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(54) **HEAT-INDUCIBLE SELF-ASSEMBLING PROTEIN DOMAINS**

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(71) Applicant: **The University of Chicago**, Chicago, IL (US)

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(72) Inventors: **D. Allan Drummond**, Chicago, IL (US); **Joshua Riback**, Chicago, IL (US); **Jamie Scott**, Chicago, IL (US); **Alexandra Rojek**, Chicago, IL (US); **Pawel Laskowski**, Chicago, IL (US); **Ronald Rock**, Chicago, IL (US); **Jagoda Rokicka**, Chicago, IL (US); **Jakub Kucharczyk**, Chicago, IL (US)

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(57) **ABSTRACT**

(22) PCT Filed: **Oct. 23, 2015**

A heat-inducible self-assembling fusion protein that includes a self-assembly domain and a target protein, wherein the self-assembly domain remains folded during assembly. The aggregate forming fusion protein can be induced to form protein aggregates conjugated to a target protein. The aggregates can be used similarly to beads in many laboratory protocols and other applications. Also disclosed are methods of making and using the protein aggregates.

(86) PCT No.: **PCT/US15/57139**

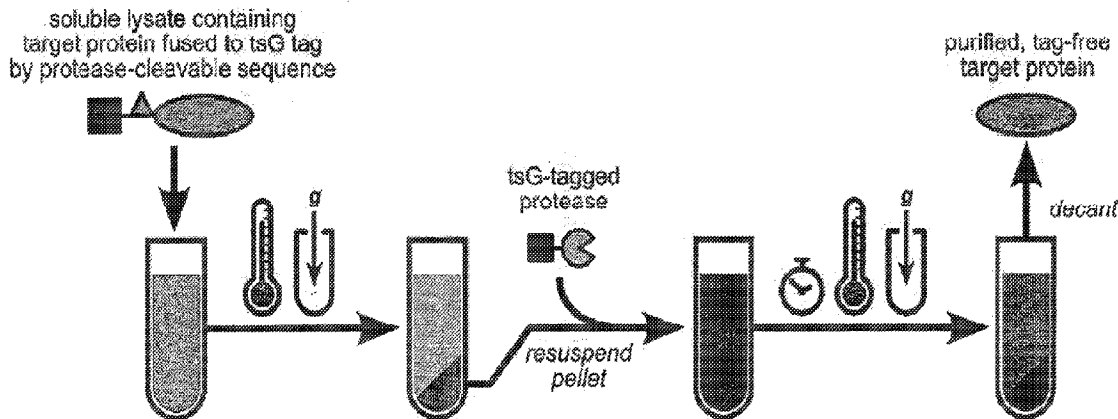
§ 371 (c)(1),

(2) Date: **Apr. 24, 2017**

Related U.S. Application Data

(60) Provisional application No. 62/068,390, filed on Oct. 24, 2014.

Specification includes a Sequence Listing.



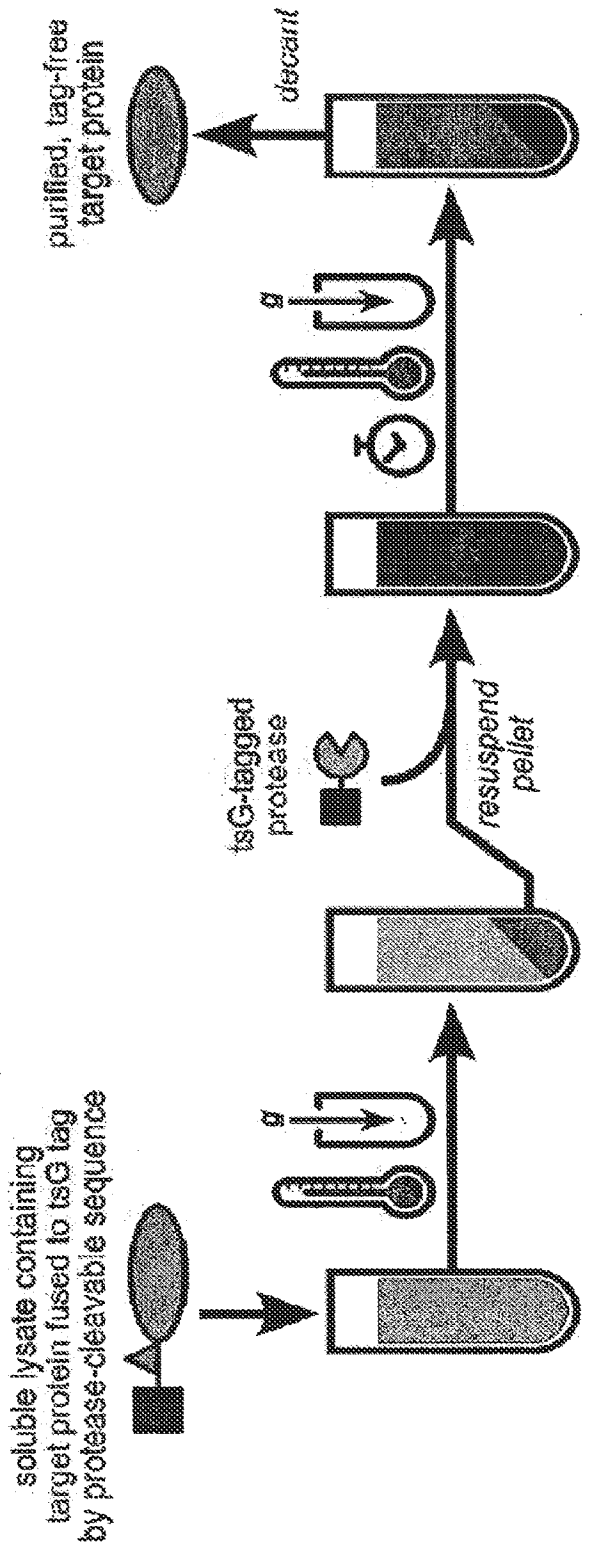


FIG. 1

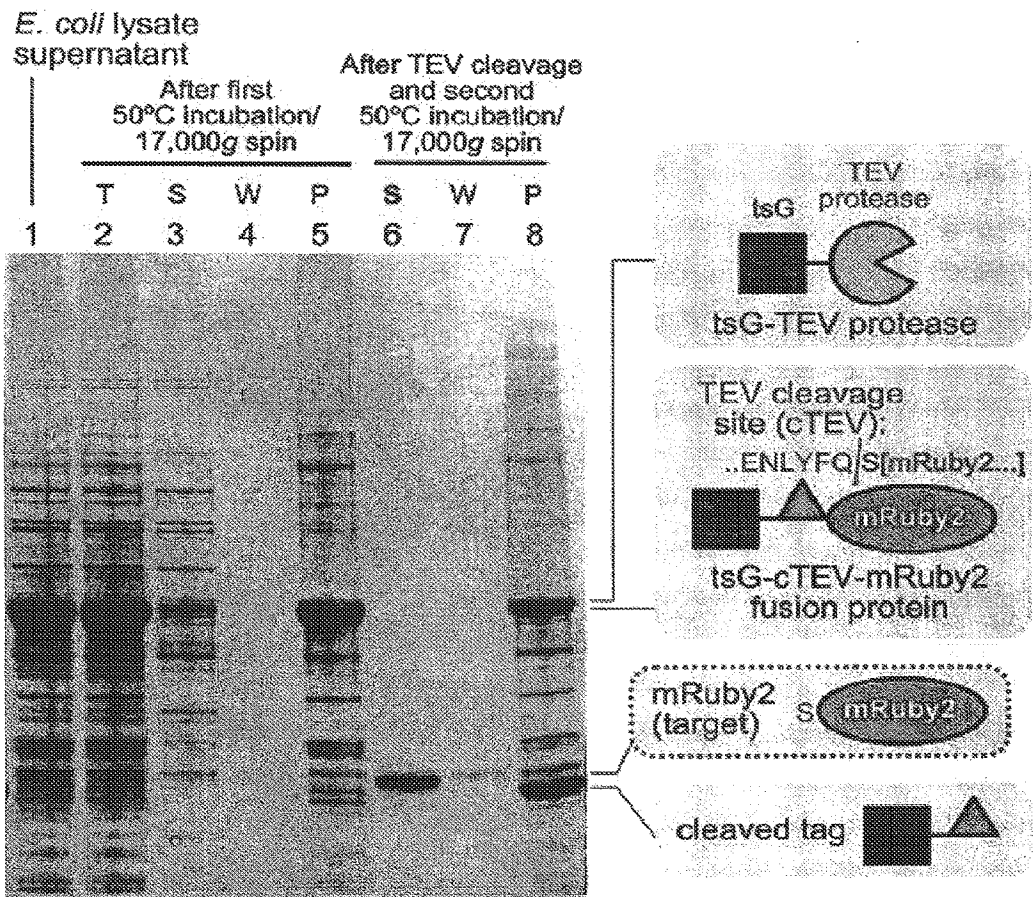


FIG. 2

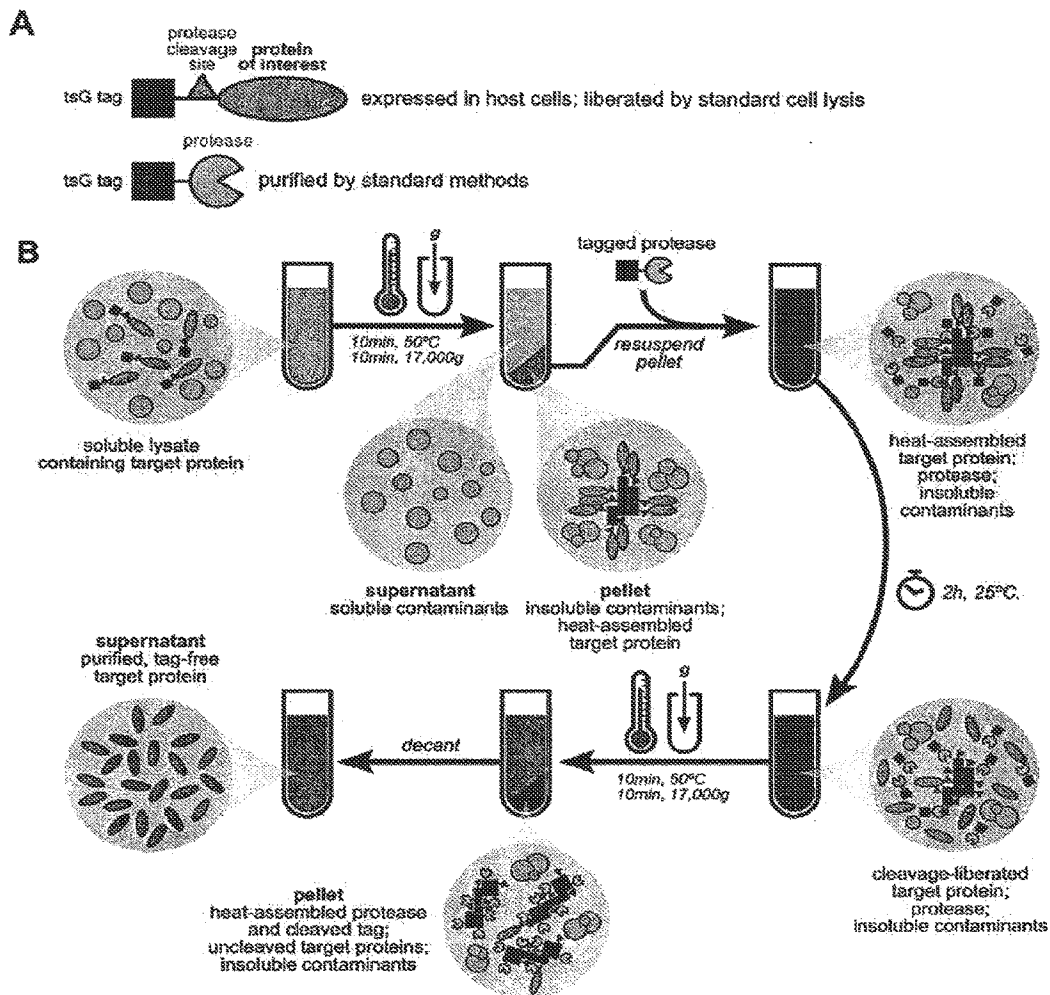


FIG. 3

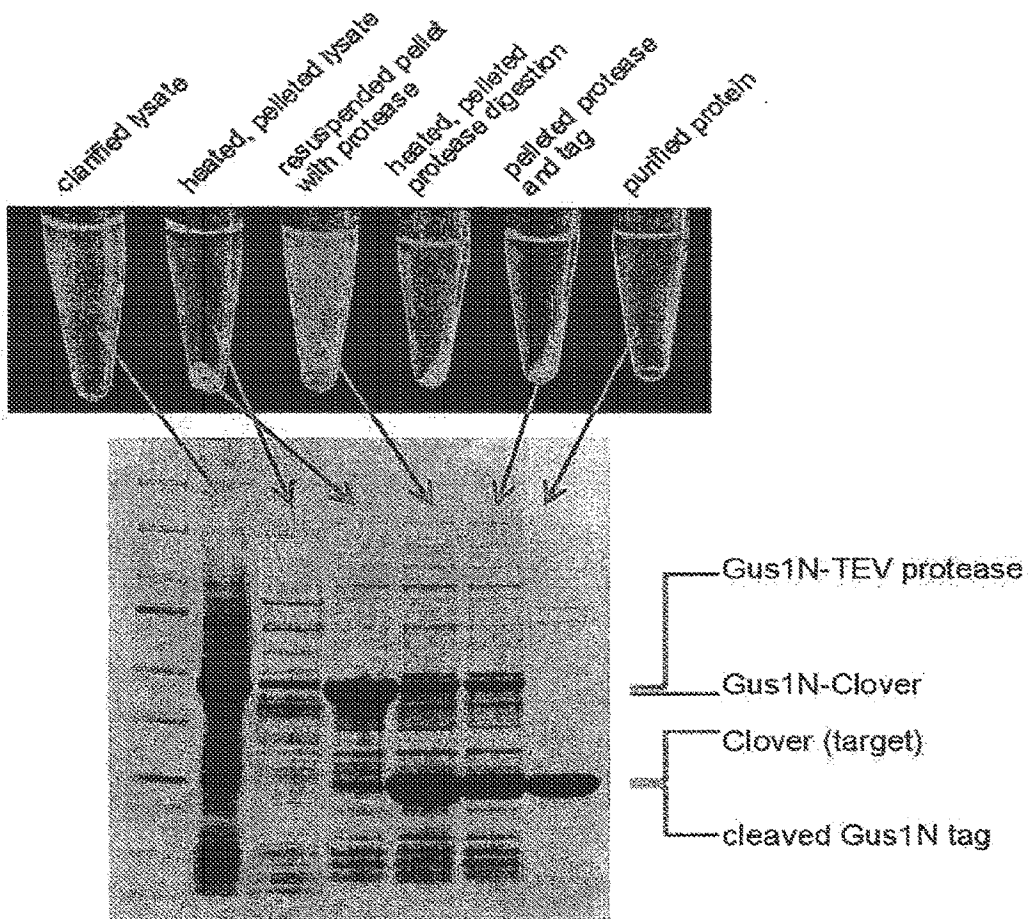


FIG. 4A

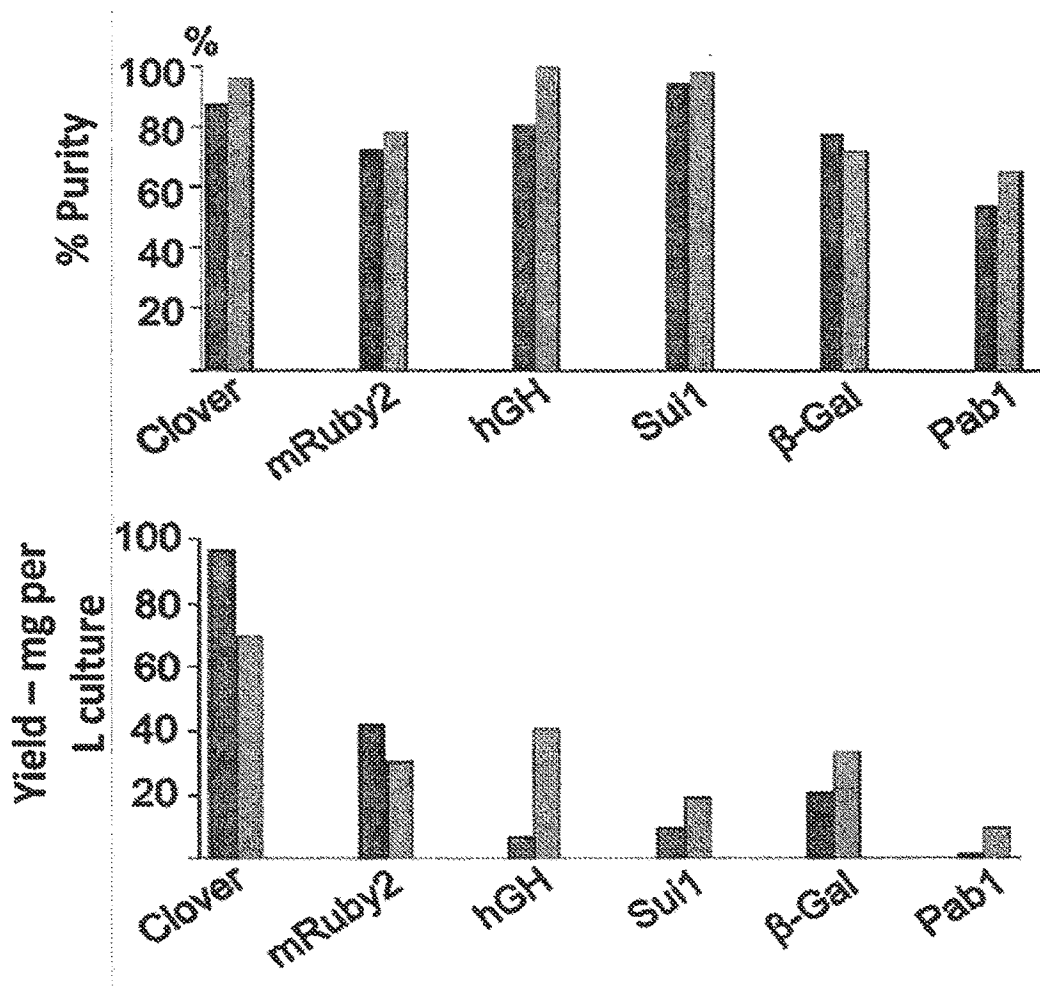


FIG. 4B

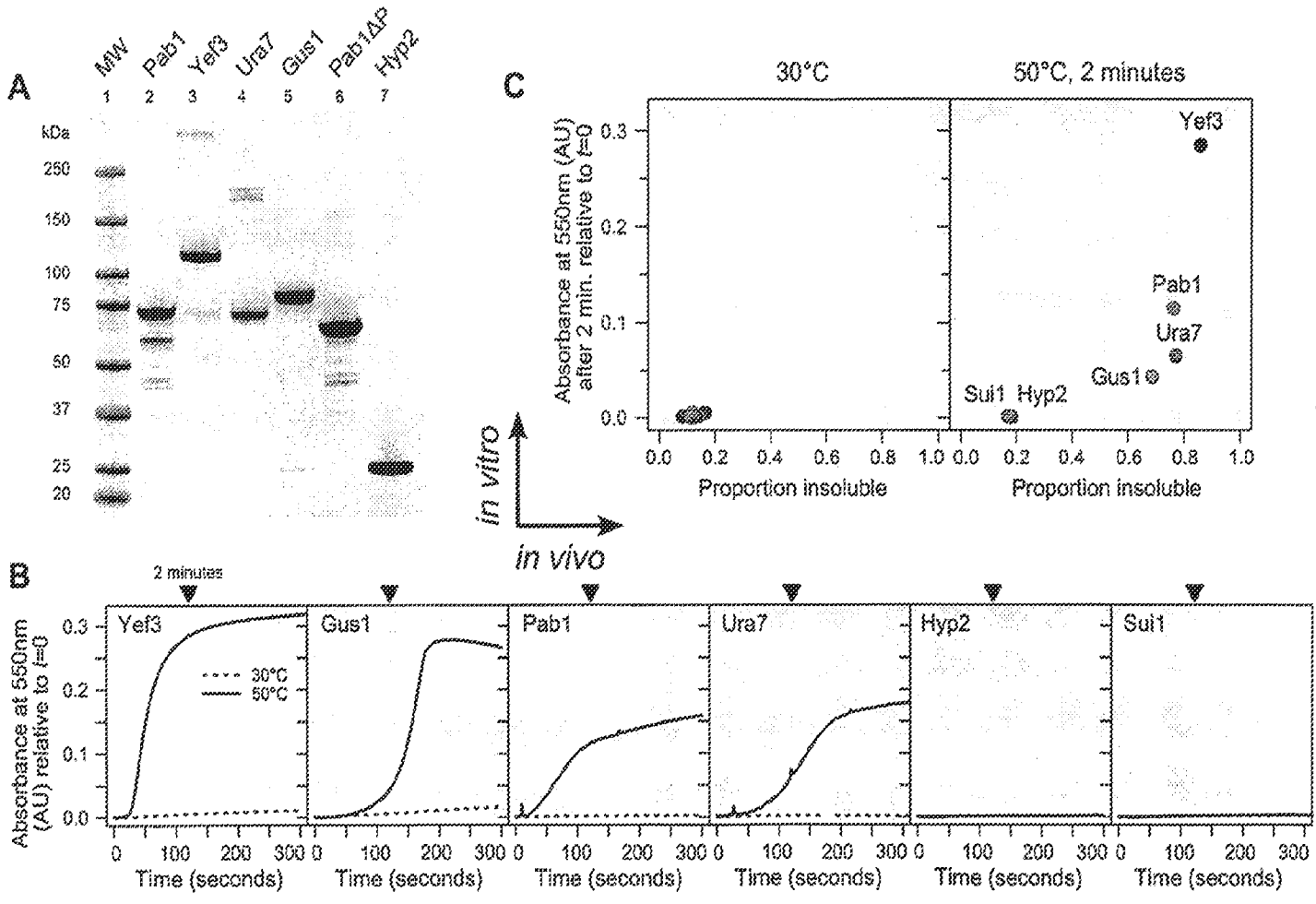


FIG. 6

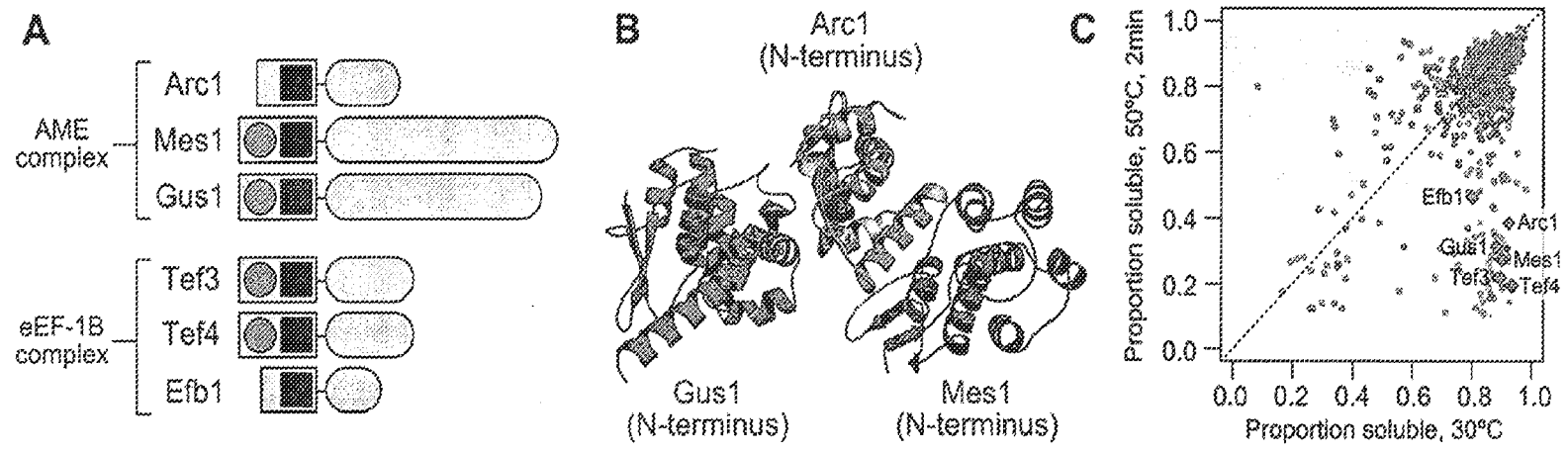


FIG. 7

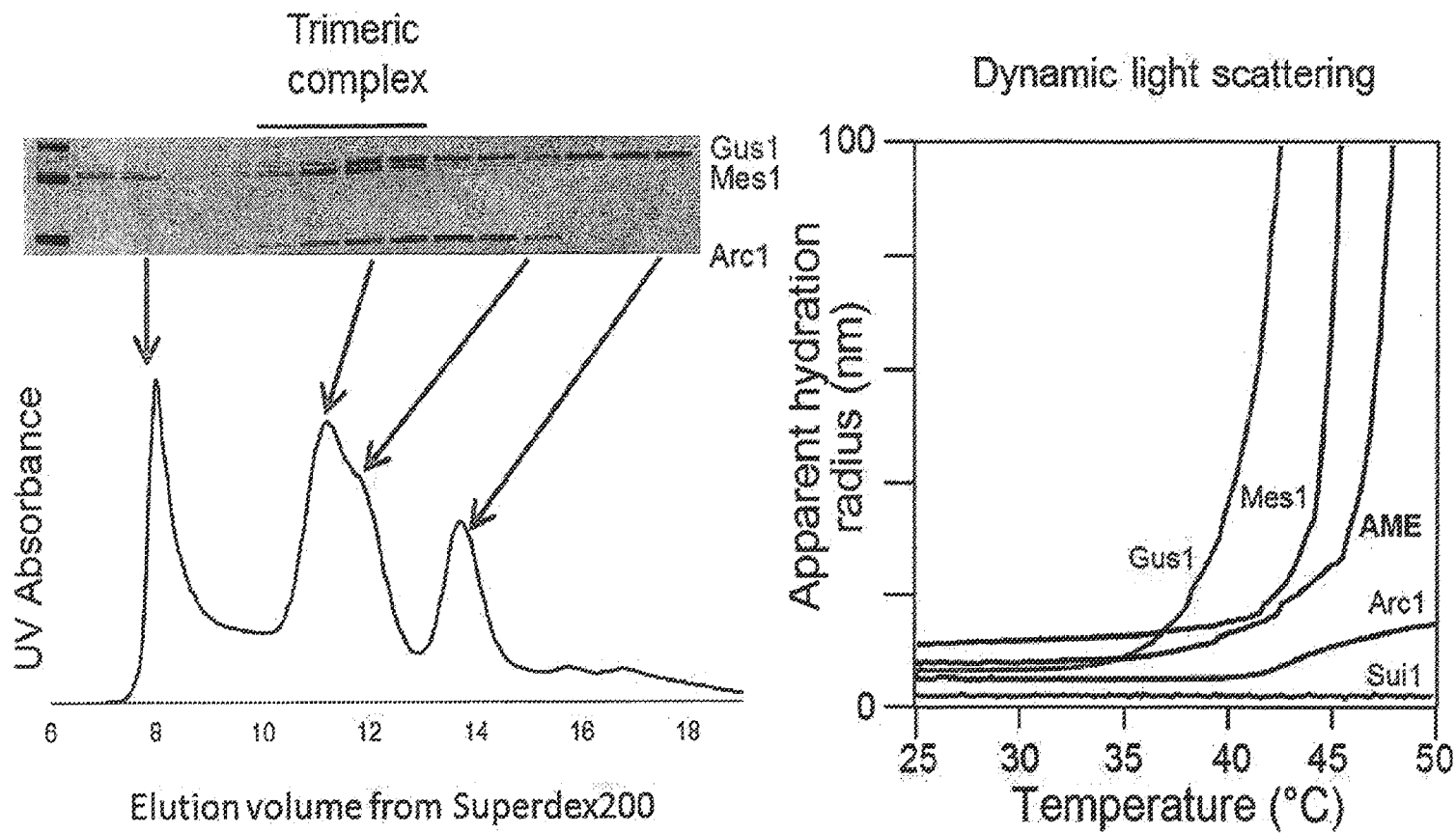


FIG. 8

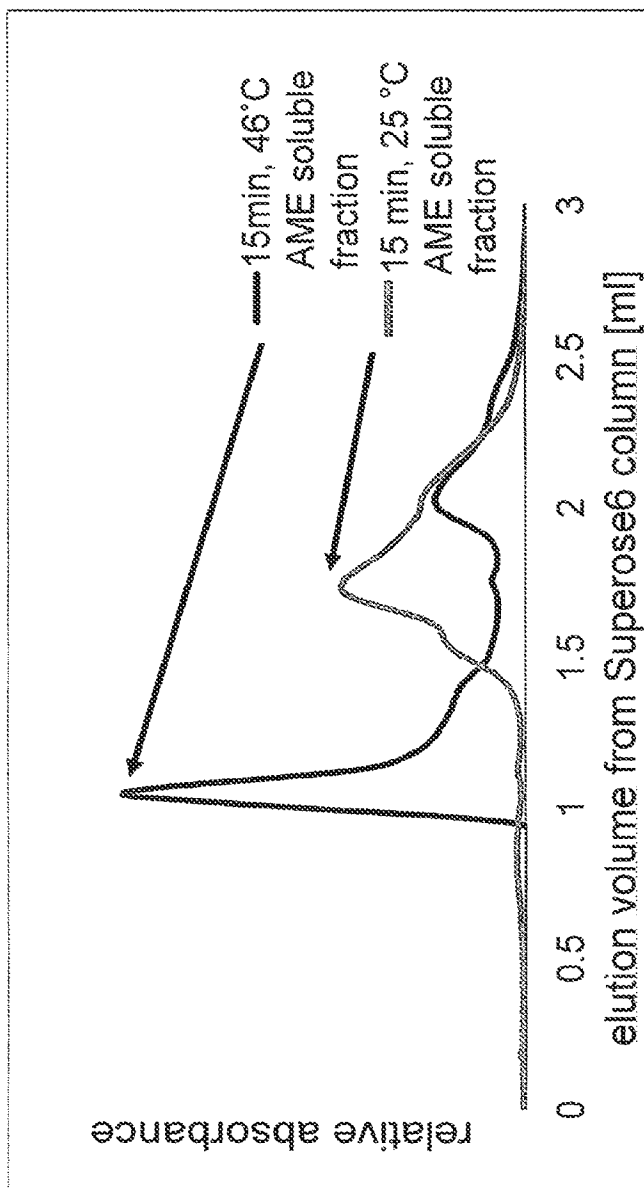


FIG. 9

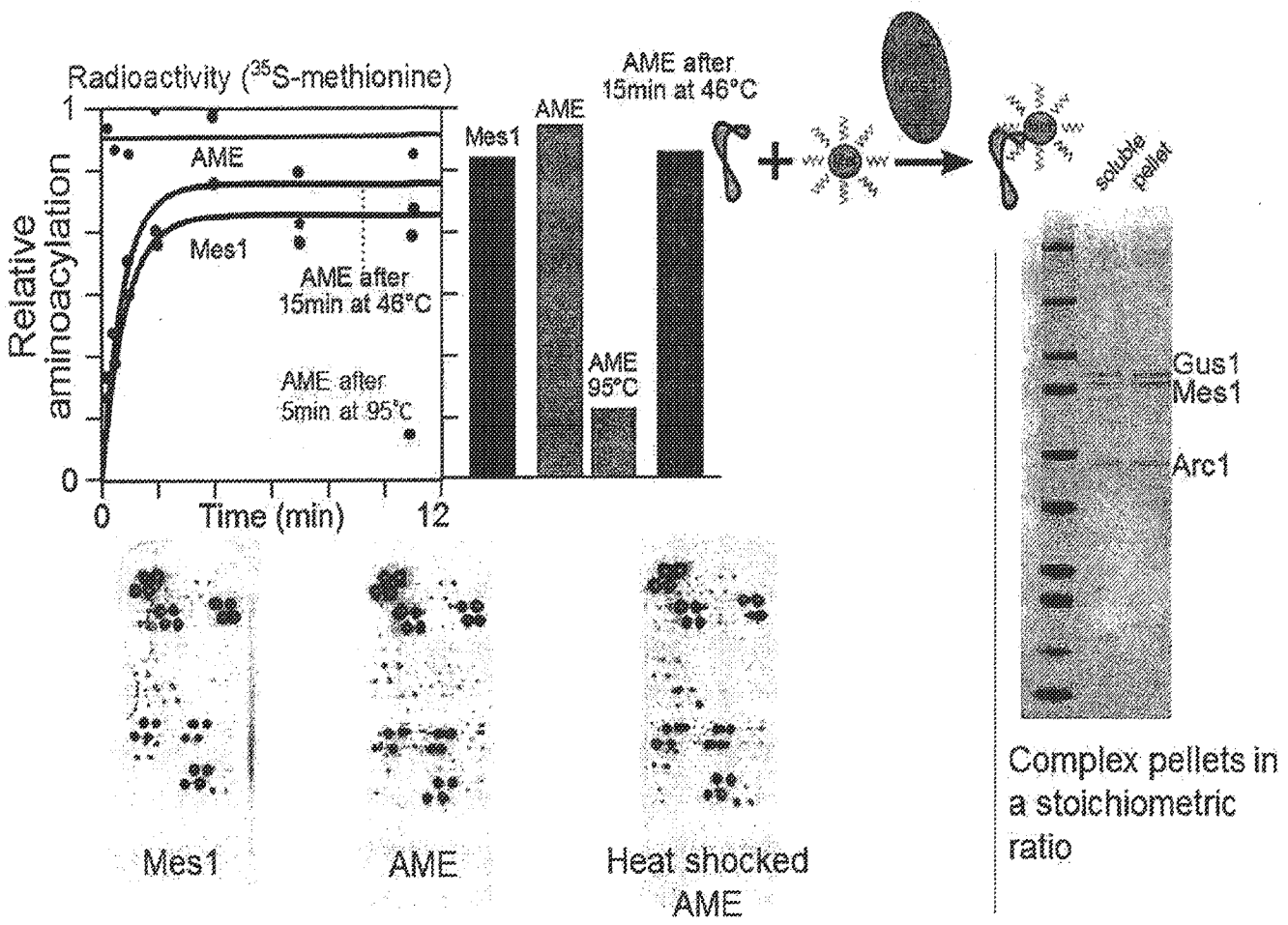


FIG. 10

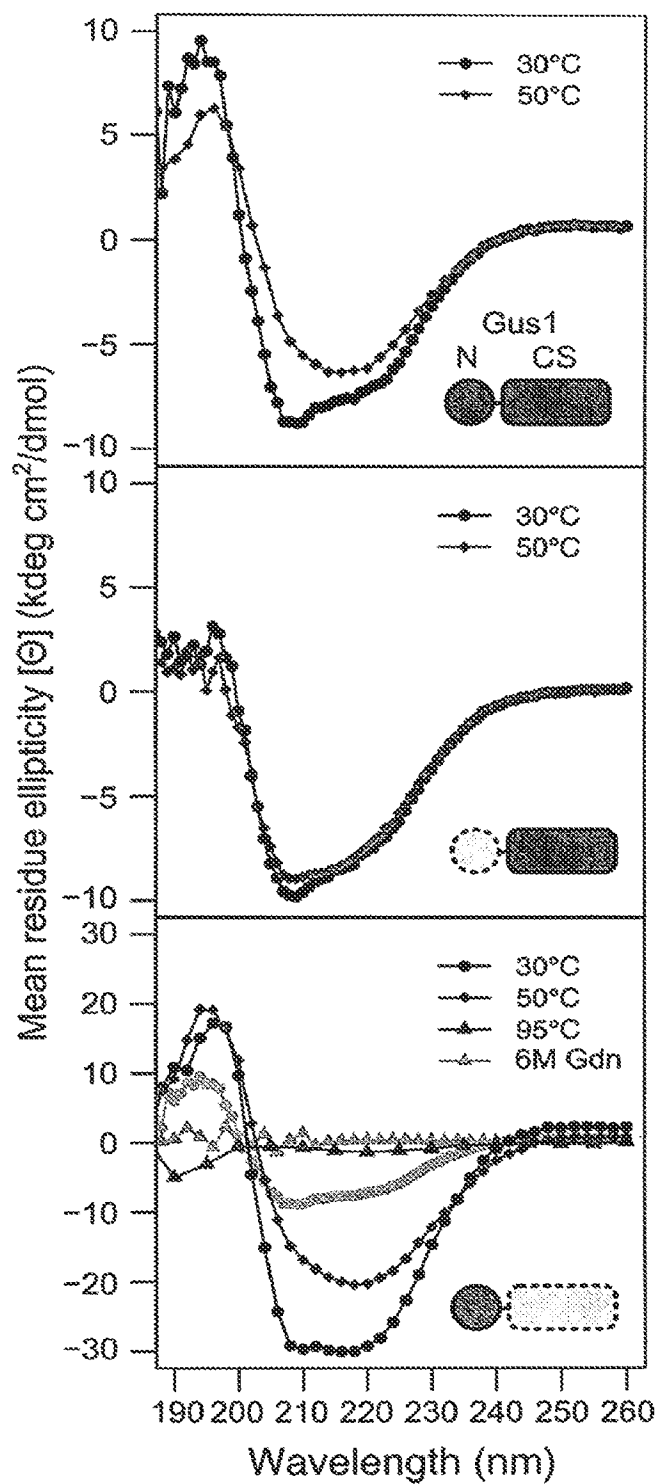


FIG. 11

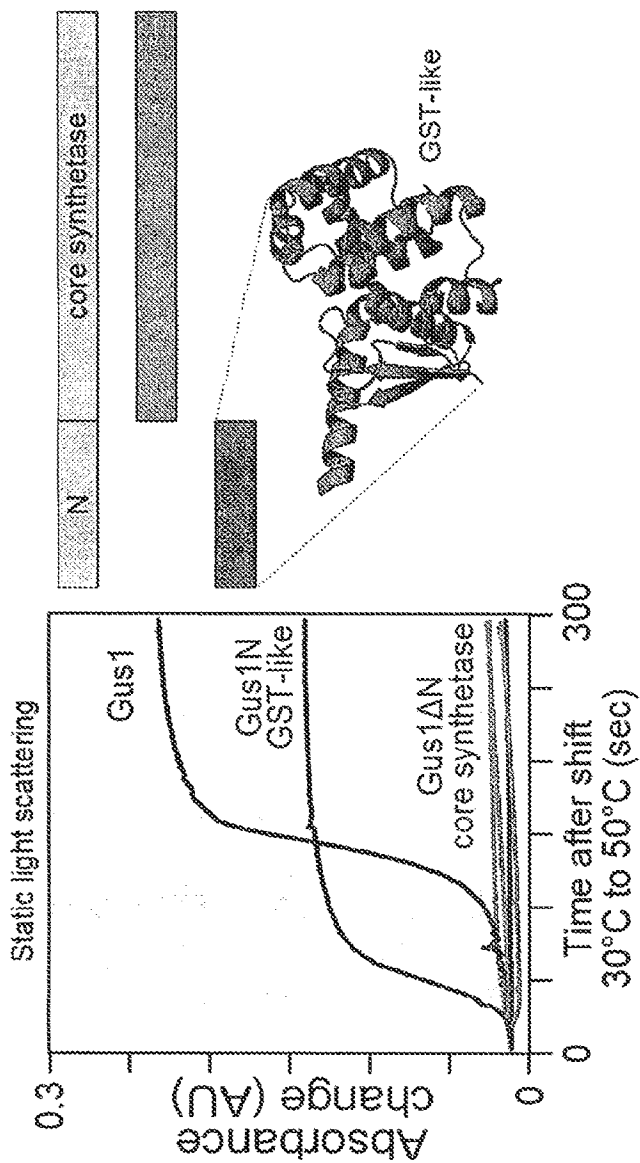


FIG. 12

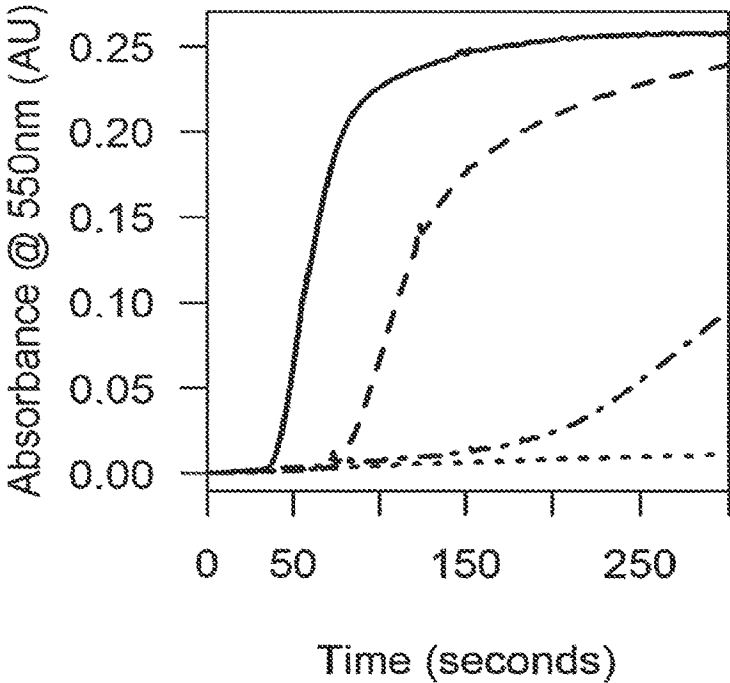


FIG. 13

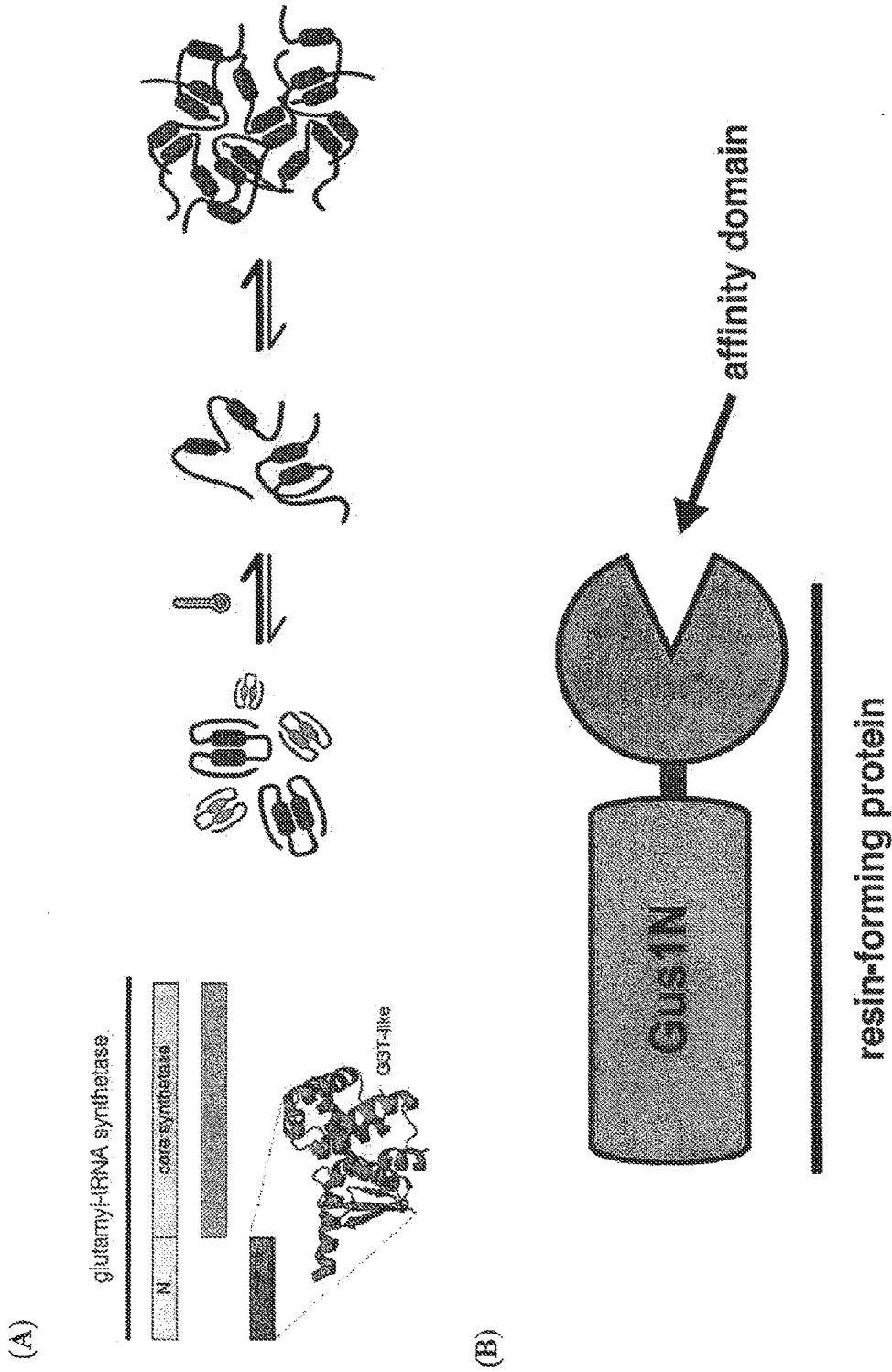


FIG. 14A-B

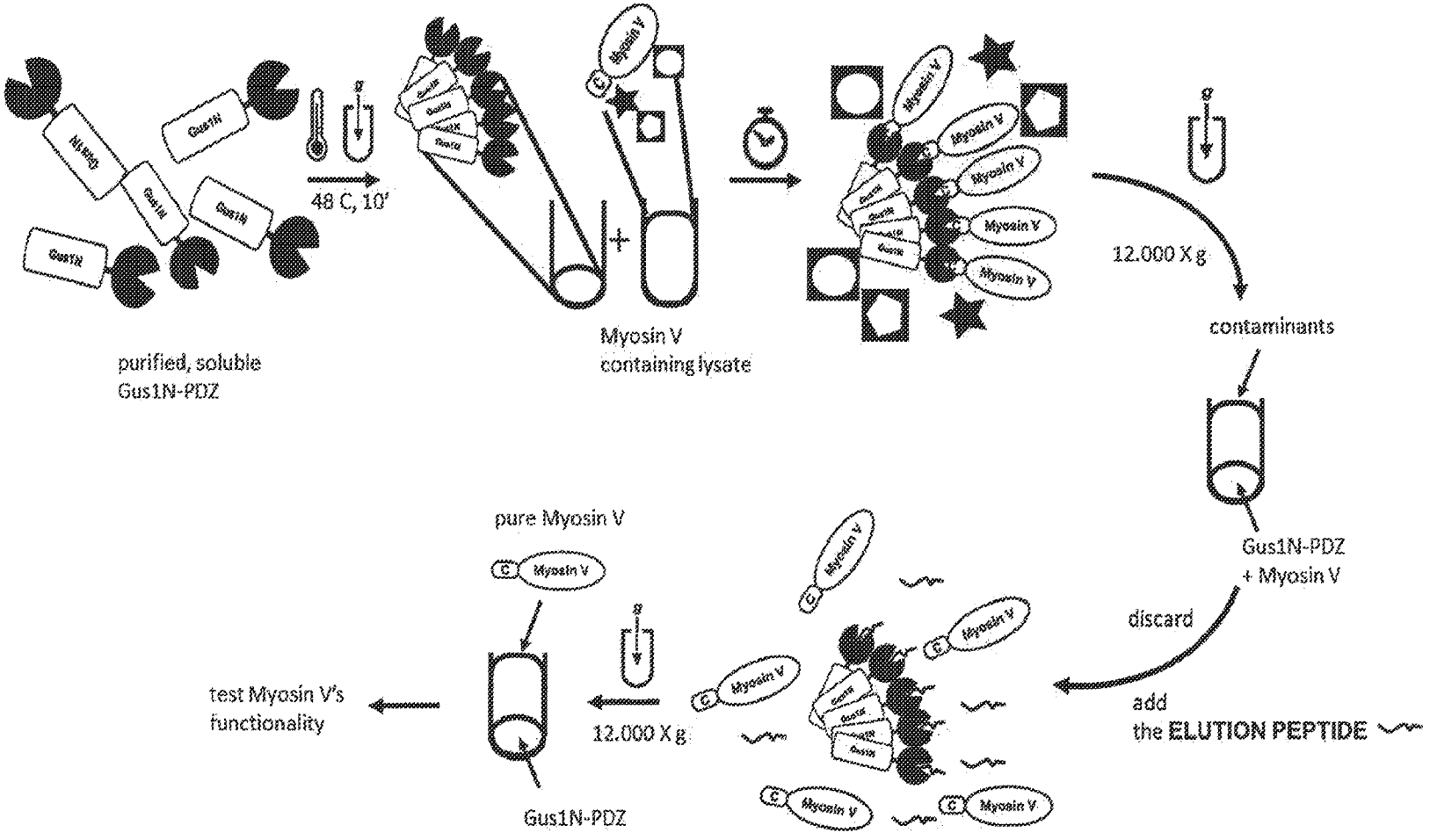


FIG. 15

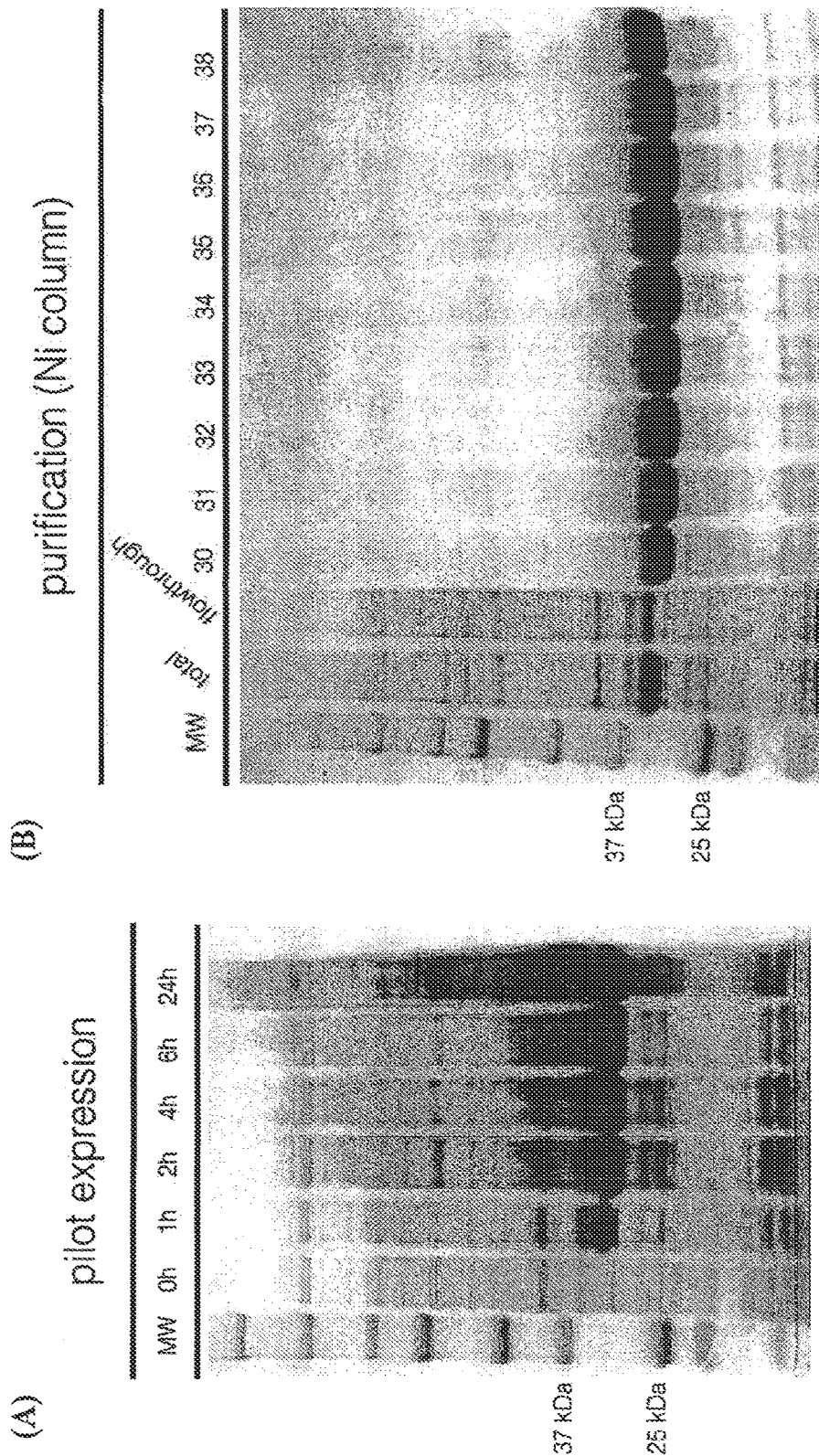


FIG. 16A-B

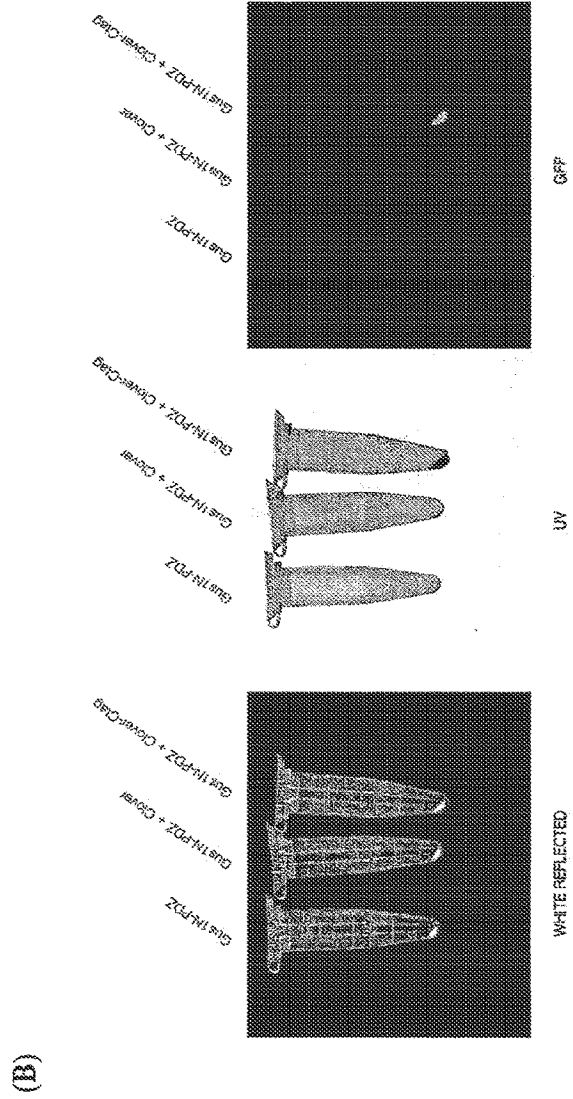
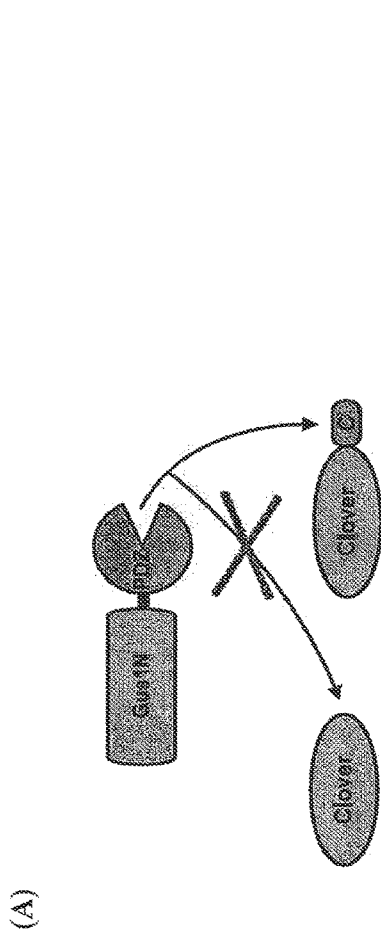


FIG. 17A-B

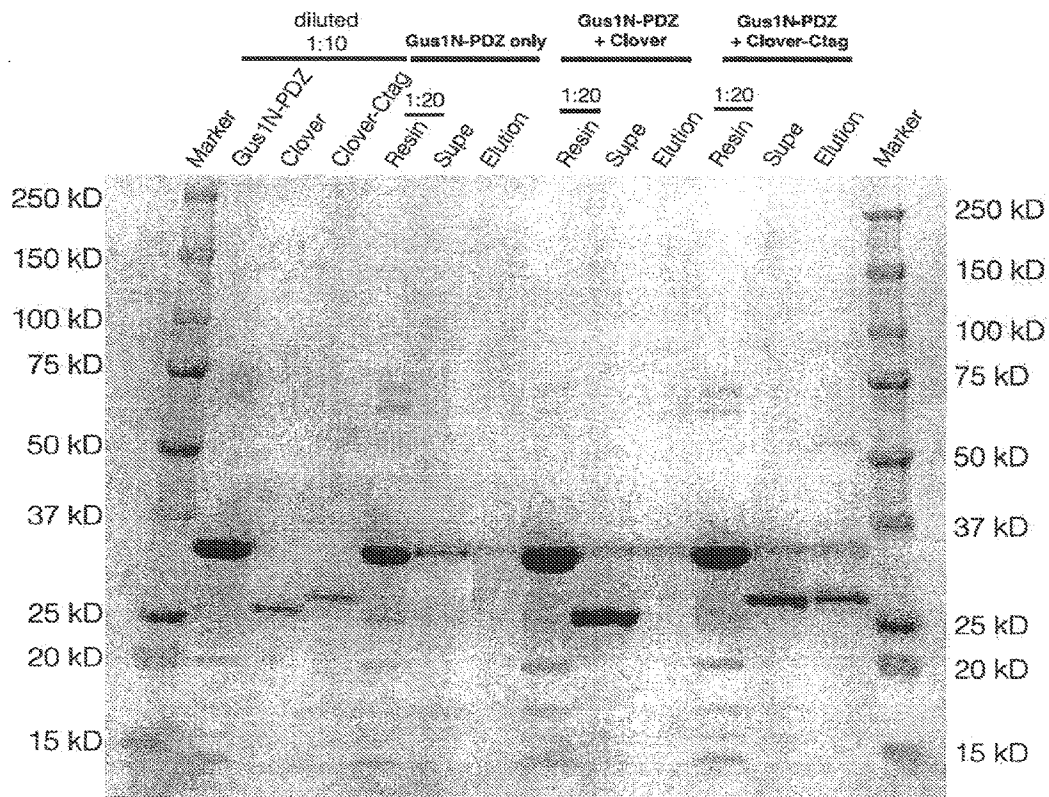


FIG. 18

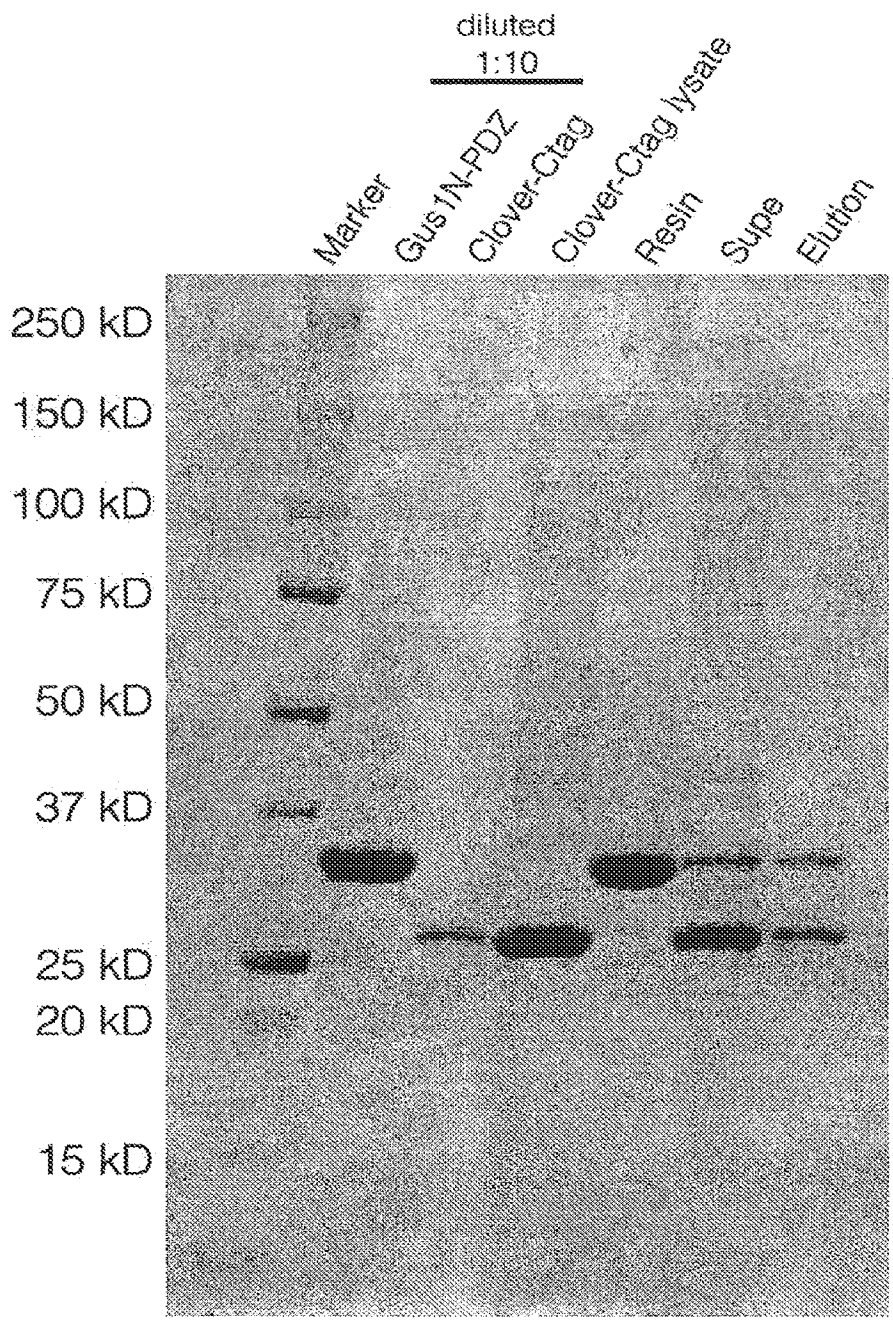


FIG. 19

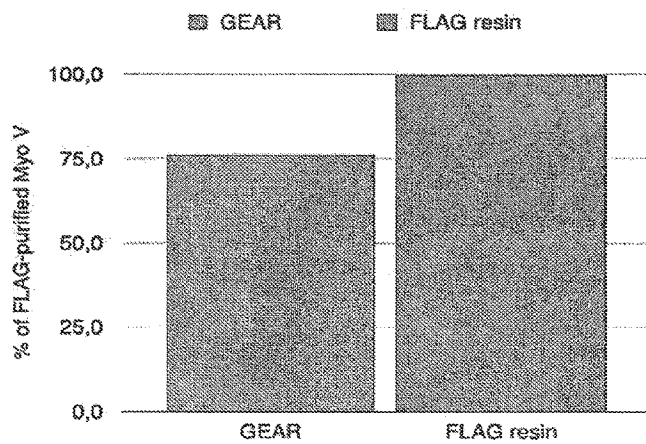
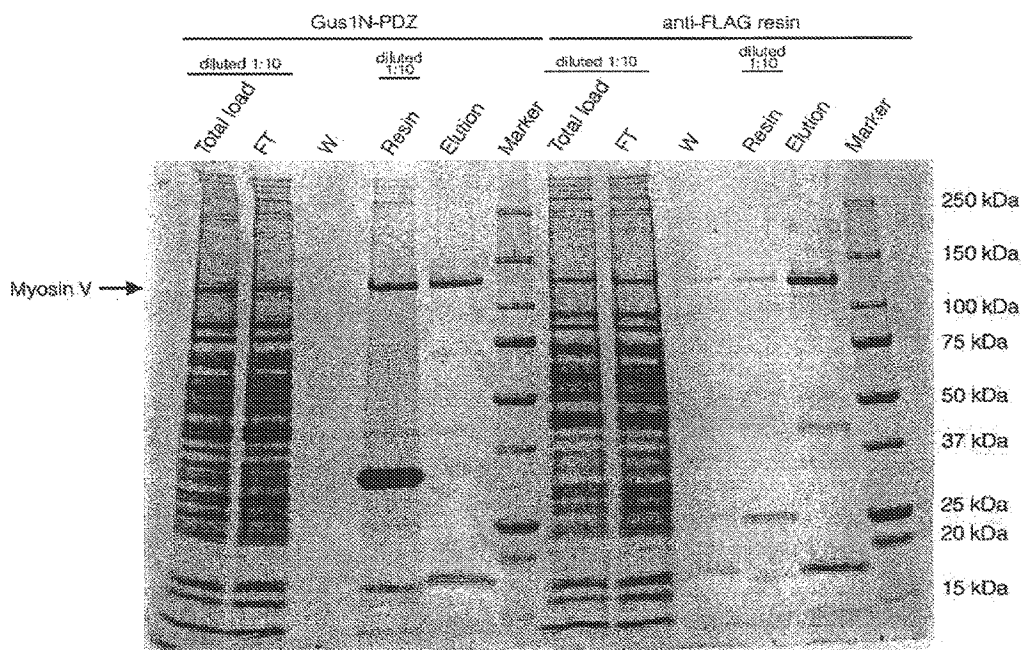


FIG. 20

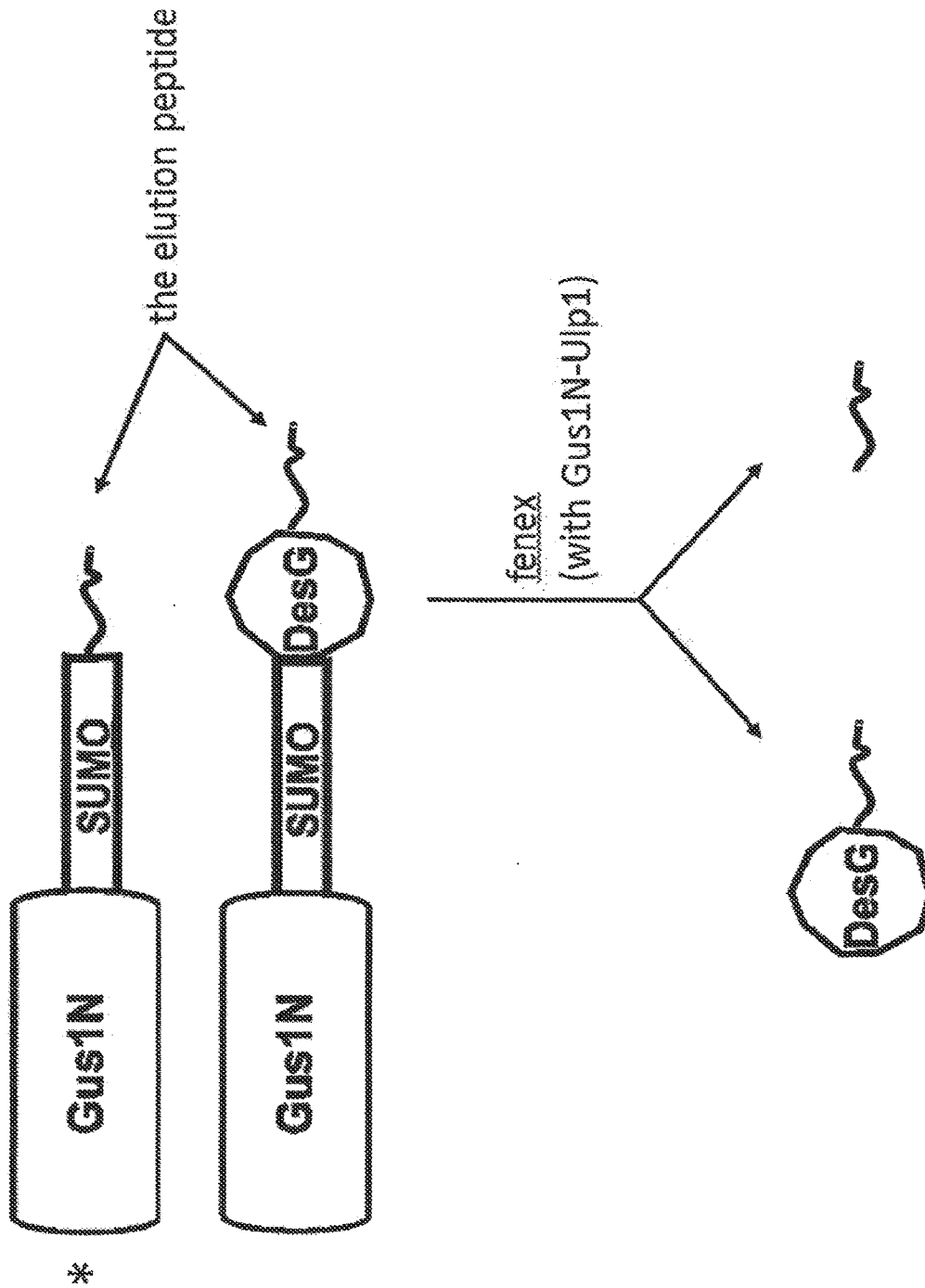


FIG. 21

HEAT-INDUCIBLE SELF-ASSEMBLING PROTEIN DOMAINS

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application claims the benefit of priority of U.S. Provisional Patent Application No. 62/068,390, filed Oct. 24, 2014, which is hereby incorporated by reference in its entirety.

BACKGROUND OF THE INVENTION

A. Field of the Invention

[0002] The invention generally concerns molecular biology and protein engineering. In particular, it involves proteins that are able to self-assemble in a heat-inducible fashion.

B. Description of Related Art

[0003] In many scientific applications, conjugation of molecules to larger albeit still microscopic particles (e.g., spheres, beads, rods, nanoparticles) is used to immobilize, control, partition, or otherwise manipulate other molecules. Sigma-Aldrich, Thermo Fisher Scientific, and many other companies sell particles made of sepharose, agarose, or other polymers which either come pre-conjugated with molecules (for example, antibodies, peptides, protein A, streptavidin, biotin) or can be conjugated by the customer using crosslinking chemistry. These particles can also be magnetic, allowing their rapid removal with a magnet.

[0004] Genetically encodable, heat-inducible, particle-forming protein domain would be valuable, as it would allow genetic engineering techniques that are now commonplace to be used to construct conjugated particles.

SUMMARY OF THE INVENTION

[0005] The present application provides heat-inducible, self-assembling protein domains and fusion proteins including such domains that can be used in many different applications described herein. For example, fusion proteins incorporating a self-assembly domain and a target protein provide improved, rapid purification methods. Such fusion proteins can be genetically encoded, expressed, and purified using conventional laboratory techniques. The fusion proteins can form protein aggregates upon heat induction, and can be used in the place of beads or other conjugated particles in many laboratory protocols.

[0006] Proteins containing a self-assembly domain form insoluble aggregates rapidly upon heat treatment, but are essentially absent and soluble at temperatures at or below 30° C., including room temperature. A self-assembly domain, fused to other proteins, confers self-assembling ability on these proteins. Folded proteins retain function within the assembled protein aggregates, and RNA, DNA, and small molecules can be stably bound within the assembled protein aggregates.

[0007] Disclosed herein is a self-assembling fusion protein comprising: (a) a heat-inducible self-assembly domain; and (b) a target protein; wherein the self-assembly domain remains folded (at least partially or mostly) during assembly. In some embodiments, the fusion protein is capable of self-assembling into protein aggregates by being heated to a temperature of between about 35 and 50° C. or any range

derivable therein. In some embodiments, the heat induction is at a temperature greater than, less than, or between any two of about 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, and 55° C.

[0008] In some embodiments of the disclosure, the fusion protein forms aggregates in less than or exactly 1200, 1100, 1000, 900, 800, 700, 600, 500, 400, 300, 275, 250, 225, 200, 175, 150, 125, 100, 75, 60, or 30 seconds upon heat induction, or any derivable range therein.

[0009] In some embodiments, the self-assembly domain is a GST-like domain or a polypeptide with at least 90% identity to a GST-like domain. A GST-like domain refers to a conserved protein domain known in the art as a Glutathione S-transferase C-terminal-like domain. This conserved domain is described in the NCBI database of conserved domains (See also Marchler-Bauer A. et al. (2013), "CDD: conserved domains and protein three-dimensional structure." *Nucleic Acids Res.* 41(D1):D384-52), which is hereby incorporated by reference. Furthermore, a protein can be determined to have this conserved domain by inputting the protein sequence into the NCBI conserved domain database, which can be found on the world wide web at ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi. The domain is capable of making stable protein-protein interactions between itself and another GST-like domain, in one of two orientations, as shown in Simader H et al. (2006), "Structural basis of yeast aminoacyl-tRNA synthetase complex formation revealed by crystal structures of two binary sub-complexes," *Nucleic Acids Res.* 34(14):3968-79.

[0010] The term "identity," "homology" or "similarity" refers to sequence similarity between two peptides or between two nucleic acid molecules. A degree of identity can be determined by comparing a position in each sequence which may be aligned for purposes of comparison. When a position in the compared sequence is occupied by the same base or amino acid, then the molecules are homologous at that position. A degree of identity between sequences is a function of the number of matching positions shared by the sequences. An "unrelated" or "non-homologous" sequence shares less than 40% identity, though preferably less than 25% identity, with one of the sequences of the present invention. The percent identity can be calculated by the formula: $(\text{Matches} \times 100) / \text{Length of aligned region (with gaps)}$. Note that only internal gaps are included in the length, and not gaps at the sequence ends.

[0011] A polynucleotide or polynucleotide region (or a polypeptide or polypeptide region) having a certain percentage (for example, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 98% or 99%, or any range derivable therein) of "sequence identity" or "homology" to another sequence means that, when aligned, that percentage of bases (or amino acids) are the same in comparing the two sequences. This alignment and the percent homology or sequence identity can be determined using software programs known in the art, for example those described in Ausubel et al. eds. (2007) *Current Protocols in Molecular Biology*.

[0012] In some embodiments, a polypeptide may have at least 65, 70, 75, 80, 85, 90, 95, 96, 97, 98, 99, or 100% identity (or any range derivable thereof) with another polypeptide.

[0013] In some embodiments, the self-assembly domain comprises a polypeptide from Arc1, Mes1, Gus1, or a polypeptide with at least 90% identity to Arc1, Mes1, or Gus1. Arc1, Mes1, and Gus1 are proteins with GST-like

domains. In some embodiments, the self-assembly domain comprises a polypeptide that is at least 20 amino acids in length and has at least 90% identity to the first 250 amino acids to Arc1, Mes1, or Gus1. In this case, the percent identity is calculated specifically as described above, wherein the sequence is aligned, and internal gaps are used to calculate sequence identity, but gaps at the end of the alignment are not used in calculating sequence identity. In some embodiments, the self-assembly domain comprises a polypeptide that is at least 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190, or 200 amino acids in length and has at least 90% identity to a polypeptide of similar or the same length from the first 250 amino acids to Arc1, Mes1, or Gus1. In some embodiments, the self-assembly domain comprises a polypeptide that is at least 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190, or 200 amino acids in length, or any derivable range thereof. The polypeptide may have a certain degree of identity to the first (N-terminus) 50, 75, 100, 125, 150, 175, 191, 200, 225, 250, 300, or 300 amino acids (or any derivable range thereof) of Arc1, Mes1, or Gus1. In some embodiments, the self-assembly domain comprises a polypeptide from Gus1 or a polypeptide with at least 90, 95, 97, or 99% identity (or any range derivable thereof) to Gus1.

[0014] In some embodiments, the self-assembly domain comprises a polypeptide from Tef3, Tef4, Efb1, or a polypeptide with at least 90% identity to Tef3, Tef4, or Efb1. Tef3, Tef4, and Efb1 are proteins with GST-like domains. In some embodiments, the self-assembly domain comprises a polypeptide that is at least 20 amino acids in length and has at least 90% identity to 20 amino acids of the first 250 amino acids from Tef3, Tef4, or Efb1. In this case, the percent identity is calculated specifically as described above, wherein the sequence is aligned, and internal gaps are used to calculate sequence identity, but gaps at the end of the alignment are not used in calculating sequence identity. In some embodiments, the self-assembly domain comprises a polypeptide that is at least 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190, or 200 amino acids in length, or any derivable range thereof. The polypeptide may have a certain degree of identity to the first (N-terminus) 50, 75, 100, 125, 150, 175, 191, 200, 225, 250, 300, or 300 amino acids (or any derivable range thereof) of Tef3, Tef4, or Efb1.

[0015] In some embodiments, the self-assembly domain comprises a polypeptide from Yef3, Ura7, or a polypeptide with at least 90% identity to Yef3 or Ura7.

[0016] In some embodiments, the polypeptide is from a *Saccharomyces cerevisiae* protein. In some embodiments, the polypeptide of the assembly domain is from a homolog of Arc1, Mes1, Gus1, Tef3, Tef4, Efb1, Yef3, or Ura7 from another organism. For example, the polypeptide may be a homolog of Arc1, Mes1, Gus1, Tef3, Tef4, Efb1, Yef3, or Ura7 from *B. dendrobatidis*, *U. hordei*, *U. maydis*, *S. reilianum*, *P. triticina*, *P. graminis*, *W. sebi*, *M. globosa*, *M. larici-populi*, *F. radiculosa*, *P. indica*, *S. lacrymans*, *C. cinerea*, *P. carnosa*, *A. bisporus*, *Allavus*, *C. neoformans*, *C. gattii*, *A. oligospora*, *Lelongisporus*, *N. tetrasperma*, *C.*

parapsilosis, *C. albicans*, *D. hansenii*, *C. dubliniensis*, *C. lusitanae*, *P. sorbitophila*, *R. delemar*, *T. melanosporum*, *P. pastoris*, *S. stipitis*, *S. japonicus*, *C. tenuis*, *W. ciferrii*, *P. angusta*, *T. stipitatus*, *S. pombe*, *A. terreus*, *N. fumigata*, *P. marneffei*, *N. fischeri*, *A. clavatus*, *A. oryzae*, *P. digitatum*, *P. chrysogenum*, *A. niger*, *A. kawachii*, *T. verrucosum*, *N. dairenensis*, *P. brasiliensis*, *A. benhamiae*, *A. capsulata*, *A. gypseum*, *K. lactis*, *N. castellii*, *A. dermatitidis*, *T. rubrum*, *T. equinum*, *S. arboricola*, *S. cerevisiae*, *T. blattae*, *K. naganishii*, *C. posadasii*, *A. gossypii*, *V. polyspora*, *T. phaffii*, *A. otae*, *Z. rouxii*, *T. delbrueckii*, *K. africana*, *C. glabrata*, *L. thermotolerans*, *Y. lipolytica*, *E. aedis*, *C. militaris*, *U. reesii*, *P. froveci*, *E. nidulans*, *T. tonsurans*, *T. asahii*, *S. commune*, *L. maculans*, *P. teres*, *N. sp.*, *N. parisii*, *E. hellem*, *E. cuniculi*, *E. romaleae*, *E. intestinalis*, *P. tritici-repen*, *C. globosum*, *T. terrestris*, *G. clavigera*, *T. heterothallic*, *B. bassiana*, *M. phaseolina*, *V. corneae*, *C. thermophilum*, *N. crassa*, *M. oryzae*, *G. graminis*, *H. atroviridis*, *H. vixens*, *Hjedorina*, *V. culicis*, *T. hominis*, *S. sclerotiorum*, *B. fuckeliana*, *N. haematococca*, *M. robertsii*, *E. bienersi*, *M. acridum*, *F. oxysporum*, *C. graminicola*, *C. gloeosporioides*, *G. destructans*, *G. lozoyensis*, *C. higginsianum*, *M. brunnea*, *V. dahliae*, *F. pseudogramine*, *S. macrospora*, *N. ceranae*, *V. albo-atrum*, *S. passalidarum*, *C. tropicalis*, *M. guilliermondii*, or *E. dermatitidis*.

[0017] The systematic name and common name of proteins that are useful for the self-assembly domain are:

Systematic Name	Common Name
YGL105W	Arc1
YGR264C	Mes1
YGL245W	Gus1
YPL048W	Tef3
YKL081W	Tef4
YAL003W	Efb1
YLR249W	Yef3
YBL039C	Ura7

[0018] In some embodiments, the self-assembly domain comprises a polypeptide of SEQ ID NO:9, 10, 11, or 12, a fragment thereof, or a polypeptide with at least 90% identity to SEQ ID NO:9, 10, 11, or 12, or a fragment thereof. SEQ ID NOs:9-12 represents the N-terminal GST-like domains of Gus1, Mes1, Tef4, and Tef3.

[0019] In some embodiments, the majority of the fusion protein molecules aggregate to form direct protein-protein interactions with other fusion protein molecules upon heat-induction. "Direct" protein-protein interactions are not mediated by solvent or any other molecule, but involve the direct non-covalent interaction of amino acids with other amino acids. These direct interactions may be aromatic-aromatic, cation-aromatic, electrostatic, van der Waals, or hydrophobic interactions. The fusion protein is capable of forming the protein aggregates under relatively "mild" conditions, which is one way that the protein aggregates described herein are more useful than other types of protein aggregates for laboratory applications such as forming aggregates conjugated to target proteins, which aggregates can be used in the place of beads or solid substrates in many laboratory protocols. The term "aggregate" in the prior art may refer to mis-folded proteins under denaturing conditions (e.g., elevated temperatures, pH, high salt content). However, the term "protein aggregates," as used herein, is

not meant to refer to mis-folded protein, but instead refers to protein that substantially retains a tertiary structure, but associates with the self-assembly domain of any other proteins in the composition. The self-assembly domain, while aggregated, substantially retains function and a tertiary structure. While the self-assembly domain does undergo a conformational change after a temperature shift, the self-assembly domain is not mis-folded at the temperature shift and substantially retains a tertiary structure. Therefore, in some embodiments, the self-assembly domain remains folded at a temperature below, above, or any range derivable thereof, of 60, 55, 50, 45, 40, 35, 30, 25, or 20° C. In some embodiments, the self-assembly domain remains folded at a temperature range of 20–50° C.

[0020] In some embodiments, the self-assembly domain is not an elastin-like polymer (ELP), does not have a significant degree of homology to an ELP, and/or does not comprise an ELP polypeptide or fragment.

[0021] In some embodiments, the fusion proteins do not self-assemble at temperatures below about 35° C. In some embodiments, the fusion proteins do not self-assemble at temperatures below about 40, 35, 30, 25, 20, or 15° C. (or any range derivable therein).

[0022] In some embodiments, the self-assembly domain is at least, at most, or exactly 15, 25, 50, 75, 100, 125, 150, 175, 200, 225, 250, 300, 350, 400, 500, 600, 700, 800, 900, and 1000 amino acids in length, or any range derivable therein.

[0023] In some embodiments, the target protein or polypeptide is at least, at most, or exactly 5, 10, 15, 25, 50, 75, 100, 125, 150, 175, 200, 225, 250, 300, 350, 400, 500, 600, 700, 800, 900, and 1000 amino acids in length, or any range derivable therein.

[0024] The target protein may be a protein in which purification is desired or may be a component of an assay, such as an antibody or protease. In some embodiments, the target protein is ferritin, a fluorescent protein, an antibody, an antibody fragment, protein A, streptavidin, protein G, protein A/G, protein L, a protease, or StrepTactin. In some embodiments, the target protein is a protease. In some embodiments, the target protein is a wild-type or mutant protein from a eukaryote or a prokaryote. In some embodiments, the target protein is not a naturally occurring protein.

[0025] In some embodiments, the fusion protein further comprises a protease cleavage site. In some embodiments, the protease cleavage site is between the target protein and the self-assembly domain.

[0026] Further aspects of the disclosure relate to a protein aggregate comprising the fusion protein described herein. In some embodiments, the protein aggregate further comprises a nucleic acid or protein that is specifically bound to the target protein. In some embodiments, a small molecule is specifically bound to the target protein. Further aspects relate to an aqueous composition comprising the fusion protein or the protein aggregate described herein.

[0027] In some embodiments, the protein aggregate is in an aqueous composition, wherein the aqueous composition comprises between about 0 and 500 mM KCl or NaCl, or is about or is less than any one of or between any two of about 50, 100, 150, 200, 250, 300, 350, 400, 450, and 500 mM KCl or NaCl. In some embodiments, the aqueous composition does not include alcohol. In some embodiments the fusion protein is present in the aqueous composition at a concentration of between about 5 and 50 μ M, or at a concentration

of less than any one of or between any two of about 5, 10, 15, 20, 25, 30, 35, 40, 45, and 50 μ M. In some embodiments, the fusion protein is at least 55% pure, or is at least between any two of about 55, 60, 65, 70, 75, 80, 85, 90, 95, 98, 99, and 99.9% pure in the aqueous solution. In some embodiments, the pH of the aqueous composition is between about 6.0 and 8.0, or is about or is greater than, less than, or between any two of about 6.0, 6.1, 6.2, 6.3, 6.4, 6.5, 6.6, 6.7, 6.8, 6.9, 7.0, 7.1, 7.2, 7.3, 7.4, 7.5, 7.6, 7.7, 7.8, 7.9, and 8.0.

[0028] In some embodiments, the target protein in the fusion proteins described above is a restriction enzyme, DNA polymerase, protease, ligase, RNA polymerase, methylase, polyadenylate polymerase, topoisomerase, guanylyl transferase, ribonuclease, deoxyribonuclease, alkaline phosphatase, polynucleotide kinase or reverse transcriptase. In some embodiments, the target protein is a therapeutic protein.

[0029] Further aspects of the disclosure relate to a polynucleotide coding for the fusion protein described herein. Other embodiments of the disclosure relate to a host cell comprising the fusion protein or the polynucleotide described herein. Yet further aspects relate to cell lysate comprising the fusion protein as described herein.

[0030] Also disclosed is a method for aggregating a target protein comprising: formulating an aqueous composition comprising the fusion protein as described herein; and heating the composition to a temperature between about 35 and 50° C. or any range derivable therein. In some embodiments, the temperature is above, below, or a derivable range of about 20, 25, 30, 35, 40, 45, 50, 55, or 60° C.

[0031] Further method aspects of the disclosure relate to a method of selectively depleting a molecule from an aqueous composition comprising: formulating an aqueous composition comprising the molecule and the fusion protein as described herein, wherein the target protein is a protein that specifically binds to the molecule; heating the aqueous composition to a temperature between about 35 and 50° C. to form protein aggregates comprising the fusion protein and the molecule; and removing the protein aggregates from the aqueous composition. In some embodiments, the temperature is above, below, or a derivable range of about 20, 25, 30, 35, 40, 45, 50, 55, or 60° C. In some embodiments, the molecule is a nucleic acid. The molecule may be, for example, a DNA or a RNA. In some embodiments, removing the protein aggregates from the aqueous composition is performed by centrifuging or filtering the aqueous composition.

[0032] Further method aspects of the disclosure relate to a method of selectively depleting a molecule from an aqueous composition comprising: heating the aqueous composition comprising the fusion protein to a temperature between about 35 and 50° C. to form protein aggregates comprising the fusion protein; adding the molecule to the aqueous composition comprising the aggregated fusion protein, wherein the target protein is a protein that specifically binds to the molecule; and removing the protein aggregates from the aqueous composition. In some embodiments, the temperature is above, below, or a derivable range of about 20, 25, 30, 35, 40, 45, 50, 55, or 60° C. In some embodiments, the molecule is a nucleic acid. The molecule may be, for example, a DNA or a RNA. In some embodiments, removing the protein aggregates from the aqueous composition is performed by centrifuging or filtering the aqueous composition.

[0033] The removal or separation of the protein aggregates may be done by methods known in the art for separating soluble and insoluble (protein aggregates) fractions. These include, for example, centrifugation, filtration, and size exclusion chromatography.

[0034] Other aspects relate to a method of immunoprecipitating a molecule comprising: formulating an aqueous composition comprising the molecule and the fusion protein of any one of claims 1-20; wherein the target protein is an antibody or antigen binding fragment that specifically binds to the molecule; and heating the aqueous composition to a temperature between about 35 and 50° C. to form protein aggregates comprising the fusion protein and the molecule. In some embodiments, the temperature is above, below, or a derivable range of 20, 25, 30, 35, 40, 45, 50, 55, or 60° C. In some embodiments, the method further comprises detecting the molecule bound in the protein aggregate. In some embodiments, the method further comprises quantifying the molecule bound in the protein aggregate. In some embodiments, the method further comprises separating the protein aggregates from the soluble composition.

[0035] Further method aspects relate to a method for purifying a protein comprising: formulating an aqueous composition comprising a fusion protein as described herein; heating the aqueous composition to a temperature between about 35 and 50° C. to form protein aggregates comprising the fusion protein; and separating the protein aggregates from the aqueous composition. In some embodiments, the temperature is above, below, or a derivable range of 20, 25, 30, 35, 40, 45, 50, 55, or 60° C. In some embodiments, the fusion protein is a first fusion protein comprising a protein cleavage site between the self-assembly domain and the target protein. In some embodiments, the method further comprises: cleaving the first fusion protein by formulating an aqueous composition comprising a second fusion protein and the first fusion protein in the separated aggregate; wherein the target protein of the second fusion protein is a protease that cleaves the first fusion protein at the protein cleavage site between the self-assembly domain and the target protein of the first fusion protein; heating the aqueous composition to a temperature between about 35 and 50° C. to form protein aggregates comprising the second fusion protein and the self-assembly domain of the cleaved first fusion protein; wherein the target protein of the first fusion protein remains soluble; and separating the protein aggregates from the soluble target protein of the first fusion protein. This purification method is further described in the figures and examples.

[0036] Further aspects relate to a method of immunoprecipitating or purifying a molecule comprising the steps of: formulating a first composition comprising the fusion protein as described herein; wherein the target protein is a first target protein that specifically binds to the molecule; heating the first composition to a temperature between about 35 and 50° C. to form protein aggregates comprising the fusion protein; and contacting the first composition with a second composition comprising the molecule. In some embodiments, the first target protein that specifically binds to the molecule is an antibody, an antigen binding fragment, or an affinity tag (e.g., PDZ domain). In some embodiments, the composition is heated to at least, at most, or exactly about 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 65, or 70° C., or any derivable range therein. In some embodi-

ments, the heating of the composition is done prior to contacting of the first composition with the second composition. In some embodiments, the heating of the first composition is done after contacting the first composition with the second composition.

[0037] In some embodiments, the second composition maintains a temperature of less than 40° C. throughout the method. In some embodiments, the second composition and/or second target protein maintains a temperature of less than 45, 44, 43, 42, 41, 40, 39, 38, 37, 36, 35, 34, 33, 32, 31, 30, 29, 28, 27, 26, 25, 24, 23, 22, 21, or 20° C. throughout the method. In some embodiments, contacting the first composition with the second composition comprises mixing the compositions. The mixing may be done by mechanical means such as, for example, vortexing, pipetting, etc. In some embodiments, the molecule is a fusion protein between a second target protein and a tag that binds to the first target protein. In some embodiments, the first target protein is a PDZ domain. In some embodiments, the tag is a Ctag. In some embodiments, the molecule comprises a protease cleavage site between the second target protein and the tag. In some embodiments, the protease cleavage site is one known in the art or described herein. In some embodiments, the method further comprises purifying the molecule by separating the aggregated protein from the first and second composition. The separating may be done by methods known in the art such as pelleting the aggregated proteins (i.e. centrifugation) or other separation techniques based on size and charge, for example. In some embodiments, the method further comprises eluting the second target protein. In some embodiments the elution is done by adding an eluting peptide or peptide that competes for the binding to the first target protein. In some embodiments, the eluting peptide is a peptide of SEQ ID NO:14 or a peptide having at least 90% sequence identity to SEQ ID NO:14. In some embodiments, the method further comprises contacting the molecule with a protease that cleaves between the second target protein and the tag.

[0038] In some embodiments, the fusion protein is capable of forming the protein aggregates in aqueous buffer with salt concentrations between about 0 and 500 mM KCl or NaCl, or with salt concentrations less than any one of or between any two of about 50, 100, 150, 200, 250, 300, 350, 400, 450, and 500 mM KCl or NaCl. In some embodiments, the fusion protein is capable of forming the protein aggregates when the fusion protein is present in an aqueous solution at a concentration of less than about 5 μM or at a concentration of less than any one of or between any two of about 5, 10, 15, 20, 25, 30, 35, 40, 45, and 50 μM. In some embodiments, the fusion protein is also capable of forming the protein aggregates when the fusion protein is present in an aqueous solution at a concentration of greater than 50 μM.

[0039] In some embodiments, the target protein (or first target protein) in a fusion protein described above is ferritin or a ferritin subunit, a fluorescent protein, an antibody, an antibody fragment, protein A, streptavidin, protein G, protein A/G, protein L, StrepTactin, anti-HA antibody (IgG1 against YPYDVPDYA (SEQ ID NO:15)), anti-cMYC antibody (IgG1 against EQKLISEEDL (SEQ ID NO:16)), anti-Glutathione S-transferrase antibody (GSTs), anti-FLAG antibody (e.g., anti-DYKDDDDK (SEQ ID NO:17) or anti-DDDDDK (SEQ ID NO:18) antibody), a monobody, or an affinity clamp. In some embodiments, the target protein is an antibody fragment (e.g., Fab or scFv), monobody, or affinity

clamp that specifically binds to cMYC, GST, FLAG, or other protein “tags” known to those of skill in the art.

[0040] In some embodiments, the target protein is ferritin. Ferritin is a protein expressed in many living organisms that stores iron and releases it in a controlled fashion. In some embodiments, the ferritin causes the protein aggregates to be paramagnetic, which allows them to be pelleted, manipulated, or removed using magnets. The ferritin in the protein aggregates described herein can be from any species. It is within the capability of a person of ordinary skill in the art to select a ferritin protein to include in the protein aggregates and to select a suitable nucleic acid encoding a ferritin protein or subunit. In some embodiments, the fusion protein comprised in the protein aggregate further comprises the ferritin. That is, the fusion protein can include the self-assembly domain, a target protein, and ferritin or a ferritin subunit. The fusion protein can also comprise only the self-assembly domain and ferritin. In some embodiments, the ferritin is comprised in a second fusion protein comprising a self-assembly domain. In some embodiments, the protein aggregate further comprises a fluorescent protein. In some embodiments, the fusion protein further comprises the fluorescent protein. In some embodiments, the fluorescent protein is comprised in a second fusion protein comprising a self-assembly domain. In some embodiments, the fluorescent protein is Clover or mRuby2. In some embodiments, the protein aggregate further comprises self-assembly domain proteins fused to other protein components. The self-assembly domain proteins can be naturally occurring proteins that contain a self-assembly domain or isolated self-assembly domain sequences. It is contemplated that in some embodiments, the protein aggregate does not contain any naturally-occurring proteins. In some embodiments, the majority of the fusion protein molecules that comprise the aggregate form direct noncovalent protein-protein interactions with other fusion protein molecules. In some embodiments, the protein-protein interactions are aromatic-aromatic, cation-aromatic, or hydrophobic interactions.

[0041] In some embodiments, a fusion protein as described above is comprised in an aqueous composition comprising an aggregate nucleating agent. In some embodiments, the aggregate nucleating agent is a thermally unstable protein that unfolds or misfolds at temperatures at or below about 40, 42, 45, or 50° C. In some embodiments, the aggregate nucleating agent is firefly luciferase.

[0042] Also disclosed is a self-assembly domain covalently conjugated to one or more other polypeptides through a non-peptide bond. The self-assembly domain and the other polypeptide can be separately expressed and then conjugated together through chemical cross-linking means, which are known to persons of skill in the art. The resulting molecule can be used to form protein aggregates according to the methods described herein.

[0043] In some embodiments, any of the fusion proteins described above can comprise a second, third, fourth, or fifth target protein or more.

[0044] In some embodiments, it is contemplated that a fusion protein consists entirely of a contiguous string of amino acids. In some embodiments, the fusion protein does not have any additional chemical entity joined to the amino acid string. It is also contemplated that the fusion protein can consist of only a self-assembly domain and a target protein as a single, contiguous amino acid string.

[0045] In some embodiments, a fusion protein described above can be conjugated to another polypeptide or other molecule through peptide or non-peptide covalent bonds. The self-assembly domain itself can also be conjugated to other polypeptides or other types of molecules through peptide or non-peptide covalent bonds. For example, in some embodiments, the self-assembly domain can be conjugated to biotin. In some embodiments, the self-assembly domain is not part of a fusion protein with a target protein, but is covalently conjugated to another molecule. The other molecule can include other polypeptides, small molecules, nucleic acids, or other types of molecules.

[0046] In some embodiments, the invention comprises a method of delivering a substance to a specific body site, comprising, (a) providing a fusion protein comprising a target protein capable of binding the substance fused to a heat-inducible self-assembly domain, (b) adding the substance to the fusion protein, (c) administering the fusion protein and substance to a patient, and (d) locally heating the body site. In specific embodiments, the substance is a nucleic acid, a protein or peptide-based therapeutic.

[0047] In some embodiments, the invention comprises a method of assessing modulators of aggregation *in vitro* by (a) providing a heat-inducible self-assembly domain in an aqueous solution, (b) adding a substance, (c) administering heat to the sample, and (d) measuring the degree of aggregation of the domain.

[0048] In some embodiments, the invention comprises a method of assessing modulators of aggregation *in vitro* by (a) providing a heat-inducible self-assembly domain fused to a protein in an aqueous solution, (b) adding a substance, (c) administering heat to the sample (d) measuring the degree of aggregation of the fusion protein. In specific embodiments, the protein is a fluorescent protein. In other specific embodiments, a mixture of fusion proteins is used where each protein has a heat-inducible self-assembly domain and a fluorescent protein that can interact with other fusion proteins in the mixture to provide FRET mediated fluorescence upon assembly.

[0049] In some embodiments, the invention comprises a method of assessing modulators of aggregation *in vivo* by (a) expressing a heat-inducible self-assembly domain fused to a protein in a cell, (b) administering a substance to the cell, (c) administering heat to the sample, (d) measuring the degree of aggregation of the fusion protein. In specific embodiments, the domain is a fluorescent protein. In other specific embodiments a mixture of fusion proteins is expressed where each protein has a heat-inducible self-assembly domain and a fluorescent protein that can interact with other fusion proteins in the mixture to provide FRET mediated fluorescence upon assembly.

[0050] It is contemplated that any embodiment described herein can be combined with any other described embodiment. For example, the features described for protein aggregates or fusion proteins in one embodiment can be applied to the protein aggregate or fusion proteins of any other embodiment. Likewise, any method steps described in a given method can be included in any other described method, and any method can incorporate or use any fusion protein or protein aggregate described herein.

[0051] As used herein, a “self-assembly domain” is a polypeptide sequence that imparts to a polypeptide, of which it is a part of, the ability to form protein aggregates under certain conditions. In some embodiments, the self-assembly

domain is heat-inducible; that is, a protein that includes a heat-inducible self-assembly domain is soluble in aqueous solution at relatively low temperatures (e.g., below about 35° C.) but assembles into aggregates with other proteins that include a heat-inducible self-assembly domain upon heating to a higher temperature (e.g., at least about 35° C.).

[0052] As used herein, “target protein” means a polypeptide that is distinct from a self-assembly domain. The term “target protein” excludes the amino acids that make up the self-assembly domain itself. The term “target protein” also excludes polypeptides that naturally possess a self-assembly domain, such as some GST-like proteins from *Saccharomyces cerevisiae* or *Ogataea parapolymorpha*.

[0053] As used herein, a “fusion protein” is a single, contiguous polypeptide molecule that comprises two or more distinct amino acid sequences derived from at least two distinct sources. In some embodiments, a distinct source can be a naturally-occurring gene product sequence, a man-made polypeptide sequence, or fragments of either. In some embodiments, each distinct amino acid sequence included in a fusion protein has at least or at most 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 25, 30, 35, 40, 45, 50, 60, 70, 80, 90, or 100 amino acids or any derivable range therein. In some embodiments, each distinct amino acid sequence has a distinct function that is associated with its source and includes enough of the amino acid sequence from the source to impart that functionality to the fusion protein. For example, in some embodiments, a fusion protein includes an amino acid sequence derived from a self-assembly domain and an amino acid sequence derived from a green fluorescent protein. In such a fusion protein, the self-assembly domain amino acid sequence has self-assembly functionality and the green fluorescent protein amino acid sequence has fluorescence functionality. The functionality imparted to the fusion protein by the distinct amino acid sequence may also be, for example, binding to a specific protein, small molecule, or ligand; performing a structural role; undergoing a conformation change under certain conditions; performing an enzymatic function such as catalyzing a chemical reaction; fluorescing under certain conditions; and so forth. In some embodiments described herein, the fusion proteins comprise distinct amino acid sequences from more than two distinct sources. A fusion protein can include at least, at most, or exactly 2, 3, 4, 5, 6, 7, or more distinct proteins or polypeptides (or any derivable range therein). The self-assembly domain in the fusion proteins described herein may be derived from a naturally-occurring protein sequence or may be artificial. As used herein, “fusion protein” does not include a polypeptide that is wholly derived from a single, naturally occurring gene product. As used herein, “fusion proteins” are not naturally-occurring.

[0054] The terms “a” and “an” are defined as one or more unless this disclosure explicitly requires otherwise.

[0055] The term “substantially” is defined as being largely but not necessarily wholly what is specified (and include wholly what is specified) as understood by one of ordinary skill in the art. In any disclosed embodiment, the term “substantially” may be substituted with “within [a percentage] of” what is specified, where the percentage includes 0.1, 1, 5, and 10 percent.

[0056] The terms “comprise” (and any form of comprise, such as “comprises” and “comprising”), “have” (and any form of have, such as “has” and “having”), “include” (and any form of include, such as “includes” and “including”) and

“contain” (and any form of contain, such as “contains” and “containing”) are open-ended linking verbs. As a result, the methods and systems of the present invention that “comprises,” “has,” “includes” or “contains” one or more elements possesses those one or more elements, but is not limited to possessing only those one or more elements. Likewise, an element of a method or system of the present invention that “comprises,” “has,” “includes” or “contains” one or more features possesses those one or more features, but is not limited to possessing only those one or more features.

[0057] Throughout this application, the term “about” is used to indicate that a value includes the standard deviation of error for the device or method being employed to determine the value.

[0058] Furthermore, a structure that is capable performing a function or that is configured in a certain way is capable or configured in at least that way, but may also be capable or configured in ways that are not listed. Metric units may be derived from the English units provided by applying a conversion and rounding to the nearest millimeter.

[0059] The feature or features of one embodiment may be applied to other embodiments, even though not described or illustrated, unless expressly prohibited by this disclosure or the nature of the embodiments.

[0060] Any method or system of the present invention can consist of or consist essentially of rather than comprise/include/contain/have any of the described elements and/or features and/or steps. Thus, in any of the claims, the term “consisting of” or “consisting essentially of” can be substituted for any of the open-ended linking verbs recited above, in order to change the scope of a given claim from what it would otherwise be using the open-ended linking verb.

[0061] Details associated with the embodiments described above and others are presented below.

BRIEF DESCRIPTION OF THE DRAWINGS

[0062] FIG. 1. Overview of purification method. Recombinant target protein (oval) linked to a temperature-sensitive glutathione-S-transferase (GST)-like polypeptide tag (tsG tag) (square) by a protease-cleavable linker (triangle) is expressed in host cells, which are lysed to release the protein and many contaminants. Soluble lysate is incubated for 10 minutes at 50° C. and centrifuged at 20,000 g. The pellet is resuspended and incubated with a tsG-tagged protease, which cleaves the linker and liberates the target protein. After another 10 minutes 50° C. incubation and centrifugation, the purified, tag-free protein is recovered from the supernatant.

[0063] FIG. 2. Purification of target protein mRuby2 (red fluorescent protein) from crude *E. coli* lysate, illustrated on an SDS-PAGE gel developed with Coomassie stain. Here, the protease is tsG-TEV, tobacco etch virus protease, which cleaves the seven-amino-acid sequence ENLYFQS (SEQ ID NO:19) between Q and S, leaving a single serine residue attached to the target protein. The target protein is mRuby2, a red fluorescent protein variant. Lane 1, *E. coli* lysate supernatant after centrifuging out cell debris. Lanes 2-5, total (T) protein, supernatant (S), wash (W), and pellet (P) material after first 50° C./10 min treatment and centrifugation at 17,000 g for 10 minutes. Lanes 6-8: supernatant, wash, and pellet material after 2 h incubation with added

tsG-TEV protease and second 50° C./10 min treatment. Lane 6 contains substantially pure mRuby2 of the expected molecular weight.

[0064] FIG. 3. Detailed description of the purification scheme described in FIG. 1. A, Protein components. The target protein is expressed as a fusion protein with a cleavable temperature-sensitive GST-like tag (tsG tag). A protease capable of cleaving the tag is purified once by standard affinity methods. B, Details of the method. Soluble lysate containing the tagged target protein is passed through a set of steps to generate soluble, tag-free target protein. No affinity chromatography is employed. The method can be completed in under three hours and requires only a benchtop centrifuge.

[0065] FIG. 4A-4B. FIG. 4A shows the purification of the fluorescent protein, Clover. FIG. 4B compares purification of six different proteins (Clover, mRuby2, hGH, Suil, β -Gal, Pab1) using the current method with purification using the traditional His-tag method. The top bar graph of FIG. 4B shows the purity achieved for each protein using the FENEX purification method described herein (left bar in each pair) and the His-tag purification method (right bar in each pair). The bottom bar graph of FIG. 4B shows the yield (in mg of protein per liter of culture) achieved for each protein using the FENEX purification method (left bar in each pair) and the His-tag purification method (right bar in each pair). The purification method described herein is twice as fast as His-tag purification, allows for simultaneous purification of many proteins (high-throughput screens) and is inexpensive.

[0066] FIG. 6. Recombinant yeast proteins rapidly and autonomously self-assemble into large particles in vitro, recapitulating in vivo TRAP formation. A, Sizing gel of purified proteins. B, Thermally-triggered self-assembly of purified proteins monitored by absorbance; temperature change at $t=0$. C, Comparison of in vivo and in vitro results after two minutes at the indicated temperature.

[0067] FIG. 7. GST-like domains likely mediate thermal assembly in the AME complex and in the eEF-1B complex. A, Domain architecture. Rectangles show GST-like domains, circles show GST-N(thioredoxin-like) subdomains, and squares show GST-C subdomains. B, Ternary complex of AME GST-like interaction domains. C, Rapid response of proteins in A to thermal shift monitored by mass-spectrometric analysis.

[0068] FIG. 8 shows that purified AME complex (complex of three proteins: aminoacylation cofactor (Arc1), methionyl-tRNA synthetase (Mes1), and glutamyl-tRNA synthetase (Gus1)) forms large aggregates upon heat shock. Shown on the left panel is the elution profile from Superdex200 gel filtration column. Above the elution profile is a western blot demonstrating the trimeric complex eluted in fraction 12. This demonstrates reconstitution of a stoichiometric three-protein complex. Shown on the right panel is a dynamic light scattering plot which demonstrates the size distribution profile of the individual proteins in solution and of the AME complex at the indicated temperatures. Gus1, and Mes1 rapidly form large assemblies when heated (dynamic light scattering data), whereas Arc1 shows only modest assembly. Suil, another yeast protein, shows no assembly, as a negative control.

[0069] FIG. 9 shows that virtually all of the AME complex is assembled after incubation for 15 minutes at 46° C. Shown is the elution profile from Superose 6 size exclusion chromatography of soluble (non-pelletable) material. At 15

minutes, 46° C., the majority of the protein elutes in 1 mL volume, indicating assembly of large AME complexes. The tallest peak (the left-most peak) is soluble assembly. The second tallest peak shows AME incubated at room temperature (25° C.) for comparison.

[0070] FIG. 10 shows that AME remains functional and possesses normal fidelity after heat shock at 46° C. for 15 min. Aminoacylation of tRNA^{Met} with ³⁵S methionine is used as a functional readout of AME activity. Top left, kinetic assay comparing equimolar amounts of unheated AME (top, straight line), heated AME (middle line), and unheated Mes1 alone (lower line) reveals that heated AME has slower kinetics relative to unheated AME, yet retains higher activity than unheated Mes1 alone. Bars show results of endpoint activity assay. Right, heated AME forms large complexes which pellet after centrifugation, and silver staining reveals that the three AME components retain 1:1:1 stoichiometry in the pellet, suggesting the complex remains intact. Bottom, tRNA microarray-based misacylation assay (cf. Netzger et al. (2009), "Innate immune and chemically triggered oxidative stress modifies translational fidelity," *Nature* 462:522-526). The differences between Mes1, AME, and heat-shocked AME methionine acylation patterns are minimal, indicating minimal perturbation of fidelity and again demonstrating activity of heat-shocked AME.

[0071] FIG. 11 shows that Gus1N (N-terminus of Gus1; also called the tsG domain) retains significant structure at assembly temperatures (50° C.). Data show circular dichroism spectra at 30° C. (lines with diamond points) and 50° C. (lines with circle points). Top, full-length Gus1 remains folded and shows a temperature-dependent conformational change. Middle: Gus1 Δ N, lacking N-terminal GST-like domain, shows almost no change in response to temperature. Bottom, Gus1N, the isolated GST-like domain, is very well-structured (gray line shows full-length Gus1 for comparison), and displays a substantial temperature-dependent conformational change. Compare near-complete loss of structure with 6M Gdn (guanidinium HCl) and 95° C. heating, which denature Gus1N.

[0072] FIG. 12 shows that assembly of Gus1 is domain-specific. Shown on the left is a static light scattering plot showing that Gus1 Δ N mutant, which lacks the N terminal GST-like domain, does not assemble into large complexes after a temperature shift from 30 to 50° C. In contrast, Gus1N (a polypeptide consisting only of the the N-terminal GST-like domain) assembles into large aggregates rapidly (about 30-60 seconds) after a 30 to 50° C. temperature shift.

[0073] FIG. 13 shows that Yef3 rapidly forms large particles in response to a temperature shift from room temperature to 50° C. (solid line), 46° C. (dashed), and 42° C. (dot-dash), but remains unassembled at 30° C. (dotted).

[0074] FIG. 14A-B shows the Gus1N self-assembling domain. Shown in A is a scheme which depicts the self-assembly of the Gus1N domains. Shown in B is the Gus1N-affinity domain fusion protein.

[0075] FIG. 15 depicts a purification scheme using the methods of the disclosure. This purification scheme is further described in Example 3. Briefly, the scheme depicts the steps: (1) heat-shock Gus1N-PDZ for 10 minutes at 48° C.; (2) pellet and mix with myosin V(MV)-containing lysate; (3) incubate together to let MV bind to PDZ; (4) centrifuge and discard supernatant (contaminants); (5) add the elution peptide which outcompetes c-tagged MV and releases it

from PDZ; (6) centrifuge, product is purified MV in the supernatant and Gus1N-PDZ in the pellet.

[0076] FIG. 16A-B shows Gus1N-PDZ expression (A) and purification (B) using a Ni column. Gus1N-PDZ protein expresses very well under standard conditions, namely OD=0.6, 1 mM IPTG at 30° C. and a good level of expression is achieved after about 4 hours. Only one-step purification was required to purify the protein. A Ni column was used for the purification, and the average yield from two separate purifications is 17.53 mg/L of cell culture.

[0077] FIG. 17A-B shows the specificity of the Gus1N-PDZ for the target. Shown in A is a cartoon depiction of the specificity assay. Heat-shocked Gus1N-PDZ was incubated with either Clover-C or tagless Clover, the pellets were washed, and then the pellets were visualized by using both UV and GFP channels. Shown in B are the results that demonstrate that Gus1N-PDZ binds specifically to clover-Ctag and not untagged clover (second and third panel showing UV and GFP results). The only pellet to emit any light is the one incubated with tagged Clover.

[0078] FIG. 18 demonstrates that Gus1N-PDZ binds its targets specifically. This figure is similar to FIG. 17, however, elution was performed using the elution peptide in this experiment. Each sample was split into three fractions which are resin after elution, supernatant (fraction not bound to Gus1NPDZ), and eluted fractions. As can be seen in the figure, nothing eluted in case of tagless Clover while in the case of C-tagged Clover there is got eluted Clover-C in the eluted fraction. Comparing the lanes for Gus1N-PDZ—elution, Gus1N-PDZ+Clover—elution, and Gus1N-PDZ+Clover-Ctag—elution, only the last elution lane had the Clover protein. Therefore, the Gus1N-PDZ provides for a resin with little or no background contamination. These results are consistent with previous results (e.g., FIG. 17). The Gus1N-PDZ used was 583 µg, the Clover or Clover-Ctag used was 100 µL of 30 µM. The peptide used was 100 µL of 200 µM.

[0079] FIG. 19 shows the purification of Clover-Ctag using Gus1N-PDZ according to the methods described in Example 3. This experiment was done to determine if Gus1N-PDZ could be used to purify out a target protein from a complex lysate. Clover-C was expressed in bacteria, and the lysate was pre-shocked with Gus1N-PDZ. Following this, the elution was performed. The Gus1N-PDZ used was 583 µg, the Clover or Clover-Ctag lysate used was 100 µL. The peptide used was 100 µL of 200 µM. Clover-C was successfully purified from the lysate with just some residual Gus1N-PDZ.

[0080] FIG. 20 depicts the purification of Myosin V using the methods described in Example 3 compared to traditional methods using anti-FLAG resin. The Gus1N-PDZ used was 13 mg for the whole preparation. 250 mL of crude extract was used. The peptide used was 600 µL of 400 µM. GEAR (Genetically-Encoded Affinity Resin) refers to the Gus1N-PDZ resin. As can be seen in the elution lanes, MV was purified in an amount that was comparable to purification using an anti-FLAG tag method. The amount of MV purified by the Gus1N-PDZ method was 25% less, however, the degree of purity was higher, and the capacity of the resin was higher, since a large portion of uneluted MV can be seen.

[0081] FIG. 21—The elution peptide (EP) for competing Ctag-tagged protein off of Gus1N-PDZ resin may be produced using the Gus1N system. A Gus1N-tevC-EP construct

is expressed in *E. coli*, and purified using heat/centrifugation as described in Example 2. The elution peptide may then be used as in Example 3.

DETAILED DESCRIPTION OF THE INVENTION

[0082] Various features and advantageous details are explained more fully with reference to the non-limiting embodiments that are illustrated in the accompanying drawings and detailed in the following description. It should be understood, however, that the detailed description and the specific examples, while indicating embodiments of the invention, are given by way of illustration only, and not by way of limitation. Various substitutions, modifications, additions, and/or rearrangements will become apparent to those of ordinary skill in the art from this disclosure.

[0083] In the following description, numerous specific details are provided to provide a thorough understanding of the disclosed embodiments. One of ordinary skill in the relevant art will recognize, however, that the invention may be practiced without one or more of the specific details, or with other methods, components, materials, and so forth.

A. PROTEINS AND PROTEIN EXPRESSION

1. Protein Sequences

[0084] In some embodiments, fusion proteins and/or protein aggregates described herein include self-assembly domain sequences from a GST-like domain, as described herein or from the proteins set forth in the table below:

Systematic Name	Common Name
YGL105W	Arc1
YGR264C	Mes1
YGL245W	Gus1
YPL048W	Tef3
YKL081W	Tef4
YAL003W	Efb1,
YLR249W	Yef3
YBL039C	Ura7

[0085] The table above gives the systematic name from *S. cerevisiae*, but it is contemplated that the homolog from other species may be used.

[0086] In some embodiments, the fusion proteins and/or protein aggregates described herein include fluorescent proteins. One fluorescent protein that can be used is Clover, a sequence for which is set forth in GenBank accession number AFR60231, which is hereby incorporated by reference. Further fluorescent proteins are described in Lee et al. (*PLoS One* 8:367902 (2013)), which is hereby incorporated by reference.

2. Polypeptide Production

[0087] In specific embodiments, all or part of proteins described herein can also be synthesized in solution or on a solid support in accordance with conventional techniques. Various automatic synthesizers are commercially available and can be used in accordance with known protocols. See, for example, Stewart and Young, (1984); Tam et al., (1983); Merrifield, (1986); and Barany and Merrifield (1979). Alternatively, recombinant DNA technology may be employed

wherein a nucleotide sequence that encodes a peptide or polypeptide is inserted into an expression vector, transformed or transfected into an appropriate host cell and cultivated under conditions suitable for expression.

[0088] One embodiment includes the use of gene transfer to cells, including microorganisms, for the production and/or presentation of proteins. The gene for the protein of interest may be transferred into appropriate host cells followed by culture of cells under the appropriate conditions. A nucleic acid encoding virtually any polypeptide may be employed. The generation of recombinant expression vectors, and the elements included therein, can be performed by routine techniques known to those of skill in the art.

[0089] In some embodiments, fusion proteins can be expressed from a nucleotide construct that encodes the entire fusion protein. Alternatively, fusion proteins can be formed by covalently joining different proteins after they have already been produced.

3. Protein Purification or Isolation

[0090] In certain embodiments a protein or peptide or a composition comprising such a protein or peptide may be isolated or purified. Protein purification techniques are well known to those of skill in the art. These techniques involve, at one level, the homogenization and crude fractionation of the cells, tissue or organ in to polypeptide and non-polypeptide fractions. The protein or polypeptide of interest may be further purified using chromatographic and electrophoretic techniques to achieve partial or complete purification (or purification to homogeneity). Analytical methods particularly suited to the preparation of a pure peptide are ion-exchange chromatography, gel exclusion chromatography, polyacrylamide gel electrophoresis, affinity chromatography, immunoaffinity chromatography and isoelectric focusing. An example of receptor protein purification by affinity chromatography is disclosed in U.S. Pat. No. 5,206,347, the entire text of which is incorporated herein by reference. A particularly efficient method of purifying peptides is fast performance liquid chromatography (FPLC) or even high performance liquid chromatography (HPLC).

[0091] A purified protein or peptide is intended to refer to a composition, isolatable from other components, wherein the protein or peptide is purified to any degree relative to its naturally-obtainable state. An isolated or purified protein or peptide, therefore, also refers to a protein or peptide free from the environment in which it may naturally occur.

[0092] Generally, "purified" will refer to a protein or peptide composition that has been subjected to fractionation to remove various other components, and which the composition substantially retains its expressed biological activity. Where the term "substantially purified" is used, this designation will refer to a composition in which the protein or peptide forms the major component of the composition, such as constituting about 50%, about 60%, about 70%, about 80%, about 90%, or more of the proteins in the composition.

[0093] A peptide, polypeptide or protein that is "purified to homogeneity," as applied to the present invention, means that the peptide, polypeptide or protein has a level of purity where the peptide, polypeptide or protein is substantially free from other proteins and biological components. For example, a purified peptide, polypeptide or protein will often be sufficiently free of other protein components so that degradative sequencing may be performed successfully.

[0094] Various methods for quantifying the degree of purification of the protein or peptide are known to those of skill in the art in light of the present disclosure. These include, for example, determining the specific activity of an active fraction, or assessing the amount of polypeptides within a fraction by SDS/PAGE analysis. A particular method for assessing the purity of a fraction is to calculate the specific activity of the fraction, to compare it to the specific activity of the initial extract, and to thus calculate the degree of purity therein, assessed by a "-fold purification number." The actual units used to represent the amount of activity will, of course, be dependent upon the particular assay technique chosen to follow the purification, and whether or not the expressed protein or peptide exhibits a detectable activity.

[0095] To purify a desired protein, polypeptide, or peptide a natural or recombinant composition comprising at least some specific proteins, polypeptides, or peptides may be subjected to fractionation to remove various other components from the composition. Various techniques suitable for use in protein purification are well known to those of skill in the art. These include, for example, precipitation with ammonium sulfate, PEG, antibodies and the like, or by heat denaturation, followed by: centrifugation; chromatography steps such as ion exchange, gel filtration, reverse phase, hydroxylapatite and affinity chromatography; isoelectric focusing; gel electrophoresis; and combinations of these and other techniques. As is generally known in the art, it is believed that the order of conducting the various purification steps may be changed, or that certain steps may be omitted, and still result in a suitable method for the preparation of a substantially purified protein or peptide.

[0096] Another example is the purification of a specific fusion protein using a specific binding partner. Such purification methods are routine in the art. Certain aspects of the present invention provide DNA sequences for the specific proteins, and any fusion protein purification method may be practiced. However, given many DNA and proteins are known, or may be identified and amplified using the methods described herein, any purification method can now be employed.

[0097] There is no general requirement that the protein or peptide always be provided in their most purified state. Indeed, it is contemplated that less substantially purified products will have utility in certain embodiments. Partial purification may be accomplished by using fewer purification steps in combination, or by utilizing different forms of the same general purification scheme. For example, it is appreciated that cation-exchange column chromatography performed utilizing an HPLC apparatus will generally result in a greater "-fold" purification than the same technique utilizing a low pressure chromatography system. Methods exhibiting a lower degree of relative purification may have advantages in total recovery of protein product, or in maintaining the activity of an expressed protein.

[0098] Affinity chromatography is a chromatographic procedure that relies on the specific affinity between a substance to be isolated and a molecule to which it can specifically bind. This is a receptor-ligand type of interaction. The column material is synthesized by covalently coupling one of the binding partners to an insoluble matrix. The column material is then able to specifically adsorb the substance from the solution. Elution occurs by changing the conditions to those in which binding will not occur (e.g., altered pH,

ionic strength, temperature, etc.). The matrix should be a substance that itself does not adsorb molecules to any significant extent and that has a broad range of chemical, physical and thermal stability. The ligand should be coupled in such a way as to not affect its binding properties. The ligand should also provide relatively tight binding. And it should be possible to elute the substance without destroying the sample or the ligand.

B. SEQUENCE LISTING

[0099]

Systematic Name	Common Name
YGL105W	Arc1
YGR264C	Mes1
YGL245W	Gus1
YPL048W	Tef3
YKL081W	Tef4
YAL003W	Efb1,
YLR249W	Yef3
YBL039C	Ura7

[0100] *S. cerevisiae* YGL105W/Arc1:

(SEQ ID NO: 1)

MSDLVTKFESLIISKYPVSVFTKEQSAQAQWESVLKSGQIQPHLDQLNL
VLRDNTFIVSTLYPTSTDVHVFEVALPLIKDLVASSKDVKSTYTTYRHI
LRWIDYMQNLEEVSSTDKLEINHLDLDPHEVIEKKKKAPAGGAADAAK
ADEDVSKAKKQDHPRGKPDDETLKKLREEAKAKKAANAKQQQEQ
QNKAPKPKPSAIDFRVGFIQKAIKHPDADSLYVSTIDVGDEEGPRTVC
SGLVKHFPDAMQERYVVVNCNKPVNMRGKSTAMVLCGSNDKKEVFEV
EPPKDSKAGDKVFFEGFGDEAPMKQLNPKKKIWEHLQPHFTTNDGLEVI
FKDEEEKDHPVRKLTNA KGESFKVASI ANAQVR*

[0101] *S. cerevisiae* YGR264C/Mes1:

(SEQ ID NO: 2)

MSFLISFDKSKKPAHLQLANNLKI ALALEYASKNLKPEVDNDNAAMELR
NTKEPFLFDANAILRYVMDDFEGQTS DKYQFALASLQNLHYKELPQQH
VEVLTKAIENYLVELKEPLTTDLILFANVYALNSLVHSHKPELPSKV
HNAVALAKKHVPRDSSSFKNIGAVKIQADLTVPKPDSEILPKPNERNILI
TSALPYVNNVPHLGNIGSVLSADIFARYCKGRYNALFICGTDEYGTAT
ETKALEEGVTPRQLCDKYHKIHSVYKWFQIGFDYFGRTTTDKQTEIAQH
IFTKLNSNGYLEEQSMKQLYCPVHNSYLADRYVEGECPKCHYDDARGDQC
DKCGALLDPFELINPRCKLDDASPEPKYSDHIFLSLDKLESQISEWVEKA
SEEGNWSKNSKITITQSWLKDGLKPRCITRDLVWGTVPVLEKYKDKVLYVW
FDATIGYVSI TSNTYKWKQWNNPEHVSLEYQFMGKDNVPFHTVVPFSGQ
LGTEENWTMLHHLNTEYLQYENGKFSKSRGVGVFGNNAQDSGISPSVWR
YYLASVRPESSDHSFSDWDFVARNNSSELLANLGNFVNRLIKFVNAYKNGV
VPKFPDKKVSNDGLVKDINEILSNVYKEMELGHERRGLEIAMSLSARGN

-continued

QFLQENKLDNTLFSQSPEKSDAVVAVGLNIIYAVSSIIITPYMPEIGEKIN
KMLNAPALKIIDDRPHLAILEGHNINKAEYLFQRIDEKKIDEWRAKYGGQQ
V*

[0102] *S. cerevisiae* YGL245W/Gus1:

(SEQ ID NO: 3)

MPSTLTINGKAPIVAYAEIAARIVNALAPNSIAIKLVDDKKAPAKLDD
ATEDVFNKITSKFAATPDNGDKEQVAKVWNLAQKELVIKNFAKLSQSLET
LDSQLNLR TPTLGGLYSAADVACWGALRSNGMCGSIIKNKVDVNSRWY
TLLEMDPI FGEAHDPLSKSLLELKKSANVGGKKETHKANFEIDLPAKMG
EVTTRFPPEPSGYLHIGHAKAALLNQYFAQAYKGLIIRPDDTNP SKEKE
EFQDSILEDDLGIKGRITYSSDYFQEMYDYCVQMIKDGKAYCDDTPT
EKMREERMDGVASARRDRSVEENLRIFTEEMKNGTEEGLKN CVRAKIDYK
ALNKTLRDPVIYRCNLT PHHRTGSTWKIYPTYDFCVPIVDAIEGVTHALR
TIEYDRNAQYDWMLQALRLRKVHIWDFARINFVRTLLSKRKLQWVMDKD
LVGNWDDPRFPPTVRGVRRRGMTVEGLRNFVLSQGPSRNVINLEWNLIAWAF
NKKVIDPIAPRHTAIVNPVKIHLGSEAPQEPKIEMKPKHKKNPAVGEKK
VIYYKDIVVDKDDADVINVDEEVTLMDWGNVITTKNNDGSMVAKLNLEG
DFKKTKHKLTLWADTKDVPVLDLDFDHLITKDRLEEDESFEFLTPQTE
FHTDAIADLNVKDMKIGDIIQFERKGYRLDALPKDGKPYVFFTTIPDGKS
VNKYGAKK*

[0103] *S. cerevisiae* YPL048W/Tef3:

(SEQ ID NO: 4)

MSQGTLYANFRIRTWVPRGLVKALKLDVKKVTPDAAAQFARDFPLKKVP
AFVGPKGKYTEAMAINYYLVKLSQDDKMTQQLGADDDLNAQAQIIRWQ
SLANSDLCIQIANTIVPLKGGAPYNKKSVD SAMDAVDKIVDIFENRLKNY
TYLATENISLADLVAASIFTRYFESLFGTEWRAQHPAIVRW FNTVRSFP
LKDEYKDFKADKPLSPQKKKKEKAPAAAPAASKKKEEAKPAATETETS
SKKPKHPLLELLGKSTFVLDDWKRKYSNEDTRPVALPWFWEHYNPEEYSLW
KVTYKYNDELTLTFMSNNLVGGFFNRLSASTKYMFGCLVVYGENNNNGIV
GAVMVRGQDYVPAFVAPDWESYDYLKLDPTNDDDEKFINNMWAWDKPVS
VNGEPKEIVDGKVLK*

[0104] *S. cerevisiae* YKL081W/Tef4:

(SEQ ID NO: 5)

MSQGTLYINRSPRNYASEALI SYFKLDVKIVDLEQSSEFASLFPLKQAPA
FLGPKGLKLEALAIQFYLANQVADEKERARLLGSDVIEKSQLRWASLA
NSDMSNIARPFPSFKGLI PYNKKDVIDACFVKIDNLAAVDFARLDRDYTFV
ATENISLGDHLAAGSWAFGLATILGPEWRAKPHLMRW FNTVAASPIVKT
PFAEVKLAEKALTYTPPKKQKAEKPKAEKSKAEKKKDEAKPADDAAPAK

-continued

PKHPLEALGKSTFVLDDWKRKYSNDDTRPVALPWFEHYNPEEYSIWKVG
 YKYNDELTLTPMSNNLVGGFFNRLSASTKYMFGCLVVYGENNNGIVGAV
 MVRGQDFAPAPDVPDWESYEYTKLDPTKEEDKEFVNNMWAWDKPVVVNG
 EDKEIVDGKVLK*

[0105] *S. cerevisiae* YAL003W/Efb1:

(SEQ ID NO: 6)

MASTDFSKIETLKQLNASLADKSYIEGTAVSQADVTFKAFQSAYPEFSR
 WFNHIASKADEPDSPPAASAAAAEEEEEDDDVDFGSDDEEADAEAEKKA
 ERIAAYNAKKAAPAKPAKSAKIVTLDVKPWDEETNLEEMVANVKAIEMEG
 LTWGAHQFIPIGFICKLQINCVDVDDKVSLLDQOSIEEDEDHVQSTDI
 AAMQKL*

[0106] *S. cerevisiae* YLR249W/Yef3:

(SEQ ID NO: 7)

MSDSQQSIKVLEELFQKLSVATADNRHEIASVASFNLNGNIEHDVPEHF
 FGELAKGIKDKKTAANAMQAVAHIANQSNLSPSVEPYIVQLVPAICTNAG
 NKDKEIQSVAETLISIVNAVNPVAIKALLPHLTNAIVETNKWQEKIAIL
 AAISAMVDAAKDQVALRMPPELIPVLSETMWDTKKEVKAATAAMTKATET
 VNDKIERFIPSLIQCIADPTEVPETVHLLGATTFVAEVT PATLSIMVPL
 LSRGLNERETGIRKRSVAVIDNMCKLVEDPQVIAPFLGKLLPGLKSNFAT
 IADPEAREVTLRALKTLRRVGNVGEDDAIPEVSHAGDVSTTLQVVNELLK
 DETVAPRFKIVVEYIAAIGADLIDERIIDQQAWFTHITPYMTIFLHEKKA
 KDILDEFKRRAVDNIPVGNPFDDDEGEDLCNCEFSLAYGAKILLNKTO
 LRLKRARRYGICGPNCGKSTLMRAIANGQVDGFPTQEBCRTVYVEHDID
 GTHSDTSLDFVFESEGVGTKEAIKDKLIEFGFTDEMIAMPISALSGGWKM
 KLALARAVLRNADILLLDEPTNHLDTVNVAVLWVNYLNTCGTISITISHDS
 VFLDNVCEYIINYEGLKLRKYKGNPTEPVKCKPAKAYEELSNTDLEPKF
 PEPGYLEGVTKQKAIKVTNMEFQYPGTSKPQITDINFQCSLSSRIAVI
 GPNAGKSTLINVLGTGELLPTSGEVYTHENCRIAYIKQHAFAHIESHLDK
 TPSEYIQWRFQTGEDRETMDRANRQINENDAEAMNKIFKIEGTPPRIAGI
 HSRRKFKNTYIEYECFLLGENIGMKSERWVPMMSVDNAWIPRGELVESHS
 KMVAEVDMKEALASGQFRPLRKEIEEHCSMLGLDPEIVSHSRIRGLSGG
 QKVKLVLAAGTWQRPHLIVLDEPTNYLDRDSLGAISKALKEFEGGVIIT
 HSAEFTKNLTEEVAVKDGRMTPSGHNWVSGQAGPRIEKKEDEEDKFDA
 MGNKIAGGKKKKLLSSAELRKKKKERMKKKELGDAYVSSDEEF*

[0107] *S. cerevisiae* YBL039C/Ura7:

(SEQ ID NO: 8)

MKYVVVSGGVISIGIKGVLASSTGMLMKTGLKVTSTIKIDPYMNDAGTM
 SPLEHGECFVLDDGGETDLDLGNERYLVGVTLTKDHNIITGKIYSHVIAK

-continued

ERKGDYLGKTVQIVPHLTNAIQDWIERVAKIPVDDTGMEPDVCIIEELGGT
 VGDIESAPFVEALRQFQFKVGKFNALIHVSLVPIHGEQKTKPTQAAIK
 GLRSLGLVPDMIACRCSSETLDKPTIDKIAMPCHVGPEQVNVHVDVNSTYH
 VPLLLLEQKIMIDYLHARLKLDEISLTHEEKQRGLLELSKWKATTGNPFDES
 METVKIALVKGKYNLKD SYLSVIKALEHSSMKCRKLDIKWVEATDLEPE
 AQESNKTKEHEAWNMVSTADGILIPGGFGVVRGTEGMVLAARWARENHIPF
 LGVCLGLQIATIEPTRSVLGRKDSHSAEFYDIDEKNHVVFMPPEIDKET
 MGGSMRLGLRPTFFQNETEWSQIKKLYGDVSEVHERHRHRYEINPKMVDE
 LENNGLIFVGGKDDTGKRCEILELKNHPYIATQYHPEYTSKVLDPKPPFL
 GLVAASAGILQDVIEGKYDLEAGENKPNF*

[0108] *S. cerevisiae* YGL245W/Gus1; 191Aa N-Terminal Polypeptide with GST-Like Domain:

(SEQ ID NO: 9)

MPSTLTINGKAPIVAYAEILIAARIVNALAPNSIAIKLVDDKKAPAKLDD
 ATEDVFNKITSKFAAIFDNGDKEQVAKWVNLAQKELVIKNFAKLSQSLET
 LDSQLNLRFTFILGGLKYSAADVACWGALRSNGMCGSIIKNKVDVNSRWY
 TLLEMDPIFGEAHDFLSKSLELKKSANVGGKKEETHKANFE

[0109] *S. cerevisiae* YGR264C/Mes1; 207Aa N-Terminal Polypeptide with GST-Like Domain:

(SEQ ID NO: 10)

MSFLISFDKSKKHPAHLQLANNLKIALALEYAKNLKPEVDNDNAAMELR
 NTKEPFLFLFDANAILRYVMDDFEGQTSKDYQFALASLQNLNLYHKELPQQH
 VEVLTKNAIENYVELKEPLTTDLILFANVYALNSSLVHSKFPPELPSKV
 HNAVALAKKHVPRDSSPKNI GAVKI QADLTVKPKDSEILPKPNERNILI
 TSALPYV

[0110] *S. cerevisiae* YKL081W/Tef4; 156Aa N-Terminal Polypeptide with GST-Like Domain:

(SEQ ID NO: 11)

MSQGTLYINRSPRNYASEALI SYFKLDVKIVDLEQSSEFASLFPLKQAPA
 FLGPKGLKLEALAIQFYLANQVADEKERARLLGSDVIEKSQLRWASLA
 NSDVMSNIARPFLSPKGLIPYNKKDVDACFVKIDNLAAVFDARLRDYTFV
 ATENIS

[0111] *S. cerevisiae* YPL048W/Tef3; 159Aa N-Terminal Polypeptide with GST-Like Domain:

(SEQ ID NO: 12)

MSQGTLYANFRIRTWVPRGLVKALKLDVKVTPDAAAEQFARDPFLKPKVP
 AFVGPVKGYKLTTEAMAINYYLVKLSQDDKMKTOQLLGADDDLNAQAQIIRWQ

-continued

Example 1

SLANSDLICIQIANTIVPLKGGAPYNKKSVDSDAMDAVDKIVDIFENRLKNY

Self-Assembly Domains Induced by Heat to Form Protein Aggregates

TYLATENIS

RGSIDTWV. (SEQ ID NO: 13)

[0112] C-Tag:

EEWETWV. (SEQ ID NO: 14)

[0113] Elution Peptide:**[0114]** Exemplary tags:

YPYDVPDYA; (SEQ ID NO: 15)

EQKLISEEDL; (SEQ ID NO: 16)

DYKDDDDK and (SEQ ID NO: 17)

DDDDK. (SEQ ID NO: 18)

[0115] TEV Cleavage Site:

ENLYFQS (SEQ ID NO: 19)

C. EXAMPLES

[0116] The present invention will be described in greater detail by way of specific examples. The following examples are offered for illustrative purposes only, and are not intended to limit the invention in any manner. Those of skill in the art will readily recognize a variety of noncritical parameters which can be changed or modified to yield essentially the same results.

[0117] Following a sudden increase in temperature, cells attenuate protein synthesis and mount the heat-shock transcriptional program, and eukaryotic cells additionally sequester proteins and RNA in stress granules. How cells sense temperature remains unclear. Here, using a novel mass spectrometric method to identify protein aggregation at the proteome scale in budding yeast, Applicants show that within two minutes of a rise in temperature, a limited set of soluble proteins assemble in vivo into large particles which are molecularly distinct from stress granules. Remarkably, Applicants find that assembly is protein-autonomous: recombinant, purified proteins self-assemble in vitro with comparable kinetics in response to an equivalent thermal shift. For glutamyl-tRNA synthetase, autonomous thermal self-assembly occurs between stably folded proteins and reflects temperature-dependent conformational changes in specific protein-protein interaction domains. Applicants propose that a distributed system of sensor domains transduce temperature into autonomous protein assembly to effect rapid adjustment of diffusible protein levels without transcription, translation, or protein modifications.

[0118] Thermally-induced protein misfolding and aggregation have long been thought to trigger the heat-shock response, but the sensitivity of individual proteins to thermal aggregation has remained unclear. To examine changes in protein aggregation in response to heat stress, Applicants used a proteome-scale mass spectrometric (MS) assay to monitor the ratio of proteins found in the supernatant (aqueous-soluble) and pellet (aqueous-insoluble, detergent-soluble) fractions. To maximize effect sizes, Applicants targeted brief treatments after which only a fraction of cells survive, shifting exponentially growing cells from 30° C. to 50° C. for two, four, and eight minutes. Applicants immediately harvested supernatant and 100,000 g pellet fractions from each, combined the supernatant fraction with the pellet fraction from cells grown on stable-isotope-labeled arginine and lysine, then analyzed the mixed samples by liquid chromatography coupled to tandem mass spectrometry (LC-MS/MS). The data reveal that after two minutes, while the vast majority of proteins showed no significant aggregation, a small set of highly soluble proteins form pelletable aggregates (FIG. 7C), with 31 of 597 protein (5%) increasing at least four-fold in the insoluble fraction. Little change was observed at later times, with Pearson correlations ≥ 0.85 between supernatant/pellet ratios at 2, 4, and 8 minutes.

[0119] These observations suggested a functional connection between the aggregating proteins, consistent with a thermally triggered assembly process. Heat-induced stress granules were originally identified by a similar centrifugation procedure. Indeed, most components of stress granules induced by robust heat shock, previously studied by fluorescent imaging, are found in these rapidly assembling particles, with the exception of one protein, Dhh1, primarily associated with mRNA processing bodies (P bodies), and one small-subunit ribosomal protein. Stress granules are thought to coalesce around stalled 48S preinitiation complexes, in which the 40S small ribosomal subunit is a core component. Small-subunit proteins are universal markers for stress granules and the yeast ribosomal protein S30, but not large-subunit L25, accumulates in heat-induced granules. However, all of the 131 detected ribosomal gene products, including 56 from the small subunit, remained strongly enriched in the supernatant and were entirely separable from the aggregating proteins at 2 min. Additional initiation-complex components eIF-1A, eIF-2 (a, β , and γ subunits), and eIF-1 also remained soluble. These and further results reported below indicate that these rapidly forming particles do not co-assemble with preinitiation complexes or small ribosomal subunits, distinguishing them from stress granules. The processing-body (P-body) markers Dcp2p also remained in the supernatant, distinguishing these particles from P-bodies. Applicants therefore designated them thermosensitive rapidly assembling particles (TRAPs).

[0120] The rapid formation of TRAPs, and their independence from preinitiation complexes led Applicants to wonder whether the constituent proteins themselves possessed the intrinsic ability to transduce a thermal shift into self-assembly. To test this possibility, Applicants purified several TRAP-forming proteins in recombinant form from bacteria (Yef3, Gus1, and CTP synthase/Ura7), along with two control proteins (Sui1/eIF-1 and Hyp2/eIF-5A). Applicants

suspended them in aqueous buffer at concentrations approximating their physiological levels, and monitored formation of large particles by visible-light absorbance at 550 nm. When Applicants subjected each protein to the same 30° C. to 50° C. thermal shift *in vitro*, all TRAP-forming proteins rapidly self-assembled into large particles, and the control proteins did not. Particles grew exponentially until exhaustion of unassembled material (FIG. 6B).

[0121] Self-assembly *in vitro* approached saturation in two minutes, consistent with *in vivo* behavior (FIG. 6C). Applicants then asked whether lower temperatures triggered assembly. Yef3 forms granules at 42° C. *in vivo* which seed subsequent formation of genuine stress granules. Correspondingly, Yef3 self-assembled rapidly at 46° C. and 42° C. *in vitro* with temperature-dependent kinetics (FIG. 13).

[0122] FIG. 8 shows that purified AME complex (complex of three proteins: aminoacylation cofactor (Arc1), methionyl-tRNA synthetase (Mes1), and glutamyl-tRNA synthetase (Gus1)) forms large aggregates upon heat shock. FIG. 9 shows that virtually all of the AME complex is assembled after incubation for 15 minutes at 46° C. FIG. 10 shows AME complexes under electron microscopy. The AME assemblies increase in abundance after 15 min. 46° C., but are absent at 15 minutes, 30° C. FIG. 10 shows that AME remains functional and possesses normal fidelity after heat shock at 46° C. for 15 min.

[0123] Applicant's data and previous work indicate that thermal self-assembly *in vitro* recapitulates assembly *in vivo* with similar temperature-dependent kinetics.

[0124] Gus1 possesses a eukaryote-specific N-terminal domain, which has been previously crystallized in isolation and shown to adopt a glutathione-S-transferase-like (GST-like) fold (FIG. 12). Applicants purified this domain (Gus1N) and the remaining core synthetase domain (Gus1ΔN). Gus1ΔN did not form large particles at any tested temperature up to 50° C. (FIG. 12), and showed a largely unperturbed structure (FIG. 12). In contrast, Gus1N readily assembled *in vitro* (FIG. 12). Gus1N is therefore necessary and sufficient for thermal assembly *in vitro*.

[0125] Gus1N behaves like a thermometer, transducing a change in temperature into self-assembly. To determine the sensitivity of this phenomenon, Applicants turned to dynamic light scattering (DLS), which unlike absorbance is capable of resolving particles at the nanometer scale. DLS revealed that this GST-like domain purified Gus1N adopts a GST-like fold and mediates binding of Gus1 to the cofactor Arc1, accelerating the aminoacylation rate of Gus1. In isolation, Gus1N rapidly assembled into large particles upon temperature shift (FIG. 12). Gus1N is thus necessary and sufficient for full-length Gus1's temperature-dependent self-assembly *in vitro*.

[0126] ¹H,¹⁵N-HSQC NMR of Gus1N at 20° C. and 43° C. indicate that the environment of the amide groups in Gus1N are similar at the two temperatures suggesting that self-assembly is mediated through a small conformational change and that most of the structure remains folded during assembly.

[0127] To determine the consequences of thermal shift on protein structure, Applicants collected far-ultraviolet circular dichroism (CD) spectra. The CD spectrum of full-length Gus1 at 30° C. revealed a well-folded structure; at 50° C., the protein remained well-folded while losing some helical structure (FIG. 11). The core synthetase, Gus1ΔN, showed minimal structural change in response to the temperature

shift (FIG. 11). In marked contrast, Gus1N underwent a significant conformational change involving loss of helical structure, yet did so while preserving a highly ordered, largely α-helical structure (FIG. 11), whereas denaturation with 6M guanidine chloride (Gdn) disrupted Gus1N structure entirely (FIG. 11). These results closely match the *in vitro* assembly data, showing that at the residue and oligomer levels, the eukaryote-specific Gus1N domain is a temperature-responsive element linked to a temperature-insensitive enzyme.

[0128] Applicants speculate that the formation of large aggregates after a near-lethal heat shock results from damage to the heat-sensing system. A sensory system's necessary sensitivity to a stimulus predisposes that system to specific damage when the stimulus grows overwhelming, much as eyes are damaged by extremely bright light and ears by extremely loud sounds but not vice versa.

[0129] Repeated demonstrations that heat-shock-like responses can be generated by protein misfolding at normal growth temperatures have led to the hypothesis that heat shock is signaled and sensed by misfolded proteins. Notably, however, it has never been established that the temperature-induced misfolding of a native protein triggers the heat shock response. Applicants' results suggest the existence of an alternative channel of information provided by domain-specific thermal-shift-induced assembly of sensory proteins. These results strongly suggest that thermosensor-domain-mediated self-assembly, likely occurring in parallel in other proteins, is the first mechanistic step connecting a temperature change with stress-granule formation.

[0130] These results suggest a model in which environmental changes trigger proportionate changes in protein assembly, building up assembled proteins and reducing the population of freely diffusing proteins. Assembly may be reversible by cellular factors, or may require synthesis of new unassembled proteins. Applicants speculate that cellular chaperones, several of which are known to disaggregate misfolded proteins, also disaggregate assemblies. If so, this would suggest a remarkably simple regulatory mechanism.

Example 2: Rapid, Low-Cost Purification of Recombinant Tag-Free Proteins without Affinity Chromatography (Fenex)

[0131] A method is described for purifying a wide range of recombinant proteins employing only mild heating (<50° C.) and centrifugation (<20,000 g) achievable with standard benchtop equipment.

[0132] Purification of proteins is essential to many biological and industrial pursuits, such as characterization of protein structure and function and the development of drugs. Recombinant protein expression and purification is a common strategy, because affinity tags optimized for selective binding to columns in affinity chromatography systems can be appended to the target protein, enabling use of the same system to purify many proteins. Affinity chromatography equipment remains expensive and complex, making protein purification inaccessible to many.

[0133] Certain proteins are routinely purified without affinity chromatography. A kilogram of RNase A was famously purified by the Armour Co. by boiling bovine pancreas and centrifuging the resulting stew; RNase A is the only protein that remains in the supernatant after this treatment. Thermophilic proteins are often purified recombinantly from mesophilic hosts (such as *E. coli*) by heat-

denaturing the host lysate at temperatures intolerable to the host but tolerated by the thermophile. These examples remain rare exceptions.

[0134] Purification methods including removal of affinity tags typically require two separate purification steps. Typical generic protein purification systems require purchase or production of columns, affinity resin, and liquid-handling systems to control flow, measure properties of the flowing liquid, and collect fractions. A single separation of lysate containing affinity-tagged protein on an affinity column most often yields recombinant protein with the affinity tag still attached. Digestion with an affinity-tagged protease and a second separation is required to yield tag-free protein. The present method achieves both separation and tag removal without any affinity columns, resin, or liquid-handling systems, making it far simpler, faster, easier, and cheaper than common approaches.

[0135] A method for purifying a target protein is diagrammed in FIG. 1, and is described below. The target protein is initially tagged and expressed in an arbitrary host organism; the method produces tag-free protein.

[0136] The method exploits a temperature-sensitive GST-like polypeptide ("tsG tag") fused to two proteins: a target protein of interest, and a protease. The tsG polypeptide has the property, discovered by Applicants and as yet unreported, of rapidly self-assembling in response to increases in temperature.

[0137] One embodiment comprises sequestering the target protein away from soluble host contaminants by heat-induced tsG-tag self-assembly, releasing the target protein from heat-aggregatable host contaminants by proteolysis with a tsG-tagged protease, and finally removing heat-aggregatable host contaminants, the protease, and the cleaved tag by heat-induced tsG self-assembly.

[0138] The self-assembly of tsG results in large particles which pellet readily upon centrifugation at 10,000-20,000 g, attainable on a typical benchtop microcentrifuge.

[0139] FIG. 2 demonstrates purification of a test protein, the red fluorescent protein mRuby2, from *E. coli* using one embodiment of the method. Briefly, *E. coli* were cultured and lysed using methods known to those skilled in the art, such as mechanical or chemical lysis. The soluble lysate containing the fusion protein is heated for 10 minutes at 50 degrees Celsius and then subjected to centrifugation for 2 minutes at 17,000 g. The SDS-PAGE gel represents protein profile from samples during the process of the method. Lane 1 shows the starting material after cell lysis. Lanes 2-5, total (T) protein, supernatant (S), wash (W), and pellet (P) material after first 50° C./10 min treatment and centrifugation at 17,000 g for 10 minutes. The supernatant comprising soluble contaminants was discarded. The remaining pellet comprises insoluble contaminants and the heat-assembled Gus1N-mRuby2. Next, a second fusion protein comprising Gus1N and tobacco etch virus (TEV) protease was added to the remaining pellet, which was suspended in TEV cleavage buffer. The mixture was heated at 25 degrees Celsius for 2 hours in which the TEV protease liberated the mRuby2 from the Gus1N. The mixture was then heated for 10 minutes at 50 degrees Celsius to aggregate mRuby2-free Gus1N protein. Insoluble contaminants and Gus1N were removed from mRuby2 by centrifugation for 2 minutes at 17,000 g. The soluble mRuby2 was decanted from the pellet. Lanes 6-8 of the SDS-PAGE in FIG. 2 shows the supernatant, wash, and pellet material after 2 h incubation with added tsG-TEV

protease and second 50° C./10 min treatment. Lane 6 contains substantially pure mRuby2 of the expected molecular weight. FIG. 3 illustrates the steps involved at the molecular level. FIG. 4A shows the purification of the fluorescent protein, Clover, using the current embodiment. FIG. 4B compares purification of six different proteins (Clover, mRuby2, hGH, Suil, 13-Gal, Pab1) using the current embodiment with purification using the traditional His-tag method. Certain embodiments of the method described herein achieves comparable purity to the His-tag method.

[0140] Certain embodiments of this method allows protein purification more cheaply and rapidly, with less equipment and less effort, than any method of which Applicants are aware. They are applicable to purification of any soluble proteins, particularly for initial or high-throughput studies.

[0141] In addition to its simplicity, the method has proven to be unusually good at removing the protease and uncleaved fusion protein from the final purification. This is often a challenge for existing affinity purification schemes, which, even with a second round of purification, fail to completely remove the uncleaved protein.

[0142] In principle, a very wide range of proteins are amenable to purification by various embodiments of the invention.

[0143] Preferably, for certain embodiments, target proteins are stable at temperatures and durations necessary for assembly of the tsG domains. Reaction temperatures can be lowered by engineering of the tsG domain. In certain embodiments, however, such as the embodiment shown in FIG. 19, the target protein is not heated to the temperature necessary for assembly of the tsG domains.

[0144] Preferably, for certain embodiments, target proteins are tolerant of fusion to the tsG domain and protease cleavage site. Most affinity purification methods require fusions of some sort (alternatives are antibodies and affinity reagents designed to be specific to the protein of interest), and both the type of domain (GST-like) and the cleavage site employed in Applicants' proof-of-concept experiment are widely employed in protein purification experiments.

[0145] Preferably, the target protein is soluble under the conditions employed.

[0146] For some applications, subsequent purification steps may be used.

[0147] Engineering a wide range of proteases by fusing them to the tsG tag will enable utilization of a wide range of protease cleavage sites. Because the selectivity and activity of proteases vary, such a library of proteases would enable the purification of an increased number of protein targets.

[0148] Applicants have discovered a range of domains exhibiting temperature-triggered self-assembly. In principle, any of these can be used in place of the tsG tag demonstrated here.

[0149] The use of centrifugation to separate assemblies is not essential. The principle is separation by size, which can also be achieved by filtration.

Example 3

Genetically-Encoded Affinity Resin (Gear) for Purification of C-Tag Myosin V

[0150] Applicants have developed a system in which the Gus1N self-assembling polypeptide is genetically fused to the PDZ domain, which binds a short polypeptide called a

C-tag. The PDZ/C-tag system has been previously described (Huang et al. 2009). Using this system, Applicants have purified myosin V in its active form. Myosin V is a molecular motor protein that is responsible for intracellular cargo transportation in cells. The protein is a dimer which possesses so-called “legs” that allow it to “walk” along the actin filaments, and cargo-binding domains that bind what the myosin actually transports, for example vesicles containing RNA. The process of “walking” is driven by ATP hydrolysis. Myosin V (MV) itself is a massive protein, around 137 kDa. These features make myosin V a very difficult protein to both purify and work on. Usually, it is purified via FLAG resin, so it binds to agarose beads coated with anti-FLAG tag antibodies and then eluted with a FLAG peptide that outcompetes bound myosins. The anti-FLAG resin is very expensive, and just 10 mL of an anti-FLAG resin can cost about \$1790.

[0151] Expression of Gus1PDZ (GEAR)

[0152] Gus1PDZ expresses very well under standard conditions, namely OD=0.6, 1 mM IPTG at 30° C. and satisfactory expression is achieved after about 4 hours (FIG. 16A). One advantage of the Gus1NPDZ is that only a one-step purification is required to achieve a highly purified product. In this example, Gus1NPDZ was purified using affinity chromatography with the Ni²⁺ column. The average yield from two separate purifications is 17.53 mg/L of cell culture (FIG. 16B).

[0153] To make this process less expensive and totally lab made, Applicants created a fusion protein that would have a self-assembling domain as a resin-forming domain and an affinity domain having affinity to tagged Myo V. Gus1N was chosen as the resin-forming domain. Gus1N is an N-terminal domain of glutamyl-tRNA synthetase which causes either this enzyme or different proteins tagged with it to self-assemble upon a few minutes of heat shock. Once assembled, Gus1N-tagged molecules form complex meshworks. So now once Gus1N were linked to an affinity domain and heat-shocked it would provide lots of binding sites for the target. The domain we decided to choose to be fused to Gus1N to make GEAR (Genetically-Encoded Affinity Resin) is PDZ. The PDZ domain is a common structural domain of 80-90 amino-acids found in the signaling proteins of bacteria, yeast, plants, viruses, and animals. PDZ is an acronym combining the first letters of three proteins—post synaptic density protein (PSD95), *Drosophila* disc large tumor suppressor (Dlg1), and zonula occludens-1 protein (zo-1) which were first discovered to share the domain. PDZ domains have previously been referred to as DHR (Dlg homologous region) or GLGF (glycine-leucine-glycine-phenylalanine) domains. Engineering of its ligands allowed for the creation of a C-tag (SEQ ID NO:13). The major advantage of c-tag is that it's bound by PDZ quite tightly yet is easily releasable. And this release is possible by using the elution peptide (SEQ ID NO:14) which outcompetes proteins bound to PDZ tagged with c-tag.

[0154] The procedure for purification is depicted in FIG. 15. Briefly, the Gus1N-PDZ protein is expressed and purified from *E. coli*, then heat-shocked at 48° C. for 10 minutes to cause self-assembly, generating a solid support decorated with PDZ domains. This can be referred to as a resin. MV-Ctag is expressed in cells, which are lysed. The lysate is incubated with the resin, then centrifuged at 12,000 g for 5 minutes. The supernatant is discarded and the pellet is

washed. Then elution peptide is added. This peptide has higher affinity for PDZ than does Ctag, so the bound MV-Ctag is released from the resin. Another 12,000 g spin is performed, and the supernatant is retained. The supernatant contains highly purified MV-Ctag. In this purification method, the target protein (MV; i.e. second target protein) is never exposed to a heat shock.

[0155] Expression of Gus1PDZ

[0156] Gus1PDZ expresses very well under standard conditions, namely OD=0.6, 1 mM IPTG at 30 C and satisfactory expression is achieved after about 4 hours. One advantage of the Gus1NPDZ is that only one-step purification is required to get a highly purified product. The Ni column was used for purification of Gus1NPDZ. The average yield from two separate purifications is 17.53 mg/L of cell culture (FIG. 16B).

[0157] To make this process less expensive and totally lab made, Applicants created a fusion protein that would have a self-assembling domain as a resin-forming domain and an affinity domain having affinity to tagged Myosin V. Gus1N was chosen as the resin-forming domain. Once assembled, Gus1N-tagged molecules form complex meshworks. So now once Gus1N was linked to an affinity domain and heat-shocked it would provide a lot of binding sites for the target. Applicants fused PDZ to Gus1N to make GEAR (Genetically-Encoded Affinity Resin). The PDZ domain is a common structural domain of 80-90 amino-acids found in signaling proteins of bacteria, yeast, plants, viruses, and animals. PDZ is an acronym combining the first letters of three proteins—post-synaptic density protein (PSD95), *Drosophila* disc large tumor suppressor (Dlg1), and zonula occludens-1 protein (zo-1) which were first discovered to share the domain. PDZ domains have previously been referred to as DHR (Dlg homologous region) or GLGF (glycine-leucine-glycine-phenylalanine) domains. Engineering of the PDZ domain's natural ligand allowed for the creation of a C-tag (SEQ ID NO:13). An advantage of C-tag is that the binding to PDZ, though quite tightly, is easily releasable. This release is possible by using the elution peptide (SEQ ID NO:14) which outcompetes C-tagged proteins bound to PDZ.

[0158] The procedure for purification in one embodiment is depicted in FIG. 15. Briefly, the Gus1N-PDZ protein is expressed and purified from *E. coli*, using standard methods or FENEX. The purified Gus1N-PDZ is then heat-shocked at 48° C. for 10 minutes to cause self-assembly, generating a solid support decorated with PDZ domains. This can be referred to as a resin. MV-C-tag is expressed in cells, which are then lysed. The lysate is incubated with the resin, then centrifuged at 12,000 g for 5 minutes. The supernatant is discarded and the pellet is washed in buffered solution. Then the elution peptide is added. This peptide has a higher affinity for the PDZ domain than the C-tag, so the bound MV-Ctag is released from the resin. Another 12,000 g spin is performed, and the supernatant is retained. The supernatant contains highly purified MV-C-tag. In this purification method, the target protein (MV; i.e., a second target protein) is never exposed to a heat shock.

[0159] Purification of Myosin V Using GEAR Compared to FLAG Purification Methods

[0160] Next, GEAR purification was compared to anti-FLAG resin purification. As can be seen by comparing the elution lanes, MV was successfully purified using GEAR in an amount that was comparable to the anti-FLAG resin. The

amount of MV purified by the Gus1NPDZ method was 25% less, however, the degree of purity was higher, and the capacity of the resin was higher, since a large portion of uneluted MV can be seen in the resin fraction compared to the resin fraction using FLAG purification (FIG. 20).

[0161] To test whether the purified MV was a functional protein, a gliding filament assay was performed. In this assay, a coverslip was coated with the GEAR-purified myosins so that the legs are exposed outward, and actin filaments that interact with the myosins were added. Next, ATP-containing buffer that triggers myosin activity was added, causing the actin filaments to glide along the immobilized myosins. Most of the actin filaments moved, indicating that the purified myosins were functional.

[0162] Purification of Clover using GEAR

[0163] The binding specificity of Gus1N-PDZ was tested in two examples. In the first example, heat-shocked Gus1N-PDZ was incubated with either Clover-C-tag (Clover-C) or tagless Clover. The mixture was pelleted by centrifugation, the pellets were washed, and then visualized with both UV and GFP channels. As shown in FIG. 17, the only pellet to emit any fluorescence is the one incubated with C-tagged Clover. In the second specificity example, a similar experimental procedure is performed, but elution was achieved using the elution peptide. In FIG. 18, each sample was split into three fractions and analyzed by SDS-PAGE: (1) resin after elution, (2) supernatant or supe (the fraction not bound to Gus1NPDZ) and (3) eluted fractions (elution). In the case of tagless Clover, Clover protein was found in the supernatant fraction and nothing was eluted. C-tagged Clover yielded Clover-C in the elution fractions. Purified components diluted 1:10 were also loaded onto the gel as a control.

[0164] These results are consistent with previous results described herein using this method.

[0165] It was then determined whether Gus1N-PDZ could be used to purify out a target protein from a complex lysate. To do so, Clover-C was expressed in bacteria, the lysate was incubated with pre-heat shocked Gus1NPDZ. Next, the elution was performed (FIG. 19). Clover-C was successfully purified with only a little residual Gus1N-PDZ.

[0166] This disclosure relates to variations of the above-described purification method. In one variation, the target protein X is expressed as an X[cleavage site]Ctag, where [cleavage site] represents the recognition amino acid sequence for a protease, such as TEV. Instead of an excess of elution peptide (EP), a protease-C-tag fusion protein is added at low concentrations. The target protein is released by cleavage off of the resin, and the protease is recruited to the resin. In further variations, other affinity domains and other release peptides are used. The resin concept can be used in virtually any application where beads or other solid supports are now used, such as depletion of a target protein from a mixture. In virtually all cases, the fact that beads are spherical or separate from one another is irrelevant.

[0167] Although certain embodiments have been described above with a certain degree of particularity, or with reference to one or more individual embodiments, those skilled in the art could make numerous alterations to the disclosed embodiments without departing from the scope of this invention. As such, the illustrative embodiments are not intended to be limited to the particular forms disclosed. Rather, they include all modifications and alternatives falling within the scope of the claims, and embodiments other than those shown may include some or all of the features of

the depicted embodiment. Further, where appropriate, aspects of any of the examples described above may be combined with aspects of any of the other examples described to form further examples having comparable or different properties and addressing the same or different problems. Similarly, it will be understood that the benefits and advantages described above may relate to one embodiment or may relate to several embodiments.

[0168] The claims are not to be interpreted as including means-plus- or step-plus-function limitations, unless such a limitation is explicitly recited in a given claim using the phrase(s) “means for” or “step for,” respectively.

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Glu	Glu	Arg	Met	Asp	Gly	Val	Ala	Ser	Ala	Arg	Arg	Asp	Arg	Ser	Val
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Glu	Glu	Asn	Leu	Arg	Ile	Phe	Thr	Glu	Glu	Met	Lys	Asn	Gly	Thr	Glu
				325						330				335	
Glu	Gly	Leu	Lys	Asn	Cys	Val	Arg	Ala	Lys	Ile	Asp	Tyr	Lys	Ala	Leu
			340					345					350		
Asn	Lys	Thr	Leu	Arg	Asp	Pro	Val	Ile	Tyr	Arg	Cys	Asn	Leu	Thr	Pro
		355					360					365			
His	His	Arg	Thr	Gly	Ser	Thr	Trp	Lys	Ile	Tyr	Pro	Thr	Tyr	Asp	Phe
	370					375					380				
Cys	Val	Pro	Ile	Val	Asp	Ala	Ile	Glu	Gly	Val	Thr	His	Ala	Leu	Arg
	385				390					395					400
Thr	Ile	Glu	Tyr	Arg	Asp	Arg	Asn	Ala	Gln	Tyr	Asp	Trp	Met	Leu	Gln
				405					410					415	
Ala	Leu	Arg	Leu	Arg	Lys	Val	His	Ile	Trp	Asp	Phe	Ala	Arg	Ile	Asn
			420					425					430		
Phe	Val	Arg	Thr	Leu	Leu	Ser	Lys	Arg	Lys	Leu	Gln	Trp	Met	Val	Asp
		435					440					445			
Lys	Asp	Leu	Val	Gly	Asn	Trp	Asp	Asp	Pro	Arg	Phe	Pro	Thr	Val	Arg
	450					455					460				
Gly	Val	Arg	Arg	Arg	Gly	Met	Thr	Val	Glu	Gly	Leu	Arg	Asn	Phe	Val
	465				470					475					480
Leu	Ser	Gln	Gly	Pro	Ser	Arg	Asn	Val	Ile	Asn	Leu	Glu	Trp	Asn	Leu
				485					490					495	
Ile	Trp	Ala	Phe	Asn	Lys	Lys	Val	Ile	Asp	Pro	Ile	Ala	Pro	Arg	His
			500					505					510		
Thr	Ala	Ile	Val	Asn	Pro	Val	Lys	Ile	His	Leu	Glu	Gly	Ser	Glu	Ala
		515					520					525			
Pro	Gln	Glu	Pro	Lys	Ile	Glu	Met	Lys	Pro	Lys	His	Lys	Lys	Asn	Pro
	530					535					540				
Ala	Val	Gly	Glu	Lys	Lys	Val	Ile	Tyr	Tyr	Lys	Asp	Ile	Val	Val	Asp
	545				550					555					560
Lys	Asp	Asp	Ala	Asp	Val	Ile	Asn	Val	Asp	Glu	Glu	Val	Thr	Leu	Met
				565					570					575	
Asp	Trp	Gly	Asn	Val	Ile	Ile	Thr	Lys	Lys	Asn	Asp	Asp	Gly	Ser	Met
			580					585					590		
Val	Ala	Lys	Leu	Asn	Leu	Glu	Gly	Asp	Phe	Lys	Lys	Thr	Lys	His	Lys

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595	600	605
Leu Thr Trp Leu Ala Asp Thr Lys Asp Val Val Pro Val Asp Leu Val		
610	615	620
Asp Phe Asp His Leu Ile Thr Lys Asp Arg Leu Glu Glu Asp Glu Ser		
625	630	635
Phe Glu Asp Phe Leu Thr Pro Gln Thr Glu Phe His Thr Asp Ala Ile		
645	650	655
Ala Asp Leu Asn Val Lys Asp Met Lys Ile Gly Asp Ile Ile Gln Phe		
660	665	670
Glu Arg Lys Gly Tyr Tyr Arg Leu Asp Ala Leu Pro Lys Asp Gly Lys		
675	680	685
Pro Tyr Val Phe Phe Thr Ile Pro Asp Gly Lys Ser Val Asn Lys Tyr		
690	695	700
Gly Ala Lys Lys		
705		
<210> SEQ ID NO 4		
<211> LENGTH: 415		
<212> TYPE: PRT		
<213> ORGANISM: <i>Saccharomyces cerevisiae</i>		
<400> SEQUENCE: 4		
Met Ser Gln Gly Thr Leu Tyr Ala Asn Phe Arg Ile Arg Thr Trp Val		
1	5	10
Pro Arg Gly Leu Val Lys Ala Leu Lys Leu Asp Val Lys Val Val Thr		
20	25	30
Pro Asp Ala Ala Ala Glu Gln Phe Ala Arg Asp Phe Pro Leu Lys Lys		
35	40	45
Val Pro Ala Phe Val Gly Pro Lys Gly Tyr Lys Leu Thr Glu Ala Met		
50	55	60
Ala Ile Asn Tyr Tyr Leu Val Lys Leu Ser Gln Asp Asp Lys Met Lys		
65	70	75
Thr Gln Leu Leu Gly Ala Asp Asp Asp Leu Asn Ala Gln Ala Gln Ile		
85	90	95
Ile Arg Trp Gln Ser Leu Ala Asn Ser Asp Leu Cys Ile Gln Ile Ala		
100	105	110
Asn Thr Ile Val Pro Leu Lys Gly Gly Ala Pro Tyr Asn Lys Lys Ser		
115	120	125
Val Asp Ser Ala Met Asp Ala Val Asp Lys Ile Val Asp Ile Phe Glu		
130	135	140
Asn Arg Leu Lys Asn Tyr Thr Tyr Leu Ala Thr Glu Asn Ile Ser Leu		
145	150	155
Ala Asp Leu Val Ala Ala Ser Ile Phe Thr Arg Tyr Phe Glu Ser Leu		
165	170	175
Phe Gly Thr Glu Trp Arg Ala Gln His Pro Ala Ile Val Arg Trp Phe		
180	185	190
Asn Thr Val Arg Ala Ser Pro Phe Leu Lys Asp Glu Tyr Lys Asp Phe		
195	200	205
Lys Phe Ala Asp Lys Pro Leu Ser Pro Pro Gln Lys Lys Lys Glu Lys		
210	215	220
Lys Ala Pro Ala Ala Ala Pro Ala Ala Ser Lys Lys Lys Glu Glu Ala		
225	230	235
		240

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Lys Pro Ala Ala Thr Glu Thr Glu Thr Ser Ser Lys Lys Pro Lys His
245 250 255

Pro Leu Glu Leu Leu Gly Lys Ser Thr Phe Val Leu Asp Asp Trp Lys
260 265 270

Arg Lys Tyr Ser Asn Glu Asp Thr Arg Pro Val Ala Leu Pro Trp Phe
275 280 285

Trp Glu His Tyr Asn Pro Glu Glu Tyr Ser Leu Trp Lys Val Thr Tyr
290 295 300

Lys Tyr Asn Asp Glu Leu Thr Leu Thr Phe Met Ser Asn Asn Leu Val
305 310 315 320

Gly Gly Phe Phe Asn Arg Leu Ser Ala Ser Thr Lys Tyr Met Phe Gly
325 330 335

Cys Leu Val Val Tyr Gly Glu Asn Asn Asn Asn Gly Ile Val Gly Ala
340 345 350

Val Met Val Arg Gly Gln Asp Tyr Val Pro Ala Phe Asp Val Ala Pro
355 360 365

Asp Trp Glu Ser Tyr Asp Tyr Ala Lys Leu Asp Pro Thr Asn Asp Asp
370 375 380

Asp Lys Glu Phe Ile Asn Asn Met Trp Ala Trp Asp Lys Pro Val Ser
385 390 395 400

Val Asn Gly Glu Pro Lys Glu Ile Val Asp Gly Lys Val Leu Lys
405 410 415

<210> SEQ ID NO 5

<211> LENGTH: 412

<212> TYPE: PRT

<213> ORGANISM: *Saccharomyces cerevisiae*

<400> SEQUENCE: 5

Met Ser Gln Gly Thr Leu Tyr Ile Asn Arg Ser Pro Arg Asn Tyr Ala
1 5 10 15

Ser Glu Ala Leu Ile Ser Tyr Phe Lys Leu Asp Val Lys Ile Val Asp
20 25 30

Leu Glu Gln Ser Ser Glu Phe Ala Ser Leu Phe Pro Leu Lys Gln Ala
35 40 45

Pro Ala Phe Leu Gly Pro Lys Gly Leu Lys Leu Thr Glu Ala Leu Ala
50 55 60

Ile Gln Phe Tyr Leu Ala Asn Gln Val Ala Asp Glu Lys Glu Arg Ala
65 70 75 80

Arg Leu Leu Gly Ser Asp Val Ile Glu Lys Ser Gln Ile Leu Arg Trp
85 90 95

Ala Ser Leu Ala Asn Ser Asp Val Met Ser Asn Ile Ala Arg Pro Phe
100 105 110

Leu Ser Phe Lys Gly Leu Ile Pro Tyr Asn Lys Lys Asp Val Asp Ala
115 120 125

Cys Phe Val Lys Ile Asp Asn Leu Ala Ala Val Phe Asp Ala Arg Leu
130 135 140

Arg Asp Tyr Thr Phe Val Ala Thr Glu Asn Ile Ser Leu Gly Asp Leu
145 150 155 160

His Ala Ala Gly Ser Trp Ala Phe Gly Leu Ala Thr Ile Leu Gly Pro
165 170 175

Glu Trp Arg Ala Lys His Pro His Leu Met Arg Trp Phe Asn Thr Val
180 185 190

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Ala Ala Ser Pro Ile Val Lys Thr Pro Phe Ala Glu Val Lys Leu Ala
 195 200 205

Glu Lys Ala Leu Thr Tyr Thr Pro Pro Lys Lys Gln Lys Ala Glu Lys
 210 215 220

Pro Lys Ala Glu Lys Ser Lys Ala Glu Lys Lys Lys Asp Glu Ala Lys
 225 230 235 240

Pro Ala Asp Asp Ala Ala Pro Ala Lys Lys Pro Lys His Pro Leu Glu
 245 250 255

Ala Leu Gly Lys Ser Thr Phe Val Leu Asp Asp Trp Lys Arg Lys Tyr
 260 265 270

Ser Asn Asp Asp Thr Arg Pro Val Ala Leu Pro Trp Phe Trp Glu His
 275 280 285

Tyr Asn Pro Glu Glu Tyr Ser Ile Trp Lys Val Gly Tyr Lys Tyr Asn
 290 295 300

Asp Glu Leu Thr Leu Thr Phe Met Ser Asn Asn Leu Val Gly Gly Phe
 305 310 315 320

Phe Asn Arg Leu Ser Ala Ser Thr Lys Tyr Met Phe Gly Cys Leu Val
 325 330 335

Val Tyr Gly Glu Asn Asn Asn Asn Gly Ile Val Gly Ala Val Met Val
 340 345 350

Arg Gly Gln Asp Phe Ala Pro Ala Phe Asp Val Ala Pro Asp Trp Glu
 355 360 365

Ser Tyr Glu Tyr Thr Lys Leu Asp Pro Thr Lys Glu Glu Asp Lys Glu
 370 375 380

Phe Val Asn Asn Met Trp Ala Trp Asp Lys Pro Val Val Val Asn Gly
 385 390 395 400

Glu Asp Lys Glu Ile Val Asp Gly Lys Val Leu Lys
 405 410

<210> SEQ ID NO 6
 <211> LENGTH: 206
 <212> TYPE: PRT
 <213> ORGANISM: *Saccharomyces cerevisiae*

<400> SEQUENCE: 6

Met Ala Ser Thr Asp Phe Ser Lys Ile Glu Thr Leu Lys Gln Leu Asn
 1 5 10 15

Ala Ser Leu Ala Asp Lys Ser Tyr Ile Glu Gly Thr Ala Val Ser Gln
 20 25 30

Ala Asp Val Thr Val Phe Lys Ala Phe Gln Ser Ala Tyr Pro Glu Phe
 35 40 45

Ser Arg Trp Phe Asn His Ile Ala Ser Lys Ala Asp Glu Phe Asp Ser
 50 55 60

Phe Pro Ala Ala Ser Ala Ala Ala Ala Glu Glu Glu Glu Asp Asp Asp
 65 70 75 80

Val Asp Leu Phe Gly Ser Asp Asp Glu Glu Ala Asp Ala Glu Ala Glu
 85 90 95

Lys Leu Lys Ala Glu Arg Ile Ala Ala Tyr Asn Ala Lys Lys Ala Ala
 100 105 110

Lys Pro Ala Lys Pro Ala Ala Lys Ser Ile Val Thr Leu Asp Val Lys
 115 120 125

Pro Trp Asp Asp Glu Thr Asn Leu Glu Glu Met Val Ala Asn Val Lys

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130				135				140							
Ala	Ile	Glu	Met	Glu	Gly	Leu	Thr	Trp	Gly	Ala	His	Gln	Phe	Ile	Pro
145				150					155						160
Ile	Gly	Phe	Gly	Ile	Lys	Lys	Leu	Gln	Ile	Asn	Cys	Val	Val	Glu	Asp
			165					170						175	
Asp	Lys	Val	Ser	Leu	Asp	Asp	Leu	Gln	Gln	Ser	Ile	Glu	Glu	Asp	Glu
			180					185						190	
Asp	His	Val	Gln	Ser	Thr	Asp	Ile	Ala	Ala	Met	Gln	Lys	Leu		
		195					200						205		

<210> SEQ ID NO 7

<211> LENGTH: 1044

<212> TYPE: PRT

<213> ORGANISM: *Saccharomyces cerevisiae*

<400> SEQUENCE: 7

Met	Ser	Asp	Ser	Gln	Gln	Ser	Ile	Lys	Val	Leu	Glu	Glu	Leu	Phe	Gln
1				5					10					15	
Lys	Leu	Ser	Val	Ala	Thr	Ala	Asp	Asn	Arg	His	Glu	Ile	Ala	Ser	Glu
			20					25					30		
Val	Ala	Ser	Phe	Leu	Asn	Gly	Asn	Ile	Ile	Glu	His	Asp	Val	Pro	Glu
			35				40					45			
His	Phe	Phe	Gly	Glu	Leu	Ala	Lys	Gly	Ile	Lys	Asp	Lys	Lys	Thr	Ala
	50						55				60				
Ala	Asn	Ala	Met	Gln	Ala	Val	Ala	His	Ile	Ala	Asn	Gln	Ser	Asn	Leu
65				70						75					80
Ser	Pro	Ser	Val	Glu	Pro	Tyr	Ile	Val	Gln	Leu	Val	Pro	Ala	Ile	Cys
				85					90					95	
Thr	Asn	Ala	Gly	Asn	Lys	Asp	Lys	Glu	Ile	Gln	Ser	Val	Ala	Ser	Glu
			100					105						110	
Thr	Leu	Ile	Ser	Ile	Val	Asn	Ala	Val	Asn	Pro	Val	Ala	Ile	Lys	Ala
			115				120					125			
Leu	Leu	Pro	His	Leu	Thr	Asn	Ala	Ile	Val	Glu	Thr	Asn	Lys	Trp	Gln
		130				135					140				
Glu	Lys	Ile	Ala	Ile	Leu	Ala	Ala	Ile	Ser	Ala	Met	Val	Asp	Ala	Ala
145					150					155					160
Lys	Asp	Gln	Val	Ala	Leu	Arg	Met	Pro	Glu	Leu	Ile	Pro	Val	Leu	Ser
				165					170					175	
Glu	Thr	Met	Trp	Asp	Thr	Lys	Lys	Glu	Val	Lys	Ala	Ala	Ala	Thr	Ala
			180					185						190	
Ala	Met	Thr	Lys	Ala	Thr	Glu	Thr	Val	Asp	Asn	Lys	Asp	Ile	Glu	Arg
		195					200					205			
Phe	Ile	Pro	Ser	Leu	Ile	Gln	Cys	Ile	Ala	Asp	Pro	Thr	Glu	Val	Pro
		210				215					220				
Glu	Thr	Val	His	Leu	Leu	Gly	Ala	Thr	Thr	Phe	Val	Ala	Glu	Val	Thr
225					230					235					240
Pro	Ala	Thr	Leu	Ser	Ile	Met	Val	Pro	Leu	Leu	Ser	Arg	Gly	Leu	Asn
				245					250					255	
Glu	Arg	Glu	Thr	Gly	Ile	Lys	Arg	Lys	Ser	Ala	Val	Ile	Ile	Asp	Asn
			260					265					270		
Met	Cys	Lys	Leu	Val	Glu	Asp	Pro	Gln	Val	Ile	Ala	Pro	Phe	Leu	Gly
			275				280						285		

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Lys Leu Leu Pro Gly Leu Lys Ser Asn Phe Ala Thr Ile Ala Asp Pro
 290 295 300

Glu Ala Arg Glu Val Thr Leu Arg Ala Leu Lys Thr Leu Arg Arg Val
 305 310 315 320

Gly Asn Val Gly Glu Asp Asp Ala Ile Pro Glu Val Ser His Ala Gly
 325 330 335

Asp Val Ser Thr Thr Leu Gln Val Val Asn Glu Leu Leu Lys Asp Glu
 340 345 350

Thr Val Ala Pro Arg Phe Lys Ile Val Val Glu Tyr Ile Ala Ala Ile
 355 360 365

Gly Ala Asp Leu Ile Asp Glu Arg Ile Ile Asp Gln Gln Ala Trp Phe
 370 375 380

Thr His Ile Thr Pro Tyr Met Thr Ile Phe Leu His Glu Lys Lys Ala
 385 390 395 400

Lys Asp Ile Leu Asp Glu Phe Arg Lys Arg Ala Val Asp Asn Ile Pro
 405 410 415

Val Gly Pro Asn Phe Asp Asp Glu Glu Asp Glu Gly Glu Asp Leu Cys
 420 425 430

Asn Cys Glu Phe Ser Leu Ala Tyr Gly Ala Lys Ile Leu Leu Asn Lys
 435 440 445

Thr Gln Leu Arg Leu Lys Arg Ala Arg Arg Tyr Gly Ile Cys Gly Pro
 450 455 460

Asn Gly Cys Gly Lys Ser Thr Leu Met Arg Ala Ile Ala Asn Gly Gln
 465 470 475 480

Val Asp Gly Phe Pro Thr Gln Glu Glu Cys Arg Thr Val Tyr Val Glu
 485 490 495

His Asp Ile Asp Gly Thr His Ser Asp Thr Ser Val Leu Asp Phe Val
 500 505 510

Phe Glu Ser Gly Val Gly Thr Lys Glu Ala Ile Lys Asp Lys Leu Ile
 515 520 525

Glu Phe Gly Phe Thr Asp Glu Met Ile Ala Met Pro Ile Ser Ala Leu
 530 535 540

Ser Gly Gly Trp Lys Met Lys Leu Ala Leu Ala Arg Ala Val Leu Arg
 545 550 555 560

Asn Ala Asp Ile Leu Leu Leu Asp Glu Pro Thr Asn His Leu Asp Thr
 565 570 575

Val Asn Val Ala Trp Leu Val Asn Tyr Leu Asn Thr Cys Gly Ile Thr
 580 585 590

Ser Ile Thr Ile Ser His Asp Ser Val Phe Leu Asp Asn Val Cys Glu
 595 600 605

Tyr Ile Ile Asn Tyr Glu Gly Leu Lys Leu Arg Lys Tyr Lys Gly Asn
 610 615 620

Phe Thr Glu Phe Val Lys Lys Cys Pro Ala Ala Lys Ala Tyr Glu Glu
 625 630 635 640

Leu Ser Asn Thr Asp Leu Glu Phe Lys Phe Pro Glu Pro Gly Tyr Leu
 645 650 655

Glu Gly Val Lys Thr Lys Gln Lys Ala Ile Val Lys Val Thr Asn Met
 660 665 670

Glu Phe Gln Tyr Pro Gly Thr Ser Lys Pro Gln Ile Thr Asp Ile Asn
 675 680 685

Phe Gln Cys Ser Leu Ser Ser Arg Ile Ala Val Ile Gly Pro Asn Gly

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690			695			700									
Ala	Gly	Lys	Ser	Thr	Leu	Ile	Asn	Val	Leu	Thr	Gly	Glu	Leu	Leu	Pro
705					710				715						720
Thr	Ser	Gly	Glu	Val	Tyr	Thr	His	Glu	Asn	Cys	Arg	Ile	Ala	Tyr	Ile
				725					730						735
Lys	Gln	His	Ala	Phe	Ala	His	Ile	Glu	Ser	His	Leu	Asp	Lys	Thr	Pro
			740						745						750
Ser	Glu	Tyr	Ile	Gln	Trp	Arg	Phe	Gln	Thr	Gly	Glu	Asp	Arg	Glu	Thr
			755						760						765
Met	Asp	Arg	Ala	Asn	Arg	Gln	Ile	Asn	Glu	Asn	Asp	Ala	Glu	Ala	Met
	770					775									780
Asn	Lys	Ile	Phe	Lys	Ile	Glu	Gly	Thr	Pro	Arg	Arg	Ile	Ala	Gly	Ile
					790										800
His	Ser	Arg	Arg	Lys	Phe	Lys	Asn	Thr	Tyr	Glu	Tyr	Glu	Cys	Ser	Phe
				805						810					815
Leu	Leu	Gly	Glu	Asn	Ile	Gly	Met	Lys	Ser	Glu	Arg	Trp	Val	Pro	Met
				820						825					830
Met	Ser	Val	Asp	Asn	Ala	Trp	Ile	Pro	Arg	Gly	Glu	Leu	Val	Glu	Ser
				835											845
His	Ser	Lys	Met	Val	Ala	Glu	Val	Asp	Met	Lys	Glu	Ala	Leu	Ala	Ser
				850											860
Gly	Gln	Phe	Arg	Pro	Leu	Thr	Arg	Lys	Glu	Ile	Glu	Glu	His	Cys	Ser
					870										880
Met	Leu	Gly	Leu	Asp	Pro	Glu	Ile	Val	Ser	His	Ser	Arg	Ile	Arg	Gly
				885											895
Leu	Ser	Gly	Gly	Gln	Lys	Val	Lys	Leu	Val	Leu	Ala	Ala	Gly	Thr	Trp
				900						905					910
Gln	Arg	Pro	His	Leu	Ile	Val	Leu	Asp	Glu	Pro	Thr	Asn	Tyr	Leu	Asp
				915						920					925
Arg	Asp	Ser	Leu	Gly	Ala	Leu	Ser	Lys	Ala	Leu	Lys	Glu	Phe	Glu	Gly
				930											940
Gly	Val	Ile	Ile	Ile	Thr	His	Ser	Ala	Glu	Phe	Thr	Lys	Asn	Leu	Thr
					945										950
Glu	Glu	Val	Trp	Ala	Val	Lys	Asp	Gly	Arg	Met	Thr	Pro	Ser	Gly	His
				965											975
Asn	Trp	Val	Ser	Gly	Gln	Gly	Ala	Gly	Pro	Arg	Ile	Glu	Lys	Lys	Glu
				980											990
Asp	Glu	Glu	Asp	Lys	Phe	Asp	Ala	Met	Gly	Asn	Lys	Ile	Ala	Gly	Gly
				995											1005
Lys	Lys	Lys	Lys	Lys	Leu	Ser	Ser	Ala	Glu	Leu	Arg	Lys	Lys	Lys	
					1010										1020
Lys	Glu	Arg	Met	Lys	Lys	Lys	Lys	Glu	Leu	Gly	Asp	Ala	Tyr	Val	
					1025										1035
Ser	Ser	Asp	Glu	Glu	Phe										
					1040										

<210> SEQ ID NO 8

<211> LENGTH: 579

<212> TYPE: PRT

<213> ORGANISM: *Saccharomyces cerevisiae*

<400> SEQUENCE: 8

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Met	Lys	Tyr	Val	Val	Val	Ser	Gly	Gly	Val	Ile	Ser	Gly	Ile	Gly	Lys	1	5	10	15
Gly	Val	Leu	Ala	Ser	Ser	Thr	Gly	Met	Leu	Met	Lys	Thr	Leu	Gly	Leu	20	25	30	
Lys	Val	Thr	Ser	Ile	Lys	Ile	Asp	Pro	Tyr	Met	Asn	Ile	Asp	Ala	Gly	35	40	45	
Thr	Met	Ser	Pro	Leu	Glu	His	Gly	Glu	Cys	Phe	Val	Leu	Asp	Asp	Gly	50	55	60	
Gly	Glu	Thr	Asp	Leu	Asp	Leu	Gly	Asn	Tyr	Glu	Arg	Tyr	Leu	Gly	Val	65	70	75	80
Thr	Leu	Thr	Lys	Asp	His	Asn	Ile	Thr	Thr	Gly	Lys	Ile	Tyr	Ser	His	85	90	95	
Val	Ile	Ala	Lys	Glu	Arg	Lys	Gly	Asp	Tyr	Leu	Gly	Lys	Thr	Val	Gln	100	105	110	
Ile	Val	Pro	His	Leu	Thr	Asn	Ala	Ile	Gln	Asp	Trp	Ile	Glu	Arg	Val	115	120	125	
Ala	Lys	Ile	Pro	Val	Asp	Asp	Thr	Gly	Met	Glu	Pro	Asp	Val	Cys	Ile	130	135	140	
Ile	Glu	Leu	Gly	Gly	Thr	Val	Gly	Asp	Ile	Glu	Ser	Ala	Pro	Phe	Val	145	150	155	160
Glu	Ala	Leu	Arg	Gln	Phe	Gln	Phe	Lys	Val	Gly	Lys	Glu	Asn	Phe	Ala	165	170	175	
Leu	Ile	His	Val	Ser	Leu	Val	Pro	Val	Ile	His	Gly	Glu	Gln	Lys	Thr	180	185	190	
Lys	Pro	Thr	Gln	Ala	Ala	Ile	Lys	Gly	Leu	Arg	Ser	Leu	Gly	Leu	Val	195	200	205	
Pro	Asp	Met	Ile	Ala	Cys	Arg	Cys	Ser	Glu	Thr	Leu	Asp	Lys	Pro	Thr	210	215	220	
Ile	Asp	Lys	Ile	Ala	Met	Phe	Cys	His	Val	Gly	Pro	Glu	Gln	Val	Val	225	230	235	240
Asn	Val	His	Asp	Val	Asn	Ser	Thr	Tyr	His	Val	Pro	Leu	Leu	Leu	Leu	245	250	255	
Glu	Gln	Lys	Met	Ile	Asp	Tyr	Leu	His	Ala	Arg	Leu	Lys	Leu	Asp	Glu	260	265	270	
Ile	Ser	Leu	Thr	Glu	Glu	Glu	Lys	Gln	Arg	Gly	Leu	Glu	Leu	Leu	Ser	275	280	285	
Lys	Trp	Lys	Ala	Thr	Thr	Gly	Asn	Phe	Asp	Glu	Ser	Met	Glu	Thr	Val	290	295	300	
Lys	Ile	Ala	Leu	Val	Gly	Lys	Tyr	Thr	Asn	Leu	Lys	Asp	Ser	Tyr	Leu	305	310	315	320
Ser	Val	Ile	Lys	Ala	Leu	Glu	His	Ser	Ser	Met	Lys	Cys	Arg	Arg	Lys	325	330	335	
Leu	Asp	Ile	Lys	Trp	Val	Glu	Ala	Thr	Asp	Leu	Glu	Pro	Glu	Ala	Gln	340	345	350	
Glu	Ser	Asn	Lys	Thr	Lys	Phe	His	Glu	Ala	Trp	Asn	Met	Val	Ser	Thr	355	360	365	
Ala	Asp	Gly	Ile	Leu	Ile	Pro	Gly	Gly	Phe	Gly	Val	Arg	Gly	Thr	Glu	370	375	380	
Gly	Met	Val	Leu	Ala	Ala	Arg	Trp	Ala	Arg	Glu	Asn	His	Ile	Pro	Phe	385	390	395	400
Leu	Gly	Val	Cys	Leu	Gly	Leu	Gln	Ile	Ala	Thr	Ile	Glu	Phe	Thr	Arg				

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100	105	110
Leu Ser Phe Lys Gly Leu Ile Pro Tyr Asn Lys Lys Asp Val Asp Ala 115 120 125		
Cys Phe Val Lys Ile Asp Asn Leu Ala Ala Val Phe Asp Ala Arg Leu 130 135 140		
Arg Asp Tyr Thr Phe Val Ala Thr Glu Asn Ile Ser 145 150 155		

<210> SEQ ID NO 12
 <211> LENGTH: 159
 <212> TYPE: PRT
 <213> ORGANISM: *Saccharomyces cerevisiae*

<400> SEQUENCE: 12

Met Ser Gln Gly Thr Leu Tyr Ala Asn Phe Arg Ile Arg Thr Trp Val 1 5 10 15
Pro Arg Gly Leu Val Lys Ala Leu Lys Leu Asp Val Lys Val Val Thr 20 25 30
Pro Asp Ala Ala Ala Glu Gln Phe Ala Arg Asp Phe Pro Leu Lys Lys 35 40 45
Val Pro Ala Phe Val Gly Pro Lys Gly Tyr Lys Leu Thr Glu Ala Met 50 55 60
Ala Ile Asn Tyr Tyr Leu Val Lys Leu Ser Gln Asp Asp Lys Met Lys 65 70 75 80
Thr Gln Leu Leu Gly Ala Asp Asp Asp Leu Asn Ala Gln Ala Gln Ile 85 90 95
Ile Arg Trp Gln Ser Leu Ala Asn Ser Asp Leu Cys Ile Gln Ile Ala 100 105 110
Asn Thr Ile Val Pro Leu Lys Gly Gly Ala Pro Tyr Asn Lys Lys Ser 115 120 125
Val Asp Ser Ala Met Asp Ala Val Asp Lys Ile Val Asp Ile Phe Glu 130 135 140
Asn Arg Leu Lys Asn Tyr Thr Tyr Leu Ala Thr Glu Asn Ile Ser 145 150 155

<210> SEQ ID NO 13
 <211> LENGTH: 8
 <212> TYPE: PRT
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: Synthetic Peptide

<400> SEQUENCE: 13

Arg Gly Ser Ile Asp Thr Trp Val 1 5
--

<210> SEQ ID NO 14
 <211> LENGTH: 7
 <212> TYPE: PRT
 <213> ORGANISM: Artificial Sequence
 <220> FEATURE:
 <223> OTHER INFORMATION: Synthetic Peptide

<400> SEQUENCE: 14

Glu Glu Trp Glu Thr Trp Val 1 5

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<210> SEQ ID NO 15
<211> LENGTH: 9
<212> TYPE: PRT
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic Peptide

<400> SEQUENCE: 15

Tyr Pro Tyr Asp Val Pro Asp Tyr Ala
1 5

<210> SEQ ID NO 16
<211> LENGTH: 10
<212> TYPE: PRT
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic Peptide

<400> SEQUENCE: 16

Glu Gln Lys Leu Ile Ser Glu Glu Asp Leu
1 5 10

<210> SEQ ID NO 17
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1. A self-assembling fusion protein comprising:
 - (a) a heat-inducible self-assembly domain; and
 - (b) a target protein; wherein the self-assembly domain remains folded during assembly.
2. (canceled)
3. The fusion protein of claim 1, wherein the self-assembly domain is a GST-like domain or a polypeptide with at least 90% identity to a GST-like domain.
4. The fusion protein of claim 1, wherein the self-assembly domain comprises a polypeptide from Arc1, Mes1, Gus1, or a polypeptide with at least 90% identity to Arc1, Mes1, or Gus1.
5. The fusion protein of claim 4, wherein the self-assembly domain comprises a polypeptide that is at least 20 amino acids in length and has at least 90% identity to 20 contiguous amino acids of the first 250 amino acids from Arc1, Mes1, or Gus1.
6. The fusion protein of claim 4, wherein the self-assembly domain comprises a polypeptide from Gus1 or a polypeptide with at least 90% identity to Gus1 or a fragment thereof.
7. The fusion protein of claim 1, wherein the self-assembly domain comprises a polypeptide from Tef3, Tef4, Efb1, or a polypeptide with at least 90% identity to Tef3, Tef4, Efb1, or fragments thereof.
8. (canceled)
9. The fusion protein of claim 1, wherein the self-assembly domain comprises a polypeptide from Yef3, Ura7, or a polypeptide with at least 90% identity to Yef3 Ura7, or fragments thereof.
10. (canceled)
11. The fusion protein of claim 1, wherein the self-assembly domain is a polypeptide from a *Saccharomyces cerevisiae* protein or a polypeptide with at least 90% identity to a *Saccharomyces cerevisiae* protein.
12. The fusion protein of claim 1, wherein the self-assembly domain comprises a polypeptide of SEQ ID NO:9, 10, 11, or 12, a fragment thereof, or a polypeptide with at least 90% identity to SEQ ID NO:9, 10, 11, or 12, or a fragment thereof.
- 13-14. (canceled)
15. The fusion protein of claim 1, wherein the self-assembly domain is between 20 and 250 amino acids in length.
16. The fusion protein of claim 1, wherein the target protein is ferritin, a fluorescent protein, an antibody, an antibody fragment, protein A, streptavidin, protein G, protein A/G, protein L, a protease, PDZ domain or StrepTactin.
17. The fusion protein of claim 1, wherein the fusion protein further comprises a protease cleavage site between the target protein and the self-assembly domain.
- 18-20. (canceled)
21. A protein aggregate comprising the fusion protein of claim 1.
- 22-23. (canceled)
24. A polynucleotide encoding for the fusion protein of claim 1.
25. A host cell comprising the fusion protein of claim 1.
26. Cell lysate comprising the fusion protein of claim 1.
27. A method for aggregating a target protein comprising: formulating an aqueous composition comprising the fusion protein of claim 1; and heating the composition to a temperature between about 35 and 50° C.
- 28-30. (canceled)
31. A method of immunoprecipitating or purifying a molecule comprising the steps of:
 - formulating a first composition comprising the fusion protein of claim 1; wherein the target protein is a first target protein that specifically binds to the molecule;
 - heating the first composition to a temperature between about 35 and 50° C. to form protein aggregates comprising the fusion protein; and contacting the first composition with a second composition comprising the molecule.
- 32-45. (canceled)
46. A method for purifying a protein comprising:
 - formulating an aqueous composition comprising a fusion protein of claim 1;
 - heating the aqueous composition to a temperature between about 35 and 50° C. to form protein aggregates comprising the fusion protein;
 - separating the protein aggregates from the aqueous composition.
- 47-49. (canceled)
50. The fusion protein of claim 1, wherein the target protein is a restriction enzyme, DNA polymerase, protease, ligase, RNA polymerase, methylase, polyadenylate polymerase, topoisomerase, guanylyl transferase, ribonuclease, deoxyribonuclease, alkaline phosphatase, polynucleotide kinase or reverse transcriptase.

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