into contact with a particular metabolic state, medical disorder, disease pathology, genotype, phenotype and or other specific stimuli. One or more entrapped agents carried by one or more elements are thereby selectively triggered and released. In doing so, they form a targeted agent delivery system without exposing the entire body—or an indiscriminate area—to a similar dose of light, thermal energy, and or agents. The agents may be delivered in vivo by means known in the art.

[0489] In one illustrative embodiment, photonic energies from one or more elements thermally operate on one or more other elements that may have one or more entrapped materials, such as, but not limited to, therapeutic, diagnostic, and or therapeutic agents within an aqueous interior, and or that may have one or more entrapped nanoparticles such as liposomes, micelles, proteins, other biological and or bioengineered elements, including organic, inorganic, and synthetic materials, and or that may have one or more hydrophobic materials bound to a lipid bilayer membrane. The well-known permeability increase at the phase transition temperature provides a means to trigger release of an entrapped agent, like, for example release of a therapeutic agent in locally heated tissues. In one embodiment, efficient in vivo or in vitro release of entrapped agents at non-targeted and or targeted sites are triggered by light emitted by one or more light sources when the one or more elements comprise a photoisomerisable species.

[0490] In another embodiment, the method of one or more LuxR proteins and lux bioluminescence genes and or other luminescent causing genes known in the art are utilized and are bioengineered and incorporated into one or more elements, ligands, targeting moieties, and or vectors, which may also be conjugated with one or more other elements, materials, and substances. In one embodiment, luminescent causing genes provide optical pumping sufficient to excite one or more quantum dots and or photonic dots.

[0491] In an illustrative embodiment, in vivo release from one or more cargo elements comprised of one or more entrapped liposomal and or non-liposomal-entrapped agents are optically triggered by photons emitted by light sources of one or more types. In one illustrative embodiment, one or more light sources produce specific light wavelength emissions caused by their coming into contact with, for example, a specific disease at in vivo target site and causes diagnostic, therapeutic, and or prosthetic agents comprised in a photosensitive invention delivery system to be triggered and released from one or more invention elements, thereby forming a highly targeted drug delivery system. For example, in one embodiment, one or more cargo elements comprise an amphipathic lipid, such as a phospholipid, having two chains derived from fatty acid that allow the lipid to pack into a bilayer structure. One or more photosensitizers may be incorporated into the entrapped materials' cavity and or membranes.

[0492] In one illustrative embodiment, a phospholipid (1,2-(4'-n-butylphenyeazo-4"(-phenylbutyroyl))-glycero-3-phosphocholine ('Bis-Azo PC'), is substituted with azoben-

zene moieties in both acyl chains that can be photoisomerised by a fast nanolaser pulse. One or more other photoisomerisable species can be used in other embodiments. Agent release from one or more cargo elements occurs on the milliseconds timescale and photosensitised cargo elements thereby serve as light sensitive elements to allow for the triggered release of agents from one or more invention elements. In one embodiment, cholesterol additives may be used. The addition of cholesterol may have a marked effect on kinetics of agent release from cargo elements, and in some circumstances can result in substantial enhancement of light sensitivity in one or more photosensitised elements comprising one or more invention elements, In another embodiment, thermal and photosensitive activation systems acting together comprise one or more elements.

[0493] The invention, in one embodiment, comprises an in vitro and or in vivo nanoscale, biomolecular electronics element and or nano-electronics element, i.e., bio-molecular devices, which may be employed in a scalable, intelligent, biomolecular electronics device platform and or a nano-electronics device platform. The platform may also be comprised of one or more non-invention elements and devices, such as crystals, conductors, insulators, semiconductors, MEMS, and circuits, but not limited to such. And further, the platform may also be coated in one or more surfactants and or cosurfactants and or metals, elements, materials and substances.

[0494] In one embodiment, one or more elements and or platforms are used for biomolecular electronic and or nanoelectronic devices. Biological molecules, particularly proteins and lipids are used to perform the basic properties necessary for the functioning of biomolecular electronic devices. These biological materials conduct and transfer molecules from one location to another, are capable of major color changes on application of an electric field or light and can produce cascades that can be used for amplification of an optical or an electronic signal. All these properties can be applied to electronic switches, gates, storage devices, biosensors, biological transistors, to name just a few. In general, the electrical properties of bilayer lipid membranes are easily measurable for signal generation and transduction. In one embodiment, hybrid elements comprising cells with intact plasma membranes can be considered to act as tiny capacitors under the influence of an electric field. Whereas sufficiently high field strength may increase the membrane potential past a critical point leading to the breakdown of the membrane, experimental care must be taken. (Dielectric breakdown of biological membrane occurs at about 1 volt across the membrane.) On the other hand, the use of electrostatic potentials around the lipid molecules is very attractive, because they are controllable.

[0495] In one embodiment, one or more elements comprise nanoscale elements, components, devices, systems and or platforms, in one or more configurations, which form connectors for carrying information from a storage, processing or communications element or device to another, of one or more types.

[0496] In one embodiment, one or more elements comprise one or more information processing elements, components, devices, systems and or platforms such as, for example, but not limited to, encoders and decoders, memory, logic gates, registers, circuits, wiring and connectors, input

and output elements, analog to digital and digital to analog converters and system architectures known in the art.

[0497] In one embodiment, one or more invention elements comprise nanoscale elements, components, devices, systems and or platforms that modify, process, manipulate, encode and decode, input, output, transmit, communicate, store and read various forms and types of information using a variety of suitable techniques known in the art, in vivo and in vitro.

[0498] A scalable information-processing invention platform may also include an encoder, e.g., a predetermined or specific DNA sequence that deliberately encodes at least a subset of the elements to take the form of specified sequence, as well as a decoder for reading information from at least a subset of the protein-based information processing elements. Examples of such a bio-system decoder are, but not limited to, a dye-based protein assay, a quantum dot-based assay, or other protein assay methods known in the art. Another example of encoders/decoders is the use of NMR and ESR and other methods known in the art that can effect and discern protein behaviors and their physical characteristics. Another example of encoders/decoders is the use of photons of different wavelengths and photo detectors.

[0499] In one embodiment, one or more elements comprise in vitro and or in vivo nanoscale information processing elements, components, devices, systems and or platform, which may follow and execute algorithms of one or more types expressed by or use biological control and regulate laws, processes, and or methods, and or geometrically derived algorithms such as graphs and Lie algebras, including Clifford algebras, but not limited to.

[0500] In another embodiment, one or more elements comprise a cognitive information processing element, device, and or platform of one or more types that follow and execute algorithms expressed by or use biological control and regulate laws and or processes, and or geometrically derived algorithms such as graphs and Lie algebras, including Clifford algebras, but not limited to.

[0501] In another embodiment, one or more elements comprise a hybrid digital and analog information processing element, device, and or platform of one or more types, wherein enlisting the rich repertoire of biochemical reactions and adopting a nested hierarchical organization makes intermixing of digital an analog processing possible in bio-computing applications.

[0502] In one embodiment, one or more elements comprise one or more nanoscale information processing elements, components, devices, systems and or platform that utilize photons emitted by invention light sources of one or more types as the basis of computation and or transmission and communication.

[0503] According to one illustrative embodiment, one or more elements comprise one or more nano-computer elements, components, devices, systems and or platforms of one or more types that are programmable, and or autonomous acting, and or do cognitive processing, which bionano-computers may also utilize self-replicating, self-adapting, self-repairing, self-regulating, and or self-regenerating methods, and which are used for applications at the cellular, molecular, and nanoscale level that may include, but are not limited to, biomedical imaging, sensors, diagnostic systems, assay systems, therapeutic systems, drug delivery systems, prosthetic systems, cybernetic systems, cellular-level nanofabrication systems, and inter- and intra-cellular imaging,

repair, and engineering systems, the monitoring, sensing, imaging, diagnosing, repairing, constructing, fabricating, and or control and regulating of organic and or inorganic elements, and which bio-nano-computer elements and or platforms also may utilize and leverage biological control and regulate laws and or methods, and or geometrically derived algorithms such as graphs and Lie algebras, including Clifford algebras, but not limited to, in the performance of their tasks.

[0504] In one illustrative embodiment, one or more element chains are created via a molecular bridge group. To align the elements with respect to one another and also with respect to an external magnetic or electrical field. In one embodiment, one or more elements and or platforms and in one or more configurations are embedded in another material, like liquid crystal.

[0505] In one embodiment, one or more elements and or platforms and in one or more configurations are coated completely and or partially in a metal.

[0506] In another embodiment, one or more elements and or platforms and in one or more configurations are coated completely and or partially in reflective and or non-reflective coatings.

[0507] In one embodiment, one or more elements and or platforms and in one or more configurations are used to coat completely and or partially metals, crystals, insulators, conductors, semiconductor components, wires, and devices.

[0508] In another illustrative embodiment, one or more elements and or platforms and in one or more configurations facilitate the externally and or mechanistically directed alignment of, for example, but not limited to, biological elements, various other non-invention nanoparticles, carbon nanotubes, crystals, conductors, semiconductors, insulators, and or other devices, materials and substances, which aligned assemblies may further be coated in one or more surfactants and or metals, elements, materials and substances

[0509] In one embodiment, one or more elements in one or more configurations include other types of nanoparticle elements such as, but not limited to, polymer-based, polybutylcyanoacrylate-based, and cetyl alcohol-based nanoparticles, empty cage Fullerenes, endohedral Fullerenes, carbon nanotubes, cells, liposomes, capsids, dendrimers, micelles, and the like.

[0510] In another illustrative embodiment, one or more elements and or platforms of one or more types in whole or in part enable a shape programmable and or scaffolding system to which one or elements of one or more types, including natural and or non-invention elements are affixed and or further form more one or more structures of one more types

[0511] In one embodiment, one or more elements and or platforms in one or more configurations form and or include optical elements such as, but not limited to, optics; optoelectronic elements; photoelectric elements; photodetectors; and photosensitive elements, which optical elements may also be coated or treated in whole or in part with materials that affect their optical properties.

[0512] In one embodiment, one or more elements and or platforms and in one or more configurations form and or include imaging elements and sensors, such as, but not limited to, CCDs and CMOS optical elements.

[0513] In one embodiment, one or more elements and or platforms, in one or more configurations include and or comprise photonic to electrical energy conversion elements.

[0514] In one embodiment, one or more elements and or platforms form one or more electronic circuits, which circuit may also be comprised of one or more other elements such as empty Fullerenes, endohedral Fullerenes, nanotubes, crystals, insulators, conductors, semiconductors, and or other materials, substances and devices, which circuits also may be coated in one or more surfactants and or cosurfactants and or other materials and substances.

[0515] In one embodiment, one or more elements and or platforms are switched on or off and or change states by applying an electric field, and may also comprise one or more transistors or devices in another embodiment.

[0516] In another embodiment, one or more elements and or platforms and in one or more configurations; self-assemble, and or are shape-programmed, and or use biological control and regulate laws, processes and methods, and or use geometrically derived algorithms such as graphs and Lie algebras, including Clifford algebras, but not limited to, and or are mechanically assembled via lithography, and or utilize other externally directed techniques and methods known the art, and or some combination thereof; form natural positions that are associated with electronic circuits and or information processing devices, such as atomic and molecular scale device design, their interconnection, nanofabrication and circuit architectures.

[0517] According to one illustrative embodiment, one or more elements and or platforms comprise one or more crystal structures and elements, of one or more types.

[0518] According to one illustrative embodiment, one or more elements and or platforms comprise one or more desiccated elements, of one or more types.

[0519] According to one illustrative embodiment, one or more invention comprise one or more hydrated and or rehydrated elements and or platforms, of one or more types.

[0520] According to one illustrative embodiment, one or more elements and or platforms comprise one or more rehydration elements and or platforms, of one or more types.

[0521] According to one illustrative embodiment, one or more elements and or platforms are embedded and or incorporated into one or more materials, substances, devices, agents, devices, systems, organisms, and or mechanisms of one or more types.

[0522] In another illustrative embodiment, one or more elements and or platforms comprise one or more magnetic nanoparticles of one or more types.

[0523] In one embodiment, one or more elements and or platforms are nanoscale recording memory media or components, which may incorporate metals, ferromagnetic materials, and or ferroelectric materials and elements, and or may form into magnetic rings, and or may form vertically polarized magnetic domains and or form magnetic domains on isolated islands of one or more types.

[0524] In one embodiment, one or more elements and or platforms are nanoscale photovoltaic cells or components of one or more types.

[0525] In one embodiment, one or more elements are nanoscale batteries or components of one or more type for storing electronic charge.

[0526] In one embodiment, one or more elements and or platforms comprise a nanoscale environmental hazard-

screening device, and or comprise an in situ remediation, removal and or sequestration component or system of one or more types.

[0527] In one embodiment, one or more elements and or platforms comprise an opto-electronic device, system or component of one or more types.

[0528] In one illustrative embodiment, embodiment, one or more elements comprise one or more nanoscale passive and or active linear or nonlinear optic components, and or particle detectors, and or other elements sufficient to implement in vivo or in vitro optical system arrays and methods.

[0529] In another embodiment, one or more elements comprise in vivo or in vitro detection, diagnostic and tracking agents for chemical, biological, and or nuclear elements and activities, but not limited to such.

[0530] In one embodiment, one or more elements and or platforms comprise a spin-based electronics element or system of one or more types.

[0531] In one embodiment, one or more elements and or platforms exploit the Coulomb blockade-like properties of self-assembled proteins, wherein a single particle at a time may move through a transmembrane protein-based channel. [0532] In one embodiment, one or more elements and or platforms utilize and or exploit the Casimir effect, which is a small attractive force that acts between two closely parallel, uncharged conducting elements. It is due to quantum vacuum fluctuations of the electromagnetic field.

[0533] In some illustrative embodiments, one or more elements and or platforms and in one or more configurations are physically linked via molecular addends of one or more types, but are not limited to such addend types.

[0534] In other illustrative configurations, one or more elements and or platforms are functionally linked via photonic, chemical, electromagnetic, electrical and/or quantum (non-classical) interactions of one or more types, including the Internet, to work and cooperate locally and/or remotely. [0535] One or more elements and or platforms of one or more types may be encapsulated, packaged, stored, incorporated, and or utilize one or more methods known in the art, including for example, but not limited to: catheters; injections, including intramuscular injections; syringes; droppers and bulbs; pills; intravenous means; oral means; anal means; capsules; nanocapsules; nanoparticles; nano-devices; prescriptions; hospital and medical supplies; dental supplies; non-prescriptions; medications; over the counter products and remedies; alternative medicine supplies, systems, products and devices; hair care products; splints, casts, walkers, crutches, canes, wheelchairs, and other ambulatory aids; natural foods; vitamin and mineral supplements; first aid products; emergency health care procedures, systems, devices, and products, including combat medicine; health care products; grafts; skin patches; bandages; adhesives; wraps; masks; markers; powders; granules; geriatric care products; pediatric care products; diagnostic devices, systems, and products; medical imaging devices, systems, and products; telemedicine devices, systems, and products; in vivo monitoring systems, products, systems, and devices; in vitro monitoring systems, products, systems, and devices; laundry products; chemical, nuclear and biological sensors; sensors; bio-sensors; environmental sensors; combat systems, clothing, uniforms, and protective gear; food preparation products; food testing and safety devices, systems, and products; food storage wraps, systems, devices, and products; water treatment devices, systems and products; waste storage, management, and treatment systems and products; sewerage systems and products; plumbing systems and products; bed and bath products; animal care and veterinary products; animal feed; animal slaughter systems and products; cooking products; cookware; forensic devices, systems and products; home and office cleaning products; home products; office products; personal products; industrial products; home and office care products; paper products; personal hygiene products; sexual hygiene and safety products; sexual reproduction devices, systems, and products; sexual arousal products and devices; dental and dental care products; oral hygiene products, devices, and systems; robotic products, systems and devices; cybernetic devices; jewelry; novelties; solvents; agro-products; plants; animals; vehicles; biologicals; chemicals; cells; tissue; organs; proteins; liposomes; phages; micelles; peptides; antibodies; monoclonal antibodies; DNA; RNA; IRNA; siRNA; RISC; cloning; human contact; micro-electromechanical systems (MEMS) and other types of nano-systems; food utensils; tools; appliances; consumer electronics; paints and finishes; heating, ventilation and air conditioning systems; construction, building, home and office materials; water; milk; food and other edible or chewable substances and items; pros-

<160> NUMBER OF SEQ ID NOS: 30

theses; food and drink additives and supplements; drinks; beverages; soaps; creams; ointments; salves; topical agents; cosmetics; beautifying agents; liquids; fluids; oils; gels; adhesives; aerosols; vapors; airborne methods; pumps; fragrances and perfumes; textiles; sporting and athletic goods and devices; physical work out and training systems, devices, and products; sports medicine systems, devices, and products; recreational products and gear; shoes, clothing, and apparel; eyewear; sprays; dyes; biological elements; organ; implants; stents; prosthetic devices; artificial skin, blood, limbs, joints, bones, cells, eyes, organs, and other artificial body parts and biological elements; subcutaneous means; incisions; surgical means; and in-patient and outpatient medical procedures.

[0536] The above-described embodiments have been set forth to describe more completely and concretely the present invention, and are not to be construed as limiting the invention. It is further intended that all matter and the description and drawings be interpreted as illustrative and not in a limiting sense. That is, while various embodiments of the invention have been described in detail, other alterations, which will be apparent to those skilled in the prior art, are intended to be embraced within the spirit and scope of the invention.

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Gln 865	Ile	Gln	Glu	Gly	Сув 870	Glu	Glu	Pro	Ala	Thr 875	His	s Asr	n Ala	Leu	880
Lys	Ile	Tyr	Ile	Asp 885	Ser	Asn	Asn	Ser	Pro 890		Суя	Phe	e Leu	Arg 895	g Glu
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Arg 945	Tyr	Leu	Val	Cys	Arg 950	Lys	Asp	Pro	Glu	Leu 955	Trp	Ala	a His	Val	. Leu 960
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ГÀа	Ala	Phe 995	Met	Thr	Ala	Asp	Leu 1000		aA c	n Gl	u Le		le G	lu I	eu Leu
Glu	Lys 1010		e Val	l Leu	ı Asp	Asr 101		er V	al P	he S		lu .020	His	Arg	Asn
Leu	Gln 1025		ı Lev	ı Lev	ı Ile	Leu 103		nr A	la I	le L		la .035	Asp	Arg	Thr
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Asp	Ile 1055		s Sei	: Ile	e Ala	Val 106		er S	er A	la L		'yr .065	Glu	Glu	Ala
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Val	Leu 1085		e Glu	ı His	; Ile	Gl _y 109		n L	eu A	.sp A	-	Ala .095	Tyr	Glu	Phe
Ala	Glu 1100	_	g Cys	s Asr	ı Glu	Pro 110		La V	al T	rp S		3ln .110	Leu	Ala	Gln
Ala	Gln 1115		ı Glr	ı Lys	a Asp	Leu 112		al L	ys G	lu A		le 125	Asn	Ser	Tyr
Ile	Arg 1130	-	Asl) Asp) Pro	Sei 113		er T	yr L	eu G		al .140	Val	Gln	Ser
Ala	Ser 1145		g Sei	Asr	ı Asn	Trp		lu A	ap L	eu V		ys .155	Phe	Leu	Gln
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Leu	Ala 1595		Pro	э Туз	Phe	11e		ln V	al M	et A	Arg	Glu 1605	Tyr	Leu	Ser
ràs	Val 1610		Ly:	s Lev	ı Asp	Ala 161		∋u G	lu S	er I	₋eu	Arg 1620	Lys	Gln	Glu
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)> SE														
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Met 65 Ala Thr	Ile 50 Asn Ile	Ile 35 Arg Asp Met	Glu Pro Asn Ile	Phe Lys Ser Pro 85	Ser Val Asn 70 Ala Asn	Thr Gly 55 Pro Ser	Leu 40 Glu Ile Lys Glu	25 Thr Gln Arg Val Met 105	Gln Met Ala Arg Ile 90 Lys	Glu Gln Pro 75 Ala	Va. 60 Ile	u Gly 45 45 Val Val E Sen u Lys	30 Lyss Lyss Ala Lyss Ala Lyss Lyss Lyss Lyss Lyss Lyss Lyss Lys	15 Asi Phe Ile Asi Asi 95 Ala	n Pro e Ile e Asp o Ser 80 v Lys
Met 65 Ala Thr	Ile 50 Asn Ile Leu	Ile 35 Arg Asp Met Gln Thr	20 Gly Glu Pro Asn Ile 100 Asp	Phe Lys Ser Pro 85 Phe Asp	Ser Val Asn 70 Ala Asn Val	Thr Gly 55 Pro Ser Ile Thr	Leu 40 Glu Ile Lys Glu Phe 120	25 Thr Gln Arg Val Met 105 Trp	Gln Met Ala Arg Ile 90 Lys	Glu Gln Pro 75 Ala Ser	Va. Va. 600 Ile	Glywar Glywar Asp 45 Val Val Lys Met Serial Lys	7 Ile 30 Lys 1 Ile 2 Als 3 Als 110	15 Asi Phe Ile Ile Ile Ile Ile Ile Ile Ile Ile Il	n Pro in
Met 65 Ala Thr Thr	Ile 50 Asn Ile Leu Met Ala 130 Ser	Ile 35 Arg Asp Met Gln Thr 115	20 Gly Glu Pro Asn Ile 100 Asp	Phe Lys Ser Pro 85 Phe Asp	Ser Val Asn 70 Ala Asn Val	Thr Gly 55 Pro Ser Ile Thr Asn 135	Leu 40 Glu Ile Lys Glu Phe 120 Ala	25 Thr Gln Arg Val Met 105 Trp Val	Gln Met Ala Arg Ile 90 Lys Tyr	Glu Gln Pro 75 Ala Ser Trp	Vaide Control of the	u Gly Asp 45 1 Val Exist Met Exist Series 129 2 Series 2 Series 2 Series 3 Series 4 Series 4 Series 5 Series 6 Series 6 Series 7 Series 7 Series 7 Series 8 Series 7 Series 8 Series 8 Series 9 Series	7 Ile 30 Lys Lys L Ile Als Als 110 Lev S	15 Asi Phe Ile Ile Ile Ile Ile Ile Ile Ile Ile Il	n Pro
Met 65 Ala Thr Val	Ile 50 Asn Ile Leu Met Ala 130 Ser	Ile 35 Arg Asp Met Gln Thr 115 Leu Gln	20 Gly Glu Pro Asn Ile 100 Asp Val	Phe Lys Ser Pro 85 Phe Asp Thr	Ser Val Asn 70 Ala Asn Val Lys 150	Thr Gly 55 Pro Ser Ile Thr Asn 135 Met	Leu 40 Glu Ile Lys Glu Phe 120 Ala Phe	25 Thr Gln Arg Val Met 105 Trp Val Asp	Gln Met Ala Arg Ile 90 Lys Tyr Arg	Glu Gln Pro 75 Ala Ser Trp His	Vai 600 Ile Leu Ly: Ile Trp 140 Sec	uu Gly 45 45 Val 45 Val Lys Met 25 125 P Sei	7 Ile 30 Lyss Ala	15 Ası Phe	n Pro

Leu	Tyr		Val	Asp	Arg	Lys		Ser	Gln	Pro	Ile	Glu	Gly	His	Ala
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Gly	Ala 370	Glu	Glu	Leu	Phe	Ala 375	Arg	Lys	Phe	Asn	Ala 380	Leu	Phe	Ala	Gln
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Arg	Arg	Pro	Leu 980	Ile	Asp	Gln	Val	Val 985	Gln	Thr	Ala	Leu	Ser 990	Glu	Thr
Gln	Asp	Pro	Glu	Glu	Val	Ser	Val	Thi	r Val	Lys	s Ala	a Phe	∋ М∈	et Th	nr Ala

														~
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Ala	Ile 1070		Asn	Glu	Leu	Phe 1075		Glu	Ala	Phe	Ala 1080	Ile	Phe	Arg
Lys	Phe 1085	_	Val	Asn	Thr	Ser 1090		Val	Gln	Val	Leu 1095	Ile	Glu	His
Ile	Gly 1100		Leu	Asp	Arg	Ala 1105		Glu	Phe	Ala	Glu 1110		CAa	Asn
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Lys	Thr 1190		Arg	Leu	Ala	Glu 1195		Glu	Glu	Phe	Ile 1200	Asn	Gly	Pro
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Lys	Met 1220		Asp	Ala	Ala	Lys 1225		Leu	Tyr	Asn	Asn 1230	Val	Ser	Asn
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Ala	Leu 1490	Arg	Thr	Ser	Ile	Asp 1495	Ala	Tyr	Asp	Asn	Phe 1500	Asp	Asn	Ile
Ser	Leu 1505	Ala	Gln	Arg	Leu	Glu 1510	Lys	His	Glu	Leu	Ile 1515	Glu	Phe	Arg
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Сув	Leu 1580	Phe	Thr	Сув	Tyr	Asp 1585	Leu	Leu	Arg	Pro	Asp 1590	Val	Val	Leu
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Glu	Ser	Asp 35	Lys	Phe	Ile	Cys	Ile 40	Arg	Glu	Lys	Val	Gly 45	Glu	Gln	Ala
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Pro 65	Ile	Ser	Ala	Asp	Ser 70	Ala	Ile	Met	Asn	Pro 75	Ala	Ser	Lys	Val	Ile 80
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Ser	Lys	Met	Lys 100	Ala	His	Thr	Met	Thr 105	Asp	Asp	Val	Thr	Phe 110	Trp	Lys
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His	Trp 130	Ser	Met	Glu	Gly	Glu 135	Ser	Gln	Pro	Val	Lys 140	Met	Phe	Asp	Arg
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Glu	Lys 1010		e Val	l Leı	ı Asp	Asr 101		er V	al P	he S		lu 020	His	Arg	Asn
Leu	Gln 1025		ı Lev	ı Lev	ı Ile	Let 103		hr A	la I	le L		la 035	Asp	Arg	Thr
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Phe	Ala 1070		Phe	e Arg	J Lys	Phe 107		sp V	al A	sn Tl		er 080	Ala	Val	Gln
Val	Leu 1085		e Glu	ı His	; Ile	Gly 109		sn L	eu A	sp A:	_	la 095	Tyr	Glu	Phe
Ala	Glu 1100	_	g Cys	a Asr	ı Glu	110		la V	al T	rp S		ln 110	Leu	Ala	Lys
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Ile	Phe 1175		ı Let	ı Ala	a Lys	Th:		sn A	rg L	eu Ai		lu 185	Leu	Glu	Glu
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Pro	Asp	Val	Val	Leu	Glu	Thr	Ala	Trp	Arg	His	Asn	Ile	Met	Asp

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•		1580)				158	85				1	590			
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۲	/al	Ala 1655		l Pro	o Pro	o Glı	n Ala 160		ro Pl	he G	ly T		ly 665	Tyr	Thr	Ala
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1	Met L	Ala	Gln	Ile	Leu 5	Pro	Ile	Arg	Phe	Gln 10	Glu	His	Leu	Gln	Leu 15	Gln
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Pro	Gln	Val 595	Ala	Asp	Ala	Ile	Leu 600	Gly	Asn	Gln	Met	Phe 605	Thr	His	Tyr
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Leu	Gln 1025		ı Le	ı Lev	ı Ile	e Le:		nr Al	la I	le L	-	la 2 035	Asp A	Arg :	Thr
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	Leu (-			Pro P	Ala A	sn II 25	le G		ne Se	er Thi	Leu 30		Met
Glu				Phe I	Ile (Cys II	le Aı		lu Ly	ia As	al Gly 45		ı Glr	n Ala
Gln			Ile I	Ile A	_			sp Pi	ro Se	er As	en Pro	o Ile	e Arg	g Arg

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Cys	Glu	Lys 915	Arg	Asp	Pro	His	Leu 920	Alá	а Су	rs V	al A	la Ty: 92!		ı Arç	g Gly
Gln	Cys 930	Asp	Leu	Glu	Leu	Ile 935	Asn	Val	l Cy	rs A		lu Ası 40	n Sei	r Lei	ı Phe
Lys 945	Ser	Leu	Ser	Arg	Tyr 950	Leu	Val	Arg	g Ar		ys A 55	sp Pro	o Glu	ı Lev	ı Trp 960
Gly	Ser	Val	Leu	Leu 965	Glu	Ser	Asn	Pro	о Ту 97		rg A	rg Pro	Le:	ı Ile 975	
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Ser	Val	Thr 995	Val	ГÀа	Ala	Phe	Met 100		nr A	Ala	Asp :		ro <i>I</i> 005	Asn (Glu Leu
Ile	Glu 1010		ı Lev	ı Glu	ı Lys	101		al I	Leu	Asp	Asn	Ser 1020	Val	Phe	Ser
Glu	His 1025	-	J Asr	ı Lev	ı Gln	Asr 103		eu I	Leu	Ile	Leu	Thr 1035	Ala	Ile	Lys
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Tyr	Asp 1055		a Pro) Asp) Ile	106		sn I	Ile	Ala	Ile	Ser 1065	Asn	Glu	Leu
Phe	Glu 1070		ı Ala	e Phe	e Ala	11e 107		he A	Arg	Lys	Phe	Asp 1080	Val	Asn	Thr
Ser	Ala 1085		. Glr	ı Val	. Leu	11e 109		lu F	His	Ile	Gly	Asn 1095	Leu	Asp	Arg
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Gln	Leu 1115		ı Lys	a Ala	Gln	112		ln I	ŗya	Gly	Met	Val 1125	Lys	Glu	Ala
Ile	Asp 1130		туг	: Ile	e Lys	Ala 113		sp A	Aap	Pro	Ser	Ser 1140	Tyr	Met	Glu
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Lys	Leu 1220		і Туг	: Asr	ı Asn	Va]		er A	Asn	Phe	Gly	Arg 1230	Leu	Ala	Ser
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	1640)				164	15				16	550			
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Gly	Lys 1505	_	Glu	ı Cys	Phe	Ala 151		la C	ys.	Leu	Phe	Th:		Cys	Tyr	Asp
Leu	Leu 1520	_	Pro) Asp) Met	Va]		eu G	lu	Leu	Ala	15:	_	Arg	His	Asn
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Gly Ser His Ala Ala Pro Ala Gln Pro Gly Pro Thr Ser Gly Ala Gly 50 \, 55 \, 60 \,
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Ser	Ser	Pro 195	Gly	Thr	Glu	Trp	Glu 200	Arg	Val	Ala	Arg	Leu 205	CAa	Asp	Phe
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Ile 785	Pro	Asp	Ile	Asp	Pro 790	Asn	Ala	Lys	Leu	Leu 795	Gln	Pro	Pro	Ala	Pro 800
Ile	Met	Pro	Leu	Asp 805	Thr	Asn	Trp	Pro	Leu 810	Leu	Thr	Val	Ser	Lys 815	Gly
Phe	Phe	Glu	Gly 820	Thr	Ile	Ala	Ser	Lys 825	Gly	Lys	Gly	Gly	Ala 830	Leu	Ala
Ala	Asp	Ile 835	Asp	Ile	Asp	Thr	Val 840	Gly	Thr	Glu	Gly	Trp 845	Gly	Glu	Asp
Ala	Glu	Leu	Gln	Leu	Asp	Glu	Asp	Gly	Phe	Val	Glu	Ala	Thr	Glu	Gly

850		855	81	60
Leu Gly Asp A 865	Asp Ala Leu 870	Gly Lys Gl	y Gln Glu G: 875	lu Gly Gly Gly Trp 880
Asp Val Glu G	Slu Asp Leu 885	Glu Leu Pı	o Pro Glu Lo 890	eu Asp Ile Ser Pro 895
	Gly Gly Ala	Glu Asp Gl		al Pro Pro Thr Lys 910
Gly Thr Ser P	ro Thr Gln	Ile Trp Cy	s Asn Asn S	er Gln Leu Pro Val 925
Asp His Ile L 930		Ser Phe Gl 935		et Arg Leu Leu His 40
Asp Gln Val G 945	Sly Val Ile 950	Gln Phe Gl	y Pro Tyr Ly 955	ys Gln Leu Phe Leu 960
Gln Thr Tyr A	ala Arg Gly 965	Arg Thr Th	nr Tyr Gln Ai	la Leu Pro Cys Leu 975
	Tyr Gly Tyr 980	Pro Asn Ai		ys Asp Ala Gly Leu 990
Lys Asn Gly V 995	al Pro Ala	Val Gly I 1000	leu Lys Leu i	Asn Asp Leu Ile Gln 1005
Arg Leu Gln 1010	Leu Cys Tyr	Gln Leu 1015	Thr Thr Val	Gly Lys Phe Glu 1020
Glu Ala Val 1025	Glu Lys Phe	Arg Ser	Ile Leu Leu	Ser Val Pro Leu 1035
Leu Val Val 1040	Asp Asn Lys	Gln Glu 1045	Ile Ala Glu	Ala Gln Gln Leu 1050
Ile Thr Ile 1055	Cys Arg Glu	Tyr Ile	Val Gly Leu	Ser Val Glu Thr 1065
Glu Arg Lys 1070	Lys Leu Pro	Lys Glu 1075	Thr Leu Glu	Gln Gln Lys Arg 1080
Ile Cys Glu 1085	Met Ala Ala	Tyr Phe	Thr His Ser	Asn Leu Gln Pro 1095
Val His Met 1100	Ile Leu Val	Leu Arg	Thr Ala Leu	Asn Leu Phe Phe
Lys Leu Lys 1115	Asn Phe Lys	Thr Ala	Ala Thr Phe	Ala Arg Arg Leu 1125
Leu Glu Leu 1130	Gly Pro Lys	Pro Glu 1135	Val Ala Gln	Gln Thr Arg Lys 1140
Ile Leu Ser 1145	Ala Cys Glu	Lys Asn 1150	Pro Thr Asp	Ala Tyr Gln Leu 1155
Asn Tyr Asp	Met His Asn	Pro Phe	Asp Ile Cys	Ala Ala Ser Tyr 1170
Arg Pro Ile 1175	Tyr Arg Gly	Lys Pro	Val Glu Lys	Cys Pro Leu Ser 1185
Gly Ala Cys 1190	Tyr Ser Pro	Glu Phe	Lys Gly Gln	Ile Cys Arg Val 1200
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Ser Pro Leu 1220	Gln Phe Arg	ı		

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Ile	Gln	Leu 35	Trp	Asp	Tyr	Arg	Met 40	Cys	Thr	Leu	Ile	Asp 45	Lys	Phe	Asp
Glu	His 50	Asp	Gly	Pro	Val	Arg 55	Gly	Ile	Asp	Phe	His 60	Lys	Gln	Gln	Pro
Leu 65	Phe	Val	Ser	Gly	Gly 70	Asp	Asp	Tyr	Lys	Ile 75	Lys	Val	Trp	Asn	Tyr 80
Lys	Leu	Arg	Arg	Сув 85	Leu	Phe	Thr	Leu	Leu 90	Gly	His	Leu	Asp	Tyr 95	Ile
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CÀa	Val 130	Leu	Thr	Gly	His	Asn 135	His	Tyr	Val	Met	Cys 140	Ala	Gln	Phe	His
Pro 145	Thr	Glu	Asp	Leu	Val 150	Val	Ser	Ala	Ser	Leu 155	Asp	Gln	Thr	Val	Arg 160
Val	Trp	Asp	Ile	Ser 165	Gly	Leu	Arg	ГÀз	Lys 170	Asn	Leu	Ser	Pro	Gly 175	Ala
Val	Glu	Ser	Asp 180	Val	Arg	Gly	Ile	Thr 185	Gly	Val	Asp	Leu	Phe 190	Gly	Thr
Thr	Asp	Ala 195	Val	Val	Lys	His	Val 200	Leu	Glu	Gly	His	Asp 205	Arg	Gly	Val
Asn	Trp 210	Ala	Ala	Phe	His	Pro 215	Thr	Met	Pro	Leu	Ile 220	Val	Ser	Gly	Ala
Asp 225	Asp	Arg	Gln	Val	Lys 230	Ile	Trp	Arg	Met	Asn 235	Glu	Ser	Lys	Ala	Trp 240
Glu	Val	Asp	Thr	Сув 245	Arg	Gly	His	Tyr	Asn 250	Asn	Val	Ser	Cys	Ala 255	Val
Phe	His	Pro	Arg 260	Gln	Glu	Leu	Ile	Leu 265	Ser	Asn	Ser	Glu	Asp 270	ГЛЗ	Ser
Ile	Arg	Val 275	Trp	Asp	Met	Ser	Lys 280	Arg	Thr	Gly	Val	Gln 285	Thr	Phe	Arg
Arg	Asp 290	His	Asp	Arg	Phe	Trp 295	Val	Leu	Ala	Ala	His 300	Pro	Asn	Leu	Asn
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Arg	Glu	Arg	Pro	Ala 325	Tyr	Ala	Val	His	Gly 330	Asn	Met	Leu	His	Tyr 335	Val
Lys	Asp	Arg	Phe 340	Leu	Arg	Gln	Leu	Asp 345	Phe	Asn	Ser	Ser	Lys 350	Asp	Val

Ala	Val	Met 355	Gln	Leu	Arg	Ser	Gly 360	Ser	Lys	Phe	Pro	Val 365	Phe	Asn	Met
Ser	Tyr 370	Asn	Pro	Ala	Glu	Asn 375	Ala	Val	Leu	Leu	Cys 380	Thr	Arg	Ala	Ser
Asn 385	Leu	Glu	Asn	Ser	Thr 390	Tyr	Asp	Leu	Tyr	Thr 395	Ile	Pro	Lys	Asp	Ala 400
Asp	Ser	Gln	Asn	Pro 405	Asp	Ala	Pro	Glu	Gly 410	Lys	Arg	Ser	Ser	Gly 415	Leu
Thr	Ala	Val	Trp 420	Val	Ala	Arg	Asn	Arg 425	Phe	Ala	Val	Leu	Asp 430	Arg	Met
His	Ser	Leu 435	Leu	Ile	Lys	Asn	Leu 440	Lys	Asn	Glu	Ile	Thr 445	Lys	Lys	Val
Gln	Val 450	Pro	Asn	CÀa	Asp	Glu 455	Ile	Phe	Tyr	Ala	Gly 460	Thr	Gly	Asn	Leu
Leu 465	Leu	Arg	Asp	Ala	Asp 470	Ser	Ile	Thr	Leu	Phe 475	Asp	Val	Gln	Gln	Lys 480
Arg	Thr	Leu	Ala	Ser 485	Val	Lys	Ile	Ser	Lys 490	Val	Lys	Tyr	Val	Ile 495	Trp
Ser	Ala	Asp	Met 500	Ser	His	Val	Ala	Leu 505	Leu	Ala	Lys	His	Glu 510	His	Ser
Cys	Pro	Leu 515	Pro	Leu	Thr	Ala	Ile 520	Val	Ile	Сув	Asn	Arg 525	Lys	Leu	Asp
Ala	Leu 530	Сув	Asn	Ile	His	Glu 535	Asn	Ile	Arg	Val	Lys 540	Ser	Gly	Ala	Trp
Asp 545	Glu	Ser	Gly	Val	Phe 550	Ile	Tyr	Thr	Thr	Ser 555	Asn	His	Ile	Lys	Tyr 560
Ala	Val	Thr	Thr	Gly 565	Asp	His	Gly	Ile	Ile 570	Arg	Thr	Leu	Asp	Leu 575	Pro
Ile	Tyr	Val	Thr 580	Arg	Val	Lys	Gly	Asn 585	Asn	Val	Tyr	Cys	Leu 590	Asp	Arg
Glu	Сув	Arg 595	Pro	Arg	Val	Leu	Thr 600	Ile	Asp	Pro	Thr	Glu 605	Phe	Lys	Phe
ГÀа	Leu 610	Ala	Leu	Ile	Asn	Arg 615	Lys	Tyr	Asp	Glu	Val 620	Leu	His	Met	Val
Arg 625	Asn	Ala	ГÀа	Leu	Val 630	Gly	Gln	Ser	Ile	Ile 635	Ala	Tyr	Leu	Gln	Lys 640
Lys	Gly	Tyr	Pro	Glu 645	Val	Ala	Leu	His	Phe 650	Val	Lys	Asp	Glu	Lys 655	Thr
Arg	Phe	Ser	Leu 660	Ala	Leu	Glu	Cys	Gly 665	Asn	Ile	Glu	Ile	Ala 670	Leu	Glu
Ala	Ala	Lys 675	Ala	Leu	Asp	Asp	Lys	Asn	Càa	Trp	Glu	685 Lys	Leu	Gly	Glu
Val	Ala 690	Leu	Leu	Gln	Gly	Asn 695	His	Gln	Ile	Val	Glu 700	Met	Cys	Tyr	Gln
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Asn	Leu	Glu	Lys	Leu 725	Arg	Lys	Met	Met	Lys 730	Ile	Ala	Glu	Ile	Arg 735	Lys
Asp	Met	Ser	Gly 740	His	Tyr	Gln	Asn	Ala 745	Leu	Tyr	Leu	Gly	Asp 750	Val	Ser
Glu	Arg	Val	Arg	Ile	Leu	Lys	Asn	Cys	Gly	Gln	Lys	Ser	Leu	Ala	Tyr

		755					760					765	5		
Leu	Thr 770	Ala	Ala	Thr	His	Gly 775	Leu	Asp	Glı	ı Gl	u Al 78	a Glu O	ı Ser	Leu	ı Lys
Glu 785	Thr	Phe	Asp	Pro	Glu 790	Lys	Glu	Thr	: Ile	e Pr 79		p Ile	e Asp	Pro	Asn 800
Ala	Lys	Leu	Leu	Gln 805	Pro	Pro	Ala	Pro	810		t Pr	o Leu	ı Asp	Thr 815	
Trp	Pro	Leu	Leu 820	Thr	Val	Ser	Lys	Gly 825		e Ph	e Gl	u Gly	7 Thr 830		e Ala
Ser	ГХа	Gly 835	Lys	Gly	Gly	Ala	Leu 840	Ala	ı Ala	a As	p Il	e Asp 845		Asp	Thr
Val	Gly 850	Thr	Glu	Gly	Trp	Gly 855	Glu	Asp	Ala	a Gl	u Le 86	u Glr 0	Leu	Asp	Glu
Asp 865	Gly	Phe	Val	Glu	Ala 870	Thr	Glu	Gly	Lei	ı Gl 87		p Asp	Ala	Leu	1 Gly 880
Lys	Gly	Gln	Glu	Glu 885	Gly	Gly	Gly	Trp	890		1 G1	u Glu	ı Asp	Leu 895	
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Asp	Gly	Phe 915	Phe	Val	Pro	Pro	Thr 920	Lys	Gly	y Th	ır Se	r Pro		Gln	ı Ile
Trp	Cys 930	Asn	Asn	Ser	Gln	Leu 935	Pro	Val	. Asj) Hi	s Il 94	e Leu 0	ı Ala	Gly	/ Ser
Phe 945	Glu	Thr	Ala	Met	Arg 950	Leu	Leu	His	. Asl	95		l Gly	v Val	. Il∈	960
Phe	Gly	Pro	Tyr	Lys 965	Gln	Leu	Phe	Leu	970		ır Ty	r Ala	a Arg	Gly 975	_
Thr	Thr	Tyr	Gln 980	Ala	Leu	Pro	Cys	Leu 985		Se	r Me	t Tyr	Gly 990		Pro
Asn	Arg	Asn 995	Trp	Lys	Asp	Ala	Gly 100		eu Ly	ys A	sn G	ly Va	1 F	ro A	Ala Val
Gly	Leu 1010		. Leu	ı Ası	n Asp	Le:		le G	ln A	Arg	Leu	Gln 1020	Leu	Cys	Tyr
Gln	Leu 1025		Thi	r Val	L Gly	Ly:		he G	lu (Glu	Ala	Val 1035	Glu	Lys	Phe
Arg	Ser 1040		e Let	ı Lev	ı Ser	Va:		ro I	eu 1	Leu	Val	Val 1050	Asp	Asn	Lys
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Tyr	Ile 1070		l Gly	/ Let	ı Ser	Va:		lu T	hr (Glu		Lys 1080	Lys	Leu	Pro
ГÀв	Glu 1085		: Let	ı Glı	ı Glr	109		ys A	rg :	Ile		Glu 1095	Met	Ala	Ala
Tyr	Phe 1100		: His	s Sei	Asr	Le:		ln F	ro 7	/al		Met 1110	Ile	Leu	Val
Leu	Arg 1115		Ala	a Lei	ı Asr	Let 112		he F	he l	ŗÀa		Lys 1125	Asn	Phe	ГÀа
Thr	Ala 1130		a Thi	r Phe	e Ala	Arç	_	rg L	eu 1	Leu		Leu 1140	Gly	Pro	Lys
Pro	Glu 1145		l Ala	a Glr	n Glr	1 Th:		rg L	iys :	Ile		Ser 1155	Ala	Cys	Glu

Lys	Asn 1160		Thr	a Asp	Ala	Ту1 116		ln Le	eu As	n Ty		sp 170	Met 1	His 1	Asn
Pro	Phe 1175	_) Ile	e Cys	. Ala	. Ala		er T∑	r Ai	g Pı		le 185	Tyr 1	Arg (Gly
Lys	Pro 1190		. Glu	ı Lys	cys	Pro		eu Se	er Gl	Ly Al		ys 200	Tyr :	Ser :	Pro
Glu	Phe 1205		Gly	/ Glr	ılle	Cys 121		rg Va	al Th	nr Th		al 215	Thr	Glu :	Ile
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Gly	Asp	Val 35	Lys	Ser	ГÀв	Thr	Glu 40	Ala	Leu	Lys	ГÀз	Val 45	Ile	Ile	Met
Ile	Leu 50	Asn	Gly	Glu	-	Leu 55	Pro	Gly	Leu	Leu	Met 60	Thr	Ile	Ile	Arg
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Glu	Ala 130	Glu	Leu	Leu		Pro 135	Leu	Met	Pro	Ala	Ile 140	Arg	Ala	Cys	Leu
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Thr	Ile	Tyr	Arg	Asn 165	Phe	Glu	His	Leu	Ile 170	Pro	Asp	Ala	Pro	Glu 175	Leu
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Val	Glu	Glu 355	Leu	Val	Ile	Val	Leu 360	Lys	Lys	Glu	Val	Ile 365	ГÀа	Thr	Asn
Asn	Val 370	Ser	Glu	His	Glu	Asp 375	Thr	Asp	Lys	Tyr	Arg 380	Gln	Leu	Leu	Val
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Val	Ile	Pro	Val	Leu 405	Met	Glu	Phe	Leu	Ser 410	Asp	Asn	Asn	Glu	Ala 415	Ala
Ala	Ala	Asp	Val 420	Leu	Glu	Phe	Val	Arg 425	Glu	Ala	Ile	Gln	Arg 430	Phe	Asp
Asn	Leu	Arg 435	Met	Leu	Ile	Val	Glu 440	Lys	Met	Leu	Glu	Val 445	Phe	His	Ala
Ile	Lys 450	Ser	Val	Lys	Ile	Tyr 455	Arg	Gly	Ala	Leu	Trp 460	Ile	Leu	Gly	Glu
Tyr 465	Cys	Ser	Thr	Lys	Glu 470	Asp	Ile	Gln	Ser	Val 475	Met	Thr	Glu	Ile	Arg 480
Arg	Ser	Leu	Gly	Glu 485	Ile	Pro	Ile	Val	Glu 490	Ser	Glu	Ile	Lys	Lys 495	Glu
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Ser	Ser 530	Arg	Pro	Thr	rys	Lys 535	Glu	Glu	Asp	Arg	Pro 540	Pro	Leu	Arg	Gly
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ГÀв	Gln	Asn	Ser 580	Phe	Val	Ala	Glu	Ala 585	Met	Leu	Leu	Met	Ala 590	Thr	Ile
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Asp	Val 610	Asp	Arg	Ile	Ser	Leu 615	CAa	Leu	Lys	Val	Leu 620	Ser	Glu	CÀa	Ser
Pro 625	Leu	Met	Asn	Asp	Ile 630	Phe	Asn	Lys	Glu	Сув 635	Arg	Gln	Ser	Leu	Ser 640
His	Met	Leu	Ser	Ala 645	Lys	Leu	Glu	Glu	Glu 650	Lys	Leu	Ser	Gln	Lys 655	ГЛа
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Phe Met	Gln Leu 675	. Thr Ala	Lys	Asn 680	Glu	Met	Asn	CAa	685 Lys	Glu	Asp	Gln
Phe Gln :	Leu Ser	Leu Leu	Ala 695	Ala	Met	Gly	Asn	Thr 700	Gln	Arg	Lys	Glu
Ala Ala . 705	Asp Pro	Leu Ala 710		Lys	Leu	Asn	Lys 715	Val	Thr	Gln	Leu	Thr 720
Gly Phe	Ser Asp	Pro Val 725	Tyr	Ala	Glu	Ala 730	Tyr	Val	His	Val	Asn 735	Gln
Tyr Asp	Ile Val 740	_	Val	Leu	Val 745	Val	Asn	Gln	Thr	Ser 750	Asp	Thr
Leu Gln .	Asn Cys 755	Thr Lev	Glu	Leu 760	Ala	Thr	Leu	Gly	Asp 765	Leu	Lys	Leu
Val Glu 770	Lys Pro	Ser Pro	Leu 775	Thr	Leu	Ala	Pro	His 780	Asp	Phe	Ala	Asn
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Gly Asn	Ile Val	Tyr Asp 805	Val	Ser	Gly	Ala 810	Ala	Ser	Asp	Arg	Asn 815	CÀa
Val Val	Leu Ser 820		His	Ile	Asp 825	Ile	Met	Asp	Tyr	Ile 830	Gln	Pro
Ala Thr	Cys Thr 835	Asp Ala	Glu	Phe 840	Arg	Gln	Met	Trp	Ala 845	Glu	Phe	Glu
Trp Glu . 850	Asn Lys	Val Thr	Val 855	Asn	Thr	Asn	Met	Val 860	Asp	Leu	Asn	Asp
Tyr Leu 865	Gln His	Ile Leu 870	_	Ser	Thr	Asn	Met 875	Lys	Сув	Leu	Thr	Pro 880
Glu Lys .	Ala Leu	Ser Gly 885	Tyr	CÀa	Gly	Phe 890	Met	Ala	Ala	Asn	Leu 895	Tyr
Ala Arg	Ser Ile 900		Glu	Asp	Ala 905	Leu	Ala	Asn	Val	Ser 910	Ile	Glu
Lys Pro	Ile His 915	Gln Gly	Pro	Asp 920	Ala	Ala	Val	Thr	Gly 925	His	Ile	Arg
Ile Arg . 930	Ala Lys	Ser Glr	Gly 935	Met	Ala	Leu	Ser	Leu 940	Gly	Asp	Lys	Ile
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Ser Leu	Tyr Asn 35	Gly Ser	Val	Cys 40	Val	Trp	Asn	His	Glu 45	Thr	Gln	Thr

Leu	Val 50	Lys	Thr	Phe	Glu	Val 55	СЛа	Asp	Leu	Pro	Val 60	Arg	Ala	Ala	Lys
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Asp	Lys 130	ГÀа	Trp	Ser	CAa	Ser 135	Gln	Val	Phe	Glu	Gly 140	His	Thr	His	Tyr
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Ala	Ser	Leu	Asp	Arg 165	Thr	Ile	Lys	Val	Trp 170	Gln	Leu	Gly	Ser	Ser 175	Ser
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Pro Va	l Ile 835	Val	Ala	Ser	His	Thr 840	Ala	Asn	Lys	Glu	Glu 845	Lys	Ser	Leu
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Ile	Gln	Val	Thr	Lys 485	Val	Thr	Gln	Val	Asp 490	Gly	Asn	Ser	Pro	Val 495	Arg
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Thr	Ile 290	Met	Ser	Ser	Ser	Met 295	Gly	ГЛа	Arg	Thr	Ser 300	Glu	Ala	Thr	Lys
Met 305	His	Ala	Pro	Pro	Ile 310	Asn	Met	Glu	Ser	Val 315	His	Met	Lys	Ile	Glu 320
Glu	Lys	Ile	Thr	Leu 325	Thr	Cys	Gly	Arg	330	Gly	Gly	Leu	Gln	Asn 335	Met
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Arg	Ile	Arg 355	Leu	His	Val	Glu	Asn 360	Glu	Asp	Lys	Lys	Gly 365	Val	Gln	Leu
Gln	Thr 370	His	Pro	Asn	Val	Asp 375	Lys	Lys	Leu	Phe	Thr 380	Ala	Glu	Ser	Leu

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Leu	Thr	Ile	Asn 420	Cys	Trp	Pro	Ser	Glu 425	Ser	Gly	Asn	Gly	Cys 430	Asp	Val
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Glu 465	Ile	Asp	Gly	Glu	Tyr 470	Arg	His	Asp	Ser	Arg 475	Arg	Asn	Thr	Leu	Glu 480
Trp	Cys	Leu	Pro	Val 485	Ile	Asp	Ala	Lys	Asn 490	Lys	Ser	Gly	Ser	Leu 495	Glu
Phe	Ser	Ile	Ala 500	Gly	Gln	Pro	Asn	Asp 505	Phe	Phe	Pro	Val	Gln 510	Val	Ser
Phe	Val	Ser 515	Lys	ГЛа	Asn	Tyr	Cys 520	Asn	Ile	Gln	Val	Thr 525	ГЛа	Val	Thr
Gln	Val 530	Asp	Gly	Asn	Ser	Pro 535	Val	Arg	Phe	Ser	Thr 540	Glu	Thr	Thr	Phe
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Glu	Asp	Ala	Thr 180	Leu	Thr	Gln	Leu	Ala 185	Thr	Ala	Trp	Val	Ser 190	Leu	Ala
Thr	Gly	Gly 195	Glu	Lys	Leu	Gln	Asp 200	Ala	Tyr	Tyr	Ile	Phe 205	Gln	Glu	Met
Ala	Asp 210	Lys	Сув	Ser	Pro	Thr 215	Leu	Leu	Leu	Leu	Asn 220	Gly	Gln	Ala	Ala
Cys 225	His	Met	Ala	Gln	Gly 230	Arg	Trp	Glu	Ala	Ala 235	Glu	Gly	Leu	Leu	Gln 240
Glu	Ala	Leu	Asp	Lys 245	Asp	Ser	Gly	Tyr	Pro 250	Glu	Thr	Leu	Val	Asn 255	Leu
Ile	Val	Leu	Ser 260	Gln	His	Leu	Gly	Lys 265	Pro	Pro	Glu	Val	Thr 270	Asn	Arg
Tyr	Leu	Ser 275	Gln	Leu	ГЛа	Asp	Ala 280	His	Arg	Ser	His	Pro 285	Phe	Ile	Lys
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Gln	Сув	Ile 35	Asn	Glu	Ala	Gln	Arg 40	Val	Lys	Leu	Ser	Ser 45	Pro	Glu	Arg
Asp	Val 50	Glu	Arg	Asp	Val	Phe 55	Leu	Tyr	Arg	Ala	Tyr 60	Leu	Ala	Gln	Arg
Lys 65	Phe	Gly	Val	Val	Leu 70	Asp	Glu	Ile	Lys	Pro 75	Ser	Ser	Ala	Pro	Glu 80
Leu	Gln	Ala	Val	Arg 85	Met	Phe	Ala	Asp	Tyr 90	Leu	Ala	His	Glu	Ser 95	Arg
Arg	Asp	Ser	Ile 100	Val	Ala	Glu	Leu	Asp 105	Arg	Glu	Met	Ser	Arg 110	Ser	Val
Asp	Val	Thr 115	Asn	Thr	Thr	Phe	Leu 120	Leu	Met	Ala	Ala	Ser 125	Ile	Tyr	Leu
His	Asp 130	Gln	Asn	Pro	Asp	Ala 135	Ala	Leu	Arg	Ala	Leu 140	His	Gln	Gly	Asp
Ser 145	Leu	Glu	Cys	Thr	Ala 150	Met	Thr	Val	Gln	Ile 155	Leu	Leu	Lys	Leu	Asp 160
Arg	Leu	Asp	Leu	Ala 165	Arg	Lys	Glu	Leu	Lys 170	Arg	Met	Gln	Asp	Leu 175	Asp
Glu	Asp	Ala	Thr 180	Leu	Thr	Gln	Leu	Ala 185	Thr	Ala	Trp	Val	Ser 190	Leu	Ala

Thr Asp Ser Gly Tyr Pro Glu Thr Leu Val Asn Leu Ile Val Leu Ser 200 Gln His Leu Gly Lys Pro Pro Glu Val Thr Asn Arg Tyr Leu Ser Gln Leu Lys Asp Ala His Arg Ser His Pro Phe Ile Lys Glu Tyr Gln Ala Lys Glu Asn Asp Phe Asp Arg Leu Val Leu Gln Tyr Ala Pro Ser Ala <210> SEQ ID NO 25 <211> LENGTH: 257 <212> TYPE: PRT <213 > ORGANISM: Homo sapiens <300> PUBLICATION INFORMATION: <308> DATABASE ACCESSION NUMBER: NCBI/NP_955474 <309> DATABASE ENTRY DATE: 2008-09-28 <313> RELEVANT RESIDUES IN SEQ ID NO: (1)..(257) <400> SEOUENCE: 25 Met Ala Pro Pro Ala Pro Gly Pro Ala Ser Gly Gly Ser Gly Glu Val Asp Glu Leu Phe Asp Val Lys Asn Ala Phe Tyr Ile Gly Ser Tyr Gln Gln Cys Ile Asn Glu Ala Gln Arg Val Lys Leu Ser Ser Pro Glu Arg Asp Val Glu Arg Asp Val Phe Leu Tyr Arg Ala Tyr Leu Ala Gln Arg Lys Phe Gly Val Val Leu Asp Glu Ile Lys Pro Ser Ser Ala Pro Glu Leu Gln Ala Val Arg Met Phe Ala Asp Tyr Leu Ala His Glu Ser Arg Ser Thr Ala Met Thr Val Gln Ile Leu Leu Lys Leu Asp Arg Leu Asp 105 Leu Ala Arg Lys Glu Leu Lys Arg Met Gln Asp Leu Asp Glu Asp Ala Thr Leu Thr Gln Leu Ala Thr Ala Trp Val Ser Leu Ala Thr Gly Gly Glu Lys Leu Gln Asp Ala Tyr Tyr Ile Phe Gln Glu Met Ala Asp Lys Cys Ser Pro Thr Leu Leu Leu Asn Gly Gln Ala Ala Cys His Met Ala Gl
n Gly Arg Trp Glu Ala Ala Glu Gly Leu Leu Gl
n Glu Ala Leu 180 185 190 Asp Lys Asp Ser Gly Tyr Pro Glu Thr Leu Val Asn Leu Ile Val Leu 200 Ser Gln His Leu Gly Lys Pro Pro Glu Val Thr Asn Arg Tyr Leu Ser Gln Leu Lys Asp Ala His Arg Ser His Pro Phe Ile Lys Glu Tyr Gln Ala Lys Glu Asn Asp Phe Asp Arg Leu Val Leu Gln Tyr Ala Pro Ser 250

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			20					25					30		
Val	Phe	Asn	Glu	Thr	Pro	Ile	Asn	Pro	Arq	Lys	Cys	Ala	His	Ile	Leu
		35					40		J	-	-	45			
Thr		Ile	Leu	Tyr	Leu		Asn	Gln	Gly	Glu		Leu	Gly	Thr	Thr
	50					55					60				
Glu	Ala	Thr	Glu	Ala	Phe	Phe	Ala	Met	Thr	Lvs	Leu	Phe	Gln	Ser	Asn
65					70					75					80
Asp	Pro	Thr	Leu	_	Arg	Met	CAa	Tyr		Thr	Ile	Lys	Glu		Ser
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Cvs	Ile	Ala	Glu	Asp	Val	Ile	Ile	Val	Thr	Ser	Ser	Leu	Thr	Lvs	Asp
- 2			100					105					110	-2	
Met	Thr		Lys	Glu	Asp	Asn		Arg	Gly	Pro	Ala		Arg	Ala	Leu
		115					120					125			
Cvs	Gln	Tle	Thr	Agn	Ser	Thr	Met	Len	Gln	Δla	Tle	Glu	Ara	Tur	Met
СУБ	130	110	1111	лър	DCI	135	ricc	пса	GIII	AIG	140	Ola	AI 9	1 y 1	ricc
_	Gln	Ala	Ile	Val	Asp	Lys	Val	Pro	Ser	Val	Ser	Ser	Ser	Ala	Leu
145					150					155					160
Wal.	Cor	Cor	Lou	ціа	Lou	Lou	Lva	Crra	Cor	Dho	7 cm	Wal.	Val	Lva	Ara
vai	SeT	ser	пец	165	Leu	пец	цув	Сув	170	FILE	Asp	vai	vai	175	AIG
${\tt Trp}$	Val	Asn		Ala	Gln	Glu	Ala	Ala	Ser	Ser	Asp	Asn	Ile	Met	Val
			180					185					190		
C12	Trees	III a	7.7.	T 011	C1	T 011	T 011	Tr + 20	IIi a	7707	7. 20.00	T	7.00	70 0000	7.200
GIII	тут	195	Ата	пец	Gly	пец	200	тут	птъ	vai	AIG	205	Abii	Asp	AIG
Leu		Val	Asn	Lys	Met		Ser	Lys	Val	Thr	Arg	His	Gly	Leu	Lys
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Ser	Pro	Dhe	Δla	Tur	Cys	Met	Met	Tle	Δra	Val	Δla	Ser	Lare	Gln	T.e.11
225	FIO	FILE	Ата	ıyı	230	Mec	Nec	116	Arg	235	AIA	Ser	шуъ	GIII	240
Glu	Glu	Glu	Asp		Ser	Arg	Asp	Ser		Leu	Phe	Asp	Phe	Ile	Glu
				245					250					255	
Car	Carc	T. 211	Δrα	Λan	Lys	шіс	Glu	Mat	17 a 1	T/all	Tarr	Glu	Λla	Λla	Car
Ser	СуБ	пец	260	Abii	цуь	шта	GIU	265	vai	vai	тут	GIU	270	лта	Det
Ala	Ile	Val	Asn	Leu	Pro	Gly	Cys	Ser	Ala	Lys	Glu	Leu	Ala	Pro	Ala
		275					280					285			
Val		Val	Leu	Gln	Leu		CAa	Ser	Ser	Pro	_	Ala	Ala	Leu	Arg
	290					295					300				
Tarr	Δla	Δla	Val	Δra	Thr	T 11	Δan	Lare	Val	Δla	Met	Lare	Hie	Pro	Ser
305	- ALG	n L d	val	y	310	Leu	11011	-ya	Val	315	1150	Lys	1113	110	320
Ala	Val	Thr	Ala	Cys	Asn	Leu	Asp	Leu	Glu	Asn	Leu	Val	Thr	Asp	Ser
				325					330					335	
Asn	Arg	Ser		Ala	Thr	Leu	Ala		Thr	Thr	Leu	Leu		Thr	Gly
			340					345					350		

Ser	Glu	Ser 355	Ser	Ile	Asp	Arg	Leu 360	Met	Lys	Gln	Ile	Ser 365	Ser	Phe	Met
Ser	Glu 370	Ile	Ser	Asp	Glu	Phe 375	Lys	Val	Val	Val	Val 380	Gln	Ala	Ile	Ser
Ala 385	Leu	CAa	Gln	ГЛа	Tyr 390	Pro	Arg	Lys	His	Ala 395	Val	Leu	Met	Asn	Phe 400
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Thr	Val 450	Leu	Ala	Thr	Arg	Ile 455	Leu	His	Leu	Leu	Gly 460	Gln	Glu	Gly	Pro
Lys 465	Thr	Thr	Asn	Pro	Ser 470	Lys	Tyr	Ile	Arg	Phe 475	Ile	Tyr	Asn	Arg	Val 480
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Lys	Phe	Gly	Ala 500	Gln	Asn	Glu	Glu	Met 505	Leu	Pro	Ser	Ile	Leu 510	Val	Leu
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Thr	Phe 530	Tyr	Leu	Asn	Val	Leu 535	Glu	Gln	Lys	Gln	Lys 540	Ala	Leu	Asn	Ala
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Lys	Ser	Val	Pro 580	Leu	Ala	Thr	Ala	Pro 585	Met	Ala	Glu	Gln	Arg 590	Thr	Glu
Ser	Thr	Pro 595	Ile	Thr	Ala	Val	Lys	Gln	Pro	Glu	Lys	Val 605	Ala	Ala	Thr
Arg	Gln 610	Glu	Ile	Phe	Gln	Glu 615	Gln	Leu	Ala	Ala	Val 620	Pro	Glu	Phe	Arg
Gly 625	Leu	Gly	Pro	Leu	Phe 630	Lys	Ser	Ser	Pro	Glu 635	Pro	Val	Ala	Leu	Thr 640
Glu	Ser	Glu	Thr	Glu 645	Tyr	Val	Ile	Arg	Сув 650	Thr	Lys	His	Thr	Phe 655	Thr
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Ala	Сув	Thr	Phe	Ser 725	Сув	Met	Met	Lys	Phe	Thr	Val	Lys	Asp	Cys 735	Asp
Pro	Thr	Thr	Gly 740	Glu	Thr	Asp	Asp	Glu 745	Gly	Tyr	Glu	Asp	Glu 750	Tyr	Val

Leu	Glu	Asp 755	Leu	Glu	Val	Thr	Val 760	Ala	Asp	His	Ile	Gln 765	Lys	Val	Met
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Asp	ГЛа	Val	Pro 820	Asp	Asn	Lys	Asn	Thr 825	His	Thr	Leu	Leu	Leu 830	Ala	Gly
Val	Phe	Arg 835	Gly	Gly	His	Asp	Ile 840	Leu	Val	Arg	Ser	Arg 845	Leu	Leu	Leu
Leu	Asp 850	Thr	Val	Thr	Met	Gln 855	Val	Thr	Ala	Arg	Ser 860	Leu	Glu	Glu	Leu
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Thr	Lуs 50	Ile	Leu	Tyr	Leu	Leu 55	Asn	Gln	Gly	Glu	His 60	Phe	Gly	Thr	Thr
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195 200 205

Leu Lys
210

In view of the foregoing, what is claimed is:

- 1. A pharmaceutical composition for replacement therapy comprising: a man-made non-cage clathrin heavy chain protein that has an amino acid sequence of at least 50, 60, 70, 80, or 90% sequence identity when compared to SEQ ID NO: 1, and administering to a subject in need thereof the preceding protein or composition, and the protein or composition is taken up in whole or in part by a living cell in vivo or in vitro, wherein the protein or composition replaces and/or modifies at least one of cell element or process, and treats a disease, condition, or disorder comprising at least one of cell.
- 2. The pharmaceutical composition of claim 1, comprising a clathrin light chain protein that is bound to the man-made protein.
- 3. The pharmaceutical composition of claim 1, comprising one or more cargo elements that are bound to the man-made protein.
- **4**. The pharmaceutical composition of claim **1**, comprising one or more additional elements that are incorporated into the man-made protein.
- **5**. The pharmaceutical composition of claim **1**, comprising tethers that capture and bind one or more elements to the man-made protein.
- **6**. The pharmaceutical composition of claim **1**, comprising receptors that capture and position one or more elements on the man-made protein.
- 7. The pharmaceutical composition of claim 1, comprising adaptor proteins that are affixed to the man-made protein.
- **8**. The pharmaceutical composition of claim **7**, wherein the adaptor proteins are disposed between receptors and the man-made clathrin protein and binding to the receptors.
- **9**. The pharmaceutical composition of claim **1**, wherein the one or more cell element or process is located external or internal to cell.
- 10. The pharmaceutical composition of claim 1, wherein the one or more cell element or process is captured and/or positioned external or internal to cell by the composition or man-made protein.
- 11. The pharmaceutical composition of claim 1, wherein the one or more cell element or process is replaced and/or modified by the composition or man-made protein.
- 12. The pharmaceutical composition of claim 1, wherein the man-made protein comprises a multiple of the man-made protein.
- 13. The pharmaceutical composition of claim 1, wherein the one or more cell comprises a multiple of living cells.
- 14. The pharmaceutical composition of claim 1, wherein the man-made protein comprises biologically active, biologically modified, chiral, genetically modified, hybridized, isolated, chemically modified, masked, recombinant, synthetic or unmasked protein molecules, or a combination thereof.

- **15**. The pharmaceutical composition of claim **1**, wherein the man-made protein has a molecular mass of at least about 170 kilodaltons.
- **16**. The pharmaceutical composition of claim **1**, wherein the man-made protein is stable with respect to dissociation.
- 17. The pharmaceutical composition of claim 1, wherein the pharmaceutical composition is formulated for any suitable route and means of administration.
- 18. The pharmaceutical composition of claim 17, wherein the pharmaceutical composition is formulated as an adhesive, aerosols, a biologic, a capsule, a chemical compound, a coated, crystals, eye drops, gels, an injectable, liquids, oils, ointments, a polymer, powders, a prosthetic, salves, soft galantine capsules, a stent, a controlled release, subcutaneous, surgical, syrups, a tablet, a topical, a vapor, or a water soluble formula.
- 19. The pharmaceutical composition of claim 1, further comprising a therapeutic agent.
- 20. The pharmaceutical composition of claim 19, wherein the therapeutic agent is a drug for treating a disease, condition or disorder comprising at least one of cell.
- 21. The pharmaceutical composition of claim 1, wherein the pharmaceutical composition is formulated as an element for use with another element, and one or both of the preceding elements used as a drug for treating a disease, condition, or disorder comprising at least one of cell.
- 22. The pharmaceutical composition of claim 1, wherein the pharmaceutical composition is formulated for targeting a disease, condition, or disorder comprising at least one of cell.
- 23. The pharmaceutical composition of claim 1, wherein the pharmaceutical composition is formulated with acceptable masking agents for reducing immunogenicity and antigenicity.
- **24**. The pharmaceutical composition of claim **1**, wherein SEQ ID NO: 1 is substituted with an alternative protein sequence generated from the same gene that generated SEQ ID NO: 1.
- **25**. The pharmaceutical composition of claim **24**, wherein the alternative protein sequence includes but not limited to, SEQ ID NO: 2, SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO: 5, SEQ ID NO: 6, SEQ ID NO: 7 SEQ ID NO: 8, SEQ ID NO: 9, SEQ ID NO: 10 or SEQ ID NO: 11.
- 26. A method for a pharmaceutical composition for replacement therapy comprising: forming pharmaceutical composition for replacement therapy comprising: a manmade non-cage clathrin heavy chain protein that has an amino acid sequence of at least 50, 60, 70, 80, or 90% sequence identity when compared to SEQ ID NO: 1, and administering to a subject in need thereof the preceding protein or composition, and the protein or composition is taken up in whole or in part by a living cell in vivo or in vitro, wherein the protein or composition replaces and/or

modifies at least one of cell element or process, and treats a disease, condition, or disorder comprising at least one of cell.

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