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(54) Title: ORTHOGONAL PROTEIN HETERODIMERS

(57) Abstract: Disclosed herein are designed heterodimer proteins, monomeric polypeptides capable of forming heterodimer proteins, protein scaffolds including such polypeptides, and methods for using the heterodimer proteins and subunit polypeptides for designing logic gates.

WO 2020/093043 A1

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ORTHOGONAL PROTEIN HETERODIMERS**Cross Reference**

This application claims priority to U.S. Provisional Patent Application Serial Nos. 62/755,264 filed November 2, 2018 and 62/904,800 filed September 24, 2019, each
10 incorporated by reference herein in their entirety.

Federal Funding Statement

This invention was made with government support under Grant No. GM103533 awarded by the National Institutes of Health. The government has certain rights in the
15 invention

Background

Heterodimeric interaction specificity between two DNA strands, and between protein and DNA, is often achieved by varying side chains or bases coming off the protein or DNA
20 backbone—for example, the bases participating in Watson-Crick base pairing in the double helix, or the side chains of protein contacting DNA in TALEN-DNA complexes. This modularity enables the generation of an essentially unlimited number of orthogonal DNA-DNA and protein-DNA heterodimers. In contrast, protein-protein interaction specificity is often achieved through backbone shape complementarity, which is less modular and hence
25 harder to generalize.

Summary

In one aspect, the disclosure provides designed heterodimer proteins, comprising:

30 (a) a monomer A polypeptide, wherein the monomer A polypeptide is a non-naturally occurring polypeptide comprising 1-5 alpha helices connected by amino acid linkers; and

(b) a monomer B polypeptide, wherein the monomer B polypeptide is a non-naturally occurring polypeptide comprising 1-5 alpha helices connected by amino acid
35 linkers,

wherein monomer A and monomer B non-covalently interact to form the designed heterodimer protein. In one embodiment,

(i) monomer A comprises a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an odd-numbered SEQ ID NO selected from the group consisting of selected from the group SEQ ID NOS: 1-290; and

(ii) monomer B comprises a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an even-numbered SEQ ID NO selected from the group consisting of selected from the group SEQ ID NOS: 1-290, wherein the even-numbered SEQ ID NO is the binding partner of the odd-numbered SEQ ID NO. in step (i).

In another aspect, the disclosure provides non-naturally occurring polypeptide comprising a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS: 1-290.

In another aspect, the disclosure provides non-naturally occurring polypeptide comprising a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS: 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494.

In another aspect, the disclosure provides proteins comprising 2, 3, 4, or more non-naturally occurring polypeptides having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS: 1-290, wherein the 2, 3, 4, or more naturally occurring polypeptides are covalently linked. In one embodiment, each of the 2, 3, 4, or more non-naturally occurring polypeptides are different. In another embodiment, each of the 2, 3, 4, or more non-naturally occurring polypeptides are present in a fusion protein.

In another aspect, the disclosure provides proteins comprising 2, 3, 4, or more non-naturally occurring polypeptides having at least 70%, 75%, 80%, 85%, 90%, 91%,

92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS: 1-290, 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494 wherein the 2, 3, 4, or more naturally occurring polypeptides are covalently linked. In one embodiment, each of the 2, 3, 4, or more non-naturally occurring polypeptides are different. In another embodiment, each of the 2, 3, 4, or more non-naturally occurring polypeptides are present in a fusion protein. In each of these aspects, amino acid changes from the reference amino acid sequence may be conservative amino acid substitutions. In another embodiment, at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions are invariant compared to the reference amino acid sequence.

In another aspect, the disclosure provides protein scaffolds, comprising a) a first designed component comprised of any number of monomer A polypeptides and/or monomer B polypeptides, each from different heterodimers, connected into a single component by amino acid linkers.

b) a second designed component, comprising corresponding monomers for each monomer A and/or monomer B in the first designed component one;

wherein the first and second designed components interact to form the protein scaffold, and wherein each monomer A only interacts in the scaffold with its monomer B mate.

In another aspect, the disclosure provides methods of forming the designed heterodimer protein of any embodiment of the disclosure, comprising:

a) providing two of the monomers as unlinked monomers;

b) providing the other two monomers as linked monomers;

whereby the unlinked monomers associate with their respective monomer of the same heterodimer, and not with any of the other monomers.

In another aspect, the disclosure provides designed heterodimer proteins, comprising:

a) asymmetric buried hydrogen bond networks incorporated into regularly repeating backbone structures; and

b) helix hairpin helix monomers wherein the supercoil phases of the helices are fixed at 0, 90, 180, or 270 degrees and the supercoil twist (ω_0) and helical twist (ω_1) are held constant for either a two layer left handed super coil ($\omega_0=-2.85$ and $\omega_1=102.85$), or a 5 layer untwisted bundle ($\omega_0=0$ and $\omega_1=100$)

5 In another aspect, the disclosure provides fusion proteins comprising a polypeptide of the formula X-B-Z, wherein:

(a) the X domain is a non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein the X domain is capable of non-covalently binding to a first target;

10 (b) the Z domain is a non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein the Z domain is capable of non-covalently binding to either (i) a second target that differs from the first target, or (ii) a different non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices; and

(c) the B domain is an amino acid linker;

15 wherein a combined number of alpha helices from the X domain and the Z domain is 4, 5, or 6; and

wherein the X domain and the Z domain interact at a binding interface, wherein the binding interface comprises a hydrogen bond network in which at least one side chain in each alpha helix hydrogen of the X domain bonds with a side chain in an alpha helix in the Z domain, and wherein the binding interface comprises a plurality of hydrophobic residues.

20 In another aspect, the disclosure provides kits or compositions, comprising at least two fusion proteins comprising the formula X-B-Z, wherein

the B domain in each fusion protein is independently a polypeptide linker;

the X domain in each fusion protein comprises a first non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices;

25 the Z domain in each fusion protein comprises a second non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein a combined number of alpha helices from the X domain and the Z domain in each individual fusion protein is 4, 5, or 6; wherein the X domain and the Z domain interact at a binding interface, wherein the binding interface comprises a hydrogen bond network in which at least one side chain in each X domain alpha
30 helix bonds with a side chain in an alpha helix in the Z domain; wherein

the X domain in a first fusion protein is capable of non-covalently binding to a first target;

the Z domain in a second fusion protein is capable of non-covalently binding to a second target; and

the X domains and Z domains in each individual fusion protein that are not capable of non-covalently binding to the first target or the second target are capable of non-covalently binding to an X or a Z domain of a different fusion protein in the plurality of fusion proteins.

In one embodiment of the fusion proteins, kits, or compositions, each X domain and each Z domain comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide comprising the amino acid sequence selected from SEQ ID NO:1-290, with the proviso that the X domain and the Z domain do not do not form a heterodimer (a-b) pair. In another embodiment at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of each X domain and each Z domain are invariant compared to the reference amino acid sequence.

In one aspect, the disclosure provides methods, comprising:

- (i) contacting a fusion protein according to any embodiment of the disclosure with a biological sample under conditions to promote non-covalent binding of the fusion protein with first target and second target present in the sample, and
- (ii) detecting non-covalent binding of the one or more fusion proteins to the first target and/or the second target in the biological sample.

In one embodiment, the method comprises detecting cooperative non-covalently binding of the one or more fusion proteins to the first target and the second target in the biological sample. In another embodiment, the method comprises detecting non-covalent binding of the one or more fusion proteins to the first target or the second target in the biological sample.

In another aspect, the disclosure provides methods for target detection, comprising

- (a) contacting a biological sample with at least two fusion proteins, wherein each of the at least two fusion proteins comprises the formula X-B-Z, wherein
 - each B is independently a polypeptide linker;
 - each X domain comprises a first non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices;
 - each Z domain comprises a second non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein a combined number of alpha helices from the X domain and the Z domain in each individual fusion protein is 4, 5, or 6; wherein the X domain and the Z domain interact at a binding interface, wherein the binding interface comprises a hydrogen

bond network in which at least one side chain in each X domain alpha helix bonds with a side chain in an alpha helix in the Z domain; wherein

the X domain in a first fusion protein is capable of non-covalently binding to a first target;

5 the Z domain in a second fusion protein is capable of non-covalently binding to a second target; and

the X domains and Z domains in each individual fusion protein that are not capable of non-covalently binding to the first target or the second target are capable of non-covalently binding to an X or a Z domain of a different fusion protein in the plurality of fusion proteins;

10 (b) detecting non-covalent binding of the two or more fusion proteins to the first target and/or the second target in the biological sample.

In one aspect, the disclosure provides compositions comprising

(a) a first polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding a first target; and

15 (b) a second polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding to the second polypeptide, and wherein the second polypeptide is capable of non-covalently binding a second target that differs from the first target; wherein:

(i) a binding affinity of the first polypeptide for the first target is
20 approximately equal to a binding affinity of the second polypeptide for the second target; and

(ii) the binding affinity of the first polypeptide for the first target and the binding affinity of the second polypeptide for the second target are greater than the binding affinity of the first target and the second target for each other.

In one aspect, the disclosure provides compositions comprising

25 (a) a first polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding a first target; and

(b) a second polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding to the second polypeptide, and wherein the second polypeptide is capable of non-covalently binding a second target that differs from the first
30 target; wherein:

(i) a binding affinity of the first polypeptide for the second polypeptide is greater than a binding affinity of the second polypeptide for the second target;

(ii) a binding affinity of the first polypeptide for the first target is approximately equal to a binding affinity of the second polypeptide for the second target; and

(iii) the binding affinity of the first polypeptide for the first target and the binding affinity of the second polypeptide for the second target are greater than the binding affinity of the first target and the second target for each other.

In another aspect, the disclosure provides compositions comprising

5 (a) a first polypeptide comprising 4 alpha helices, wherein the first polypeptide is capable of non-covalently binding a first target; and

(b) a second polypeptide comprising 4 alpha helices, wherein the second polypeptide is capable of non-covalently binding a second target that differs from the first target; wherein:

10 (i) a binding affinity of the first target for the second target is greater than a binding affinity of the first polypeptide for the first target;

(ii) a binding affinity of the first polypeptide for the first target is approximately equal to a binding affinity of the second polypeptide for the second target; and

(iii) the sum of the binding affinity of (A) the first polypeptide for the first target and (B) the binding affinity of the second polypeptide for the second target, is greater than the binding affinity of the first target and the second target.

In various embodiments for each composition of the disclosure, the composition may further comprise the first target and the second target, and the first target and/or the second target further may comprise one or more effector polypeptide domains. In one embodiment, 20 the first polypeptide and/or the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide comprising the amino acid sequence selected from the group consisting of SEQ ID NO:1-290, or the group consisting of SEQ ID NOS:1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494. In another embodiment, the first target and/or the second target each comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide 30 selected from the group including, but not limited to a polypeptide comprising the amino acid sequence selected from the group consisting of SEQ ID NO:1-290, or the group consisting of SEQ ID NOS:1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, with the proviso that the first target forms a

heterodimer (a-b) pair with the first polypeptide, and the second target forms a heterodimer (a-b) pair with the second polypeptide. In another embodiment, the compositions are contacted with a biological sample and binding is detected, such as detecting an output signal caused by actions of effector polypeptides upon binding.

5 The disclosure also provides nucleic acids encoding the polypeptides, proteins, and fusion proteins of the disclosure; expression vectors comprising the nucleic acids operatively linked to a promoter; and host cells comprising the nucleic acids, expression vectors, and/or polypeptides, proteins, fusion proteins, scaffolds, and designed heterodimer pairs of the disclosure.

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Description of the Figures

FIGs. 1A to 1H shows modular heterodimer design. FIG. 1A shows individual helix generation: the helical phase ($\Delta\phi_1$), supercoil radius (R) and offset along the Z-axis (Z offset) were exhaustively sampled; a total of 11 free parameters since there is no z offset for the first helix. FIG. 1B shows top-down view of the parallel twisted backbone. FIG. 1C shows representative hydrogen bond networks identified using HBNETTM. FIG. 1D shows matches of multiple HBNETTM containing heptads to a single full length backbone. FIG. 1E shows addition of loops to connect the 4 helices into two helix hairpins. FIGs. 1F, 1G, and 1H show SEC trace, CD spectra and (inset) temperature melt, and SAXS (black, experimental SAXS data; red, spectra computed from the designed backbones) profile of the design DHD37_ABXB. Experiments were performed once.

FIGs. 2A to 2F show structural characterization of designed heterodimers. FIGs. 2A-2D show crystal structures superimposed on design models with monomers; cross-sections on backbones (left) indicate locations of designed hydrogen-bond networks (middle panels). Solid and dashed boxes compare networks in design model and crystal structure. Black boxes compare overall hydrophobic packing. FIG. 2A shows DHD_131, 2.4 Å resolution with 1.0 Å C α RMSD. FIG. 2B shows DHD37_1:234, 3.3 Å resolution with 1.4 Å RMSD. FIG. 2C shows DHD_127, 1.8 Å resolution with 1.7 Å RMSD. FIG. 2D shows DHD_15, 3.4 Å resolution with 0.9 Å RMSD; hydrogen bond networks were not well resolved. FIGs. 2E-2F show DHD_39 and DHD_120 backbones and designed hydrogen bond networks. Experimental SAXS data (black) are similar to spectra computed from the designed backbones.

FIGs. 3A to 3C shows new functionality from DHD combinations. FIG. 3A shows induced dimerizer formed from “b” component of DHD13_XAAA fused to “b” component

of DHD37_ABXB with an intervening flexible linker. The “a” components of the two heterodimers are brought into close proximity by the heterodimerizer. FIG. 3B shows Y2H data on 4 induced dimerization systems. For each pair of bars: left, without heterodimerizer fusion; right, with heterodimerizer fusion. Dashed line indicates background growth with unfused AD and DBD. Data are mean \pm s.d. from 3 biological repeats. FIG. 3C shows 9_a, 13_XAAA_a and 37_ABXB_a were covalently linked to form a scaffold, recruiting 9_b (hexahistidine tagged), 13_XAAA_b and 37_ABXB_b.

FIGs. 4A to 4C show all-against-all orthogonality assessment. FIG. 4A shows Y2H for 21 heterodimers show heterodimer formation with little homodimer formation. First letter at bottom indicates monomer fused to AD, second letter, to DBD. FIG. 4B shows Y2H all by all testing of 9 pairs of heterodimers, colors indicate growth. Boxes indicate designed cognate heterodimer pairs, dashed black box indicates a set of 6 orthogonal heterodimers. FIG. 4C shows Off-target binding of DHD15_a and DHD13_XAAA_b, in the absence (left) or presence (right) of DHD15_b and DHD13_XAAA_a. Data are mean \pm s.d.. Red dashed line indicates background growth with unfused AD and DBD.

FIGs. 5A to 5B show example HBNets resulting from the systematic search. FIG. 5A shows overlay of 50 backbones with different Crick parameters for each helix. FIG. 5B shows example hydrogen bond networks from the systematic search, each involving at least 4 residues and contacting all 4 helices.

FIGs. 6A to 6C show thermal and chemical denaturation of DHDs. FIGs. 6A and 6B show CD spectra for thermal denaturation of DHD_15 and DHD_20, respectively. Top, wavelength scan at 25°C, 75°C, 95°C, and final 25°C. Designs were alpha helical and stable up to 95°C. Bottom, CD temperature melts, monitoring absorption at 222 nm as temperature was increased from 25°C to 95°C. FIG. 6C shows GdnHCl denaturation of DHD_127 by CD monitoring absorption at 222 nm. All CD experiments were performed once.

FIG. 7A to 7B show backbone and hydrogen bond network permutations. FIG. 7A shows on a 2+2 backbone (left), two loops were designed to connect the 4 helices into a single monomer in 2 different ways (middle), after which 4 different cut points were introduced to generate 4 possible backbone permuted heterodimers of a single helix and a three helix bundle (3+1 heterodimers, right). For example, 2:134 refers to a heterodimer where the original helix 2 is a single helix, and helices 1, 3, and 4 were connected into a 3 helix bundle. FIG. 7B shows hydrogen bond network permutation. Each unique network was assigned a letter (Networks “A” and “B” in this case), and with the hydrophobic packing assigned X. The backbone on the left reads “ABXB”, with its first heptad accommodates

network “A”, its second and fourth heptad accommodate network “B”, and its third heptad accommodates hydrophobic packing only.

FIGs. 8A to 8H show crystal structure of the domain swapped DHD_15 and biophysical characterization of higher order oligomers. FIG. 8A shows crystal structure of DHD_15 at pH 6.5, with 2.25 Å resolution. FIG. 8B shows superposition of design models in color onto both halves of the crystal structure in white, with backbone RMSD of 1.83 Å. FIGs. 8C to 8F show SEC traces of the induced dimerization DHD_9-13 fusion, DHD_15-37 fusion, DHD_13-37 fusion, and the scaffolding complex in FIG. 3C (the peak at around 15 mL corresponds to the fully assembled complex, followed by a peak representing excess of individual components). FIG. 8G shows CD thermal melt curves for the scaffolding complex in FIG. 3C. Wavelength scan was performed at 25°C, 75°C, 95°C, and final 25°C. Design was alpha helical and stable up to 95°C. FIG. 8H, CD chemical denaturation profile of the scaffolding complex in FIG. 3C. 2 (FIG. 8C to 8F) or 1 (FIG. 8G to 8H) biologically independent repeats were performed.

FIGs. 9A to 9G show Y2H all-against-all assay of 16 DHDs. FIG. 9A shows Y2H assay with cell growth on agar plates containing 100 mM 3-AT, lacking tryptophan, leucine and histidine. Plates were imaged at Day 5. White, no growth on agar plates; grey, weak growth forming non-circular colonies; black, strong growth. FIG. 9B shows Y2H result by growing yeast culture in liquid media containing 100 mM 3-AT, lacking tryptophan, leucine and histidine. OD 600 values were measured at Day 2 to evaluate cell growth. FIG. 9C shows an additional set of DHDs tested by Y2H showing improved orthogonality. FIG. 9D shows distribution of OD 600 values for non-cognate interactions in FIG. 9B, the majority of cells grew to OD 600 values less than 0.4, indicating weak interactions for non-cognate binding. FIG. 9E shows more buried bulky polar residues strongly correlates with design success. f, Successful designs tend to have bigger polar interface surface area. FIG. 9G shows designs with better hydrophobic packing (as reported by the ROSETTA™ filter value Average Degree on Ile, Leu and Val residues) tend to have a higher chance of being constitutive heterodimers. Two (FIGs. 9A to 9C) independent experiments were performed.

FIG. 10 shows hydrogen bond network sequence motifs of the set of 6 orthogonal pairs in Y2H experiments. Letters patches mark the location of hydrogen bond network forming residues on the backbones, and indicate residue identities.

FIGs. 11A to 11H show cooperativity of CIPHR logic gates. FIG. 11A shows backbone structure of *A:A'* heterodimer building block, with hydrogen bond network detail in inset. Bottom right, condensed representation used throughout figures. FIG. 11B shows

thermodynamic cycle describing the induced dimerization system. FIG. 11C shows simulation of the induced dimerization system under thermodynamic equilibrium. A and B' monomers were held constant while titrating in various initial amounts of the A'-B dimerizer proteins. If binding is not cooperative (small c), the final amount of trimeric complexes decreases when the dimerizer protein is in excess. FIG. 11D shows equilibrium denaturation experiments monitored by CD for designs with 6- and 12- amino acid (AA) linkers. Circles represent experimental data, and lines are fits to the 3-state unimolecular unfolding model. Design models are shown on the side. FIG. 11E shows experimental SAXS profile of $I'-2'$ with a 6-residue linker (in black), fitted to the calculated profile of $I:I'$ heterodimer. FIG. 11F shows an induced dimerization system using a 6-residue linker. FIG. 11G shows a two-input AND gate schematic. FIG. 11H shows a three-input AND gate.

FIGs. 12A to 12G show CIPHR two input logic gates. FIG. 12A shows CIPHR gates are built from DHDs (top) with monomers or covalently connected monomers as inputs (left); some gates utilize only the designed cognate interactions (left side of middle panel), while others take advantage of observed inter and intramolecular binding affinity hierarchies (right side of middle panel). FIGs. 12B and 12C show two-input AND (12B) and OR (12C) CIPHR logic gates based on orthogonal DHD interactions. FIGs. 12D to 12G show NOT (12D), NOR (12E), XNOR (12F), and NAND (12G) CIPHR logic gates made from multispecific and competitive protein binding. For each gate, black dots represent individual Y2H growth measurement corrected over background growth, with their average values shown in bars. * indicates no yeast growth over background. 0s and 1s in the middle and right blocks represent different input states and expected outputs, respectively. \

FIGs. 13A to 13E show three-input CIPHR logic gates. FIG. 13A shows schematic of a three-input AND gate. FIG. 13B shows schematic of a three-input OR gate. FIG. 13C shows Y2H results confirmed activation of the 3-input OR gate with either of the inputs. FIG. 13D shows schematic of a DNF gate. FIG. 13 E shows Y2H results confirmed proper activation of the gate. For each gate, black dots represent individual measurements corrected over background growth, with their average values shown in bars.

FIG. 14A shows molecular implementation of the cooperative induced dimerization system, binding only occurs when all three components are present. FIG. 14B shows size exclusion chromatography profiles of $I'-2'$ variants with 0, 2, 6, 12, and 24 amino acids in the flexible linker connecting I' and $2'$.

FIGs. 15A and 15B show binding affinity gradient from individual Y2H experiments. FIG. 15A shows the 8:8' heterodimer binds more tightly than the homodimers of its

monomers. FIG. 15B shows binding affinity gradient among the monomers of 1:1', 9:9', and 10:10' pairs.

FIG. 16 shows exemplary heterodimer proteins comprising combinations of monomer A and monomer B.

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Detailed Description

All references cited are herein incorporated by reference in their entirety. Within this application, unless otherwise stated, the techniques utilized may be found in any of several well-known references such as: *Molecular Cloning: A Laboratory Manual* (Sambrook, et al., 10 1989, Cold Spring Harbor Laboratory Press), *Gene Expression Technology* (Methods in Enzymology, Vol. 185, edited by D. Goeddel, 1991. Academic Press, San Diego, CA), "Guide to Protein Purification" in *Methods in Enzymology* (M.P. Deutscher, ed., (1990) Academic Press, Inc.); *PCR Protocols: A Guide to Methods and Applications* (Innis, et al. 1990. Academic Press, San Diego, CA), *Culture of Animal Cells: A Manual of Basic* 15 *Technique, 2nd Ed.* (R.I. Freshney. 1987. Liss, Inc. New York, NY), *Gene Transfer and Expression Protocols*, pp. 109-128, ed. E.J. Murray, The Humana Press Inc., Clifton, N.J.), and the Ambion 1998 Catalog (Ambion, Austin, TX).

As used herein, the singular forms "a", "an" and "the" include plural referents unless the context clearly dictates otherwise.

20 As used herein, the amino acid residues are abbreviated as follows: alanine (Ala; A), asparagine (Asn; N), aspartic acid (Asp; D), arginine (Arg; R), cysteine (Cys; C), glutamic acid (Glu; E), glutamine (Gln; Q), glycine (Gly; G), histidine (His; H), isoleucine (Ile; I), leucine (Leu; L), lysine (Lys; K), methionine (Met; M), phenylalanine (Phe; F), proline (Pro; P), serine (Ser; S), threonine (Thr; T), tryptophan (Trp; W), tyrosine (Tyr; Y), and 25 valine (Val; V).

All embodiments of any aspect of the disclosure can be used in combination, unless the context clearly dictates otherwise.

Unless the context clearly requires otherwise, throughout the description and the claims, the words 'comprise', 'comprising', and the like are to be construed in an inclusive 30 sense as opposed to an exclusive or exhaustive sense; that is to say, in the sense of "including, but not limited to". Words using the singular or plural number also include the plural and singular number, respectively. Additionally, the words "herein," "above," and "below" and words of similar import, when used in this application, shall refer to this application as a whole and not to any particular portions of the application.

It is understood that wherever aspects are described herein with the language "comprising," otherwise analogous aspects described in terms of "consisting of" and/or "consisting essentially of" are also provided.

The term "interface residue" or "interface position", as used herein, means amino acid
5 residues or positions that are interacting between at least two monomers in heterodimer,
heterotrimer, heterotetramer, etc. The interaction comprises a hydrogen bond network in
which at least a hydrogen from an alpha helix in the first monomer binds to a side chain in an
alpha helix in the second monomer. In some aspects, the interaction comprises at least one
10 hydrogen bond, at least two hydrogen bonds, at least three hydrogen bonds, at least four
hydrogen bonds, at least five hydrogen bonds, at least six hydrogen bonds, at least seven
hydrogen bonds, at least eight hydrogen bonds, at least nine hydrogen bonds, and at least ten
hydrogen bonds. In some aspects, the interface residue comprises hydrophobic residues.

The description of embodiments of the disclosure is not intended to be exhaustive or
to limit the disclosure to the precise form disclosed. While the specific embodiments of, and
15 examples for, the disclosure are described herein for illustrative purposes, various equivalent
modifications are possible within the scope of the disclosure, as those skilled in the relevant
art will recognize.

In a first aspect the disclosure provide designed heterodimer proteins, comprising:

(a) a monomer A polypeptide, wherein the monomer A polypeptide is a non-
20 naturally occurring polypeptide comprising 1-5 alpha helices connected by amino acid
linkers; and

(b) a monomer B polypeptide, wherein the monomer B polypeptide is a non-
naturally occurring polypeptide comprising 1-5 alpha helices connected by amino acid
linkers,

25 wherein monomer A and monomer B non-covalently interact to form the
designed heterodimer protein.

The disclosure provides designed heterodimer proteins according to this aspect
formed by the non-covalent interaction of two different alpha-helix-containing
polypeptides (monomer A and monomer B).

30 By doubling the interaction surface area of protein coiled coils with an
additional helix, and incorporating modular hydrogen bond networks, a wide range of
heterodimeric interaction specificities can be achieved, as described herein. Millions of
helical backbones with varying degrees of supercoiling around a central axis were
generated and searched for those accommodating extensive hydrogen bond networks,

followed by connecting the helices with short loops and designing the remainder of the sequence. As disclosed in the examples that follow, designs expressed in *E coli* exclusively formed heterodimers, and crystal structures of exemplary designs fit the computational models and confirmed the designed hydrogen bond networks. Following
5 mixing of independently expressed and purified heterodimer designs, the vast majority of the interactions observed by native mass spectrometry were between the designed cognate pairs. The large sets of orthogonal polypeptide heterodimers disclosed herein can be used, for example, to generate synthetic protein logic gates, transcriptional networks and other synthetic biology applications.

10 Heterodimers are generally more useful than homodimers in bioengineering because of their ability to bring together two different entities (often fusion proteins). A long standing challenge in the field has been to come up with a set of orthogonally interacting protein heterodimers -- monomers that selectively form cognate pairs and in the meantime avoid binding to other non-cognate monomers. Disclosed herein include
15 such sets of orthogonal heterodimers, which can be programmably expanded into an even bigger set. The ability to bring together two different fusion proteins via genetically fused heterodimers allowed the design of protein-based logic gates, as also disclosed herein.

In one embodiment, monomer A and monomer B have their interaction
20 specificity determined by at least one designed hydrogen bond network at the interface between monomer A and monomer B. In some aspects, (i) monomer A comprises 1 helix, and monomer B comprises 1 helix; (ii) monomer A comprises 1 helix and monomer B comprises 2 helices; (iii) monomer A comprises 1 helix and monomer B comprises 3 helices, (iv) monomer A comprises 1 helix and monomer B comprises 4
25 helices; or (v) monomer A comprises 1 helix and monomer B comprises 5 helices. In some aspects, (i) monomer A comprises 2 helices, and monomer B comprises 1 helix; (ii) monomer A comprises 2 helices and monomer B comprises 2 helices; (iii) monomer A comprises 2 helices and monomer B comprises 3 helices, (iv) monomer A comprises 2 helices and monomer B comprises 4 helices; or (v) monomer A comprises 2 helices
30 and monomer B comprises 5 helices. In some aspects, (i) monomer A comprises 3 helices, and monomer B comprises 1 helix; (ii) monomer A comprises 3 helices and monomer B comprises 2 helices; (iii) monomer A comprises 3 helices and monomer B comprises 3 helices, (iv) monomer A comprises 3 helices and monomer B comprises 4 helices; or (v) monomer A comprises 3 helices and monomer B comprises 5 helices. In

some aspects, (i) monomer A comprises 4 helices, and monomer B comprises 1 helix; (ii) monomer A comprises 4 helices and monomer B comprises 2 helices; (iii) monomer A comprises 4 helices and monomer B comprises 3 helices, (iv) monomer A comprises 4 helices and monomer B comprises 4 helices; or (v) monomer A comprises 4 helices and monomer B comprises 5 helices. In some aspects, (i) monomer A comprises 5 helices, and monomer B comprises 1 helix; (ii) monomer A comprises 5 helices and monomer B comprises 2 helices; (iii) monomer A comprises 5 helices and monomer B comprises 3 helices, (iv) monomer A comprises 5 helices and monomer B comprises 4 helices; or (v) monomer A comprises 5 helices and monomer B comprises 5 helices.

Any suitable amino acid linkers can be used to separate the alpha helices in each monomer. The length and amino acid content may vary based on an intended use, and can be determined by one of skill in the art based on the teachings herein. The polypeptide monomers may include any other useful sequences, including detectable tags and purification tags. In one non-limiting embodiment, at least one of monomer A and monomer B comprises a hexahistidine tag.

In another embodiment, the disclosure provides heterodimers, comprising:

(a) a monomer A polypeptide, wherein the monomer A polypeptide is a non-naturally occurring polypeptide comprising 1-5 alpha helices, wherein adjacent alpha helices may optionally be connected by an amino acid linker; and

(b) a monomer B polypeptide, wherein the monomer B polypeptide is a non-naturally occurring polypeptide comprising 1-5 alpha helices, wherein adjacent alpha helices may optionally be connected by an amino acid linker;

wherein the monomer A polypeptide and the monomer B polypeptide non-covalently interact to form the designed heterodimer protein.

In one embodiment, the monomer A polypeptide and the monomer B polypeptide have their interaction specificity determined by at least one hydrogen bond network at the interface between the monomer A polypeptide and the monomer B polypeptide. In another embodiment,

(i) the monomer A polypeptide comprises 2 alpha helices, and the monomer B polypeptide comprises 3 alpha helices;

(ii) the monomer A polypeptide comprises 3 alpha helices and the monomer B polypeptide comprises 3 alpha helices;

(iii) the monomer A polypeptide comprises 3 alpha helices and the monomer B polypeptide comprises 4 alpha helices,

(iv) the monomer A polypeptide comprises 4 alpha helices and the monomer B polypeptide 3 alpha helices;

(v) the monomer A polypeptide comprises 4 alpha helices and the monomer B polypeptide comprises 4 alpha helices;

5 (vi) the monomer A polypeptide comprises 5 alpha helices and the monomer B polypeptide comprises 4 alpha helices;

(vii) the monomer A polypeptide comprises 4 alpha helices and the monomer B polypeptide comprises 5 alpha helices;

10 (viii) the monomer A polypeptide comprises 5 alpha helices and the monomer B polypeptide comprises 5 alpha helices;

(ix) the monomer A polypeptide comprises 2 alpha helices and the monomer B polypeptide comprises 2 alpha helices; or

(x) the monomer A polypeptide comprises 3 alpha helices and the monomer B polypeptide comprises 2 alpha helices.

15 In one embodiment of any of the above embodiments,

(i) monomer A comprises a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an odd-numbered SEQ ID NO selected from the group consisting of selected from the group SEQ ID NOS: 1-290;
20 and

(ii) monomer B comprises a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an even-numbered SEQ ID NO selected from the group consisting of selected from the group SEQ ID NOS: 1-290,
25 wherein the even-numbered SEQ ID NO is the binding partner of the odd-numbered SEQ ID NO. in step (i).

The amino acid sequences of SEQ ID NOS:1-290 are provided in Table 1A. The “binding partners” are sequentially numbered (and similarly named) as shown in the Table. For example, SEQ ID NO:1 (DHD9 A) and SEQ ID NO:2 (DHD9 B) are
30 binding partners, so that if monomer A comprises the polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of SEQ ID NO:1, then monomer B comprises the polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along

the length of the amino acid sequence of SEQ ID NO:2. Similarly, SEQ ID NOS:3-4 are binding partners, SEQ ID NO:5-6 are binding partners....SEQ ID NOS:289-290 are binding partners. Those of skill in the art will clearly understand what is meant by binding partner based on the teachings herein.

5

Table 1A

Design name	Oligomerization State	Chain	Design sequence
DHD9	Heterodimer	a	GSPKEEARELIRKQKELIKEQKKLIKEAKQKSDSRDAERIWKRSREINRES KKINKRIKELIKS SEQ ID NO:1
DHD9	Heterodimer	b	PKKEAEELAESEELHDRSEKLERAEQSSNSEEARKI LEDIERTSERIEE ISDRIERLLRS SEQ ID NO:2
DHD13_X AAA	Heterodimer	a	GTKEDILERQRKII ERAQEI HRRQQEIL EELERI IRKPGSSEEAMKRMLKL LEESLRLKELLELSEESAQLLYEQR SEQ ID NO:3
DHD13_X AAA	Heterodimer	b	GTEKRLLEEAERAHREQKEIIKKAQELHRRLEEIVRQSGSSEEAKKEAKKI LEETRELSKRSLELLREILYLSQEQQSLVPR SEQ ID NO:4
DHD13_X AXA	Heterodimer	a	TKEDILERQRKII ERAQEI HRRQQEIL EELERI IRKPGSSEEAMKRMLKLL EESLRLKELLELSEESAQLLYEQR SEQ ID NO:5
DHD13_X AXA	Heterodimer	b	GSTEKRLLLEEAERAHREAKEIIKKAQELHRRLEEIVRQSGSSEEAKKEAKK ILEETRELSKRLELLREILYLSQEQQ SEQ ID NO:6
DHD13_X AAX	Heterodimer	a	TKEDILERARKII ERAQEI HRRQQEIL EELERI IRKPGSSEEAMKRMLKLL EESLRLKELLELSEELAQLLYEQR SEQ ID NO:7
DHD13_X AAX	Heterodimer	b	GSTEKRLLLEEAERAIHQKEIIKKAQELHRRLEEIVRQSGSSEEAKKEAKK ILEETRELSKRSLELLREILYLSQEQQ SEQ ID NO:8
DHD13_2 :341	Heterodimer	a	TKEDILERQRKII ERAQEI HRRQQEIL EEELEYIIR SEQ ID NO:9
DHD13_2 :341	Heterodimer	b	MSEEAMKRMLKLEESLRLKELLELSEESAQLLYEQRKANNGSETEKRLL EEAERAHREQKEIIKKAQELHRRLEEIVRQSGSSEEAKKEAKKI LEEIREL SKRSLELLREILYLSQEQQ SEQ ID NO:10
DHD13_A AAA	Heterodimer	a	MTKEDILERQRKII ERAQEI HRRQQEIL KEQEKI IRKPGSSEEAMKRSLKL IEESLRLKELLELSEESAQLLYEQR SEQ ID NO:11
DHD13_A AAA	Heterodimer	b	GTEKRLLEEAERAHREQKEIIKKAQELHKKELTKIHQQSGSSEEAKKRALKI SQETRELSKRSLELLREILYLSQEQQ SEQ ID NO:12
DHD13_B AAA	Heterodimer	a	TKEDILERQRKII ERAQEI HRRQQEIL KRSEEI IRKPGSSEEAELETRELEQ EESLRLKELLELSEESAQLLYEQR SEQ ID NO:13
DHD13_B AAA	Heterodimer	b	GSTEKRLLLEEAERAHREQKEIIKKAQELHRRTEEII RQSGSSEEAKDELRR IQEETRELSKRSLELLREILYLSQEQQ SEQ ID NO:14
DHD13_4 :123	Heterodimer	a	TTKRYLEEAERAHREQKEIIKKAQELHRRLEEIVRQ SEQ ID NO:15
DHD13_4 :123	Heterodimer	b	GSSEEAKKEAKKILEETRELSKRSLELLREILYLSQQVNDVDEKALERQRK II ERAQEI HRRQQEIL EELERI IRKPGSSEEAMKRMLKLEESLRLKELL ELSEESAQLLYEAR SEQ ID NO:16
DHD13_1 :234	Heterodimer	a	EAMKRMLKLEESLRLKELLELSEESAQLLYEAR SEQ ID NO:17
DHD13_1 :234	Heterodimer	b	TTKRYLEEAERAHREQKEIIKKAQELHRRLEEIVRQSGSSEEAKKEAKKIL EEIRELSKRSLELLREILYLSQQVNDVDEKALERQRKI I ERAQEI HRRQQE ILEELERI IRKPGS SEQ ID NO:18
DHD15	Heterodimer	a	TREELLRENI ELAKEHI EIMREI LELQKMEELLEKARGADEDVAKTIKEL LRLKKEI IERNQRIAKEHEYIARERS SEQ ID NO:19
DHD15	Heterodimer	b	GTERKLLERSRRLQEESKRLLEDMAEIMRRIKLLKARGADEKVLDELRK I IERIRELLDRSRKIHESSEIAYKEE SEQ ID NO:20
DHD20	Heterodimer	a	GDRQELIRNIELKEHIKILEEISQLIEELLEDKSSSEEVVKRYKKIL ERYQQLLRKSQEIHKESSEI AKKES SEQ ID NO:21

DHD20	Heterodimer	b	GDEQKLIERSQRMQKESLELLKKEI IKLDTIEKLLDKPDS EELLDTIKKLH DTLKKIHDRNKKLLKEHEEILRQRSGSLVPR SEQ ID NO: 22
DHD21	Heterodimer	a	DKEEYKRLLEDEIKELKESKEVLKDSKRVLEDIKRKVPDDDLVKLLEKHV RLLEEHVKLLEQLIREAEKSSK SEQ ID NO: 23
DHD21	Heterodimer	b	QGSSAEELLKKIKESEKKIRDSL RKIKEI IKKS RKEGVDDKQLDLIRKVV E SHRDLRLHRDLLRLLREETS SEQ ID NO: 24
DHD25	Heterodimer	a	DIDESIKEVEKLEEEVQSLQKLD DSKKLLKLVNQPDPVDDSVRKIVKRH VEILKRHEEVLKRLIEVVKHEHTKTVK SEQ ID NO: 25
DHD25	Heterodimer	b	GSDREEVHKEIVKLIREI IKIHKKI LKIH EKIKNGEIDPSEILKLS EEEK LTDTI IKI IEDLEQLTRDLRR SEQ ID NO: 26
DHD27	Heterodimer	a	DRKEIVKRHQKVV ELLKESKLLRESSKLLQRLLDKTDGDNLQKAVDDQDK AIKRQETAIRKSQEASKKLD SEQ ID NO: 27
DHD27	Heterodimer	b	DNSEI IKKVAKTSR EVAEYSE RVAKENDKVVKTLEEGKID ESELLR LLEES IKIFDTALKLHEEAYKLHQDLVRKVS SEQ ID NO: 28
DHD30	Heterodimer	a	DESEASVAIESVQILVSVK LLEESVRILLDAVKNGVEDLLRVAQRWEK LVDEWLKVVKRWLDNVRDIQR SEQ ID NO: 29
DHD30	Heterodimer	b	GSDKAEVEKSVRKIEESI KKRKSI KKAEDAVQLLKEGKIDAKDFLRIVR EDLEVVKEDVEIVKEDVENVREFSS SEQ ID NO: 30
DHD33	Heterodimer	a	SDKEVSDKLLKASKLLK VSEELLEVRRLKALKDDELIKKIADLLRKII DKDKKFIRTSEEIVKESR SEQ ID NO: 31
DHD33	Heterodimer	b	GSDLKEVLKTV EAVKEI IKSSEELLQISRKILEISR VGDEHEYISAIRE YLKALEKHIQILKKFIEILKELIRAVS SEQ ID NO: 32
DHD34_X AAXA	Heterodimer	a	SKEEIDKIVKHKKKI EHKKKVDELK LKLV EEHDKRVSQDKDDKVKKLSEE VKKIIKRLEEVSKRLEEVSKLLKVI SDKR SEQ ID NO: 33
DHD34_X AAXA	Heterodimer	b	GSNDEELKKI LETLDRI LK KLDKILTR LIEVLKKS EDPNLDDKDYTEL VKQ FIELIKKYEEVVKYEEVVRQLIRLFS SEQ ID NO: 34
DHD34_X AXXA	Heterodimer	a	SKEEIDKIVKHKKKI EELKLVDELK LKLV EEHDKRVSQDKDDKVKKLSEE VKKIIKRVEEVAKRLEEVSKLLKVI SDKR SEQ ID NO: 35
DHD34_X AXXA	Heterodimer	b	GSNDEELKKI LETLDRI LK KLEKILTR LIEVLKKS EDPNLDDKDYTEL VKQ FIELIKKFEVIKYE E VVRQLIRLFS SEQ ID NO: 36
DHD34_X AAAA	Heterodimer	a	SKEEIDKIVKHKKKI EHKKKVDEHKKLV EEHDKRVSQDKDDKVKKLSEE LKKISKRLEEVSKRLEEVSKLLKVI SDKR SEQ ID NO: 37
DHD34_X AAAA	Heterodimer	b	GSNDEELKKI LETLDRI LK KLDKILTR LDEV LKKS EDPNLDDKDYTEL VKQ YIELVKKYEEVVKYEEVVRQLIRLFS SEQ ID NO: 38
DHD36	Heterodimer	a	DHSRKLKEIDLRLRKHVKRLKEHLDELRLDVRQVPEDKLEHVVKLSDKIL QISERAVREFTKSVDKDS SEQ ID NO: 39
DHD36	Heterodimer	b	GSDKKDELERILDEIRRLIERLDEILSR LNKLEL LKHGVPNAKEVVKDYI RLLEKEYELVKEFLKLVKRHADLVS SEQ ID NO: 40
DHD37_A BxB	Heterodimer	a	DSDEHLK LKKTFL ENLRRHLDR LDKHIKQLRDI LSENPE DERVKDVIDLSE RSVRIVKTVIKIFEDSVRKKE SEQ ID NO: 41
DHD37_A BxB	Heterodimer	b	GSDDKELDKLLDTLEKILQ TATKI ID DANKLLEKLRRSERKDPKVVET YVE LLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 42
DHD37_B BBB	Heterodimer	a	MDEEDHLK LKTHLEKLERHLK LLEDHAKKLEDILKERPDSAVKESIDEL RRSIELVRESIEIFRQSV EEEE SEQ ID NO: 43
DHD37_B BBB	Heterodimer	b	GDVKELTKILD TLTKI LETATKVIK DATK LLEEHRKSDKDPRLIETHK KL VEEHETLVRQHKE LAEEHLKRTR SEQ ID NO: 44
DHD37_X BxB	Heterodimer	a	DSDEHLK LKKTFL ENLRRHLDR LDKL LKELRDI LSENPE DERVKDVIDELE RVIRIVKTVIKIFEDSVRKKE SEQ ID NO: 45
DHD37_X BxB	Heterodimer	b	GSDDKELDKLLDTLEKILQ TATKI ID DLNKVLEKLRRSERKDPKVIETVVE LLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 46
DHD37_A XXB	Heterodimer	a	DSDEHLK LKKTFL ENLRRLEDL LDKHIKQLRDI LSENPE DERVKDVIDLSE RVVRTVKTVIKIFEDSVRKKE SEQ ID NO: 47
DHD37_A XXB	Heterodimer	b	GSDDKELDKLLDTLEKILQ TATKVVDDANKLLEKLRRSERKDPKVVET YVE LLKRLEKLIKELLEIAKTHAKKVE SEQ ID NO: 48
DHD37_3 :124	Heterodimer	a	DSDEHLK LKKTFL ENLRRHLDR LDKHIKQLRDI LSEN SEQ ID NO: 49
DHD37_3 :124	Heterodimer	b	EDERVKDVIDLSERSVRIVKTVIKI FEDSVRKLEKTKPDSKTAKE LDKLLD TLEKILQ TATKI ID DANKLLEKLRRSERKDPKVVET YVELLKRHEKAVKEL

			LEIAKTHAKKVE SEQ ID NO: 50
DHD37_1 :234	Heterodimer	a	DSDEHLYKLTFFLENLRRHLDRLDKHIKQLRDI LSENPEDEVRKDAIDLSE RSVRIVKTVIKIFEDSVRKKERPIDKRDDKELDKLLDTLEKILQTATKIIDANKLLEYLRR SEQ ID NO: 51
DHD37_1 :234	Heterodimer	b	GDPKVVETVVELLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 52
DHD37_A XBB	Heterodimer	a	DSDEHLDRLDKHLKLLKLTFFLENLRRHIKQLRDI LSENPEDEVRKDVLDLSK TVIKIFEDSVRKKERSVRIVE SEQ ID NO: 53
DHD37_A XBB	Heterodimer	b	GSDDKEATKI IDDLKLLDTLEKILQTANKLLEKLRRSERKDPKVVETVYK AVKELLEIAKTHAELLKRHEKKVE SEQ ID NO: 54
DHD37_X BBA	Heterodimer	a	DSDEHIKQLRDHLDRLDKHLKLLKLTFFLENLRRI LSENPEDEVRKTVIKIFE DSVRKKERSVRIVKVIDLSE SEQ ID NO: 55
DHD37_X BBA	Heterodimer	b	GSDDKEANKLLEKATKI IDDLKLLDTLEKILQTLRRSERKDPKAVKELLE IAKTHAELLKRHEKVVETVYKVE SEQ ID NO: 56
DHD39	Heterodimer	a	DHSRKLEELDRRLRKHVKRLEHLRELLSLVKENPEDKDLVEVLELSLAIL RRSLEAVEAPLKSVTKKDPDDEDLRRKADEIRKEVEEIKKSLAEVEKEIYK LK SEQ ID NO: 57
DHD39	Heterodimer	b	GSSADDVLEDILKIRELIEILDQILSLNQLLKLRLRHGVPNAKVVVEKYK EILELYLQVLSFLKIVKTHADAVSGKIDKKAEEEEIKKEEIKI KEKLRQAK DILKQLQEEIDKTR SEQ ID NO: 58
DHD40	Heterodimer	a	DRDAHLYKLLTFLEQLVVRHLDRLVKHITQLRDI VVKDPEDERAVDVIRQSV RSLIEIVITVLKIFVDSVSDAARSKEAEKIVRKIRKEIDEIRQKLEIDKEV KKTTS SEQ ID NO: 59
DHD40	Heterodimer	b	GSNDKVLDKILDILDRILRLATRVIDLANKLLQVKKKSTHKDPRIVETYKE LLKIHETAVRLLLELADLHRLKSKDEEANKRVETELDRI RKKVKDIEDKV RKLEDKVRKTAS SEQ ID NO: 60
DHD43	Heterodimer	a	NDSLKEVLKLEKSVVEELLRRVQKSVKEAQKRGLLSDELVDRHLKILNQLV KRHELLEQEVIKRSDKK SEQ ID NO: 61
DHD43	Heterodimer	b	GSDEAVKRVVEKSLKILDEVIKKSLDILRELIELQIRHAKDDES VIRASKS ALKDAIEALKKSLDEIKKALKRSDAEG SEQ ID NO: 62
DHD65	Heterodimer	a	SSEEVVVKVHEKVVKLHKEILELLKIKI HETAARDPDDKDSIKKLSDEIK KIVKRIEDISDQAKRESSDAQRKQS SEQ ID NO: 63
DHD65	Heterodimer	b	DKEESKELKLLKKEIKLRSEELLEESKELLKLAKNGEIDES ELADADRKL NKKHEKLVQDIQDLLREHERQDR SEQ ID NO: 64
DHD70	Heterodimer	a	DEKKKIDKIVKETEDLLQKSEKLLQKSKEAVKRIRSQV KENEIVDRLLRIS EELLKISRRLVEISRRIASTLS SEQ ID NO: 65
DHD70	Heterodimer	b	GSSKEEVI RLLKENVRLIKENLELLTRNLKLITDLVRGNSGSEEKIKTLKE LLKEYRELLKRYRKLVEDYKRLVDKHD SEQ ID NO: 66
DHD88	Heterodimer	a	EIQELIKSSRRIIEESKELIKES EEVLRRIKEILDRI RINGVDNQEDLLREI LKLLTKNLKIIQRNLKLLQD NAEILKRLVS SEQ ID NO: 67
DHD88	Heterodimer	b	GSYIEDVIKKILDVSRELIKLSRTI I KISEEINKQLQQGRDTKDLVKKYDE I IKKYTRIVQHYTELIKELQKLLS SEQ ID NO: 68
DHD89	Heterodimer	a	SPTEEAIQLSQRVIELSKRVI ELSKEILKLLKRVLDLLPDLKNEEKRLDD YDKELKEYDKELKKYERKLDLAS SEQ ID NO: 69
DHD89	Heterodimer	b	GSEEEELIKIQKELLRIQSEILDKQKKILDTLRSNGAVTEEVRSILEKVER LSEEAKELSKEAKELTKEVSKLIS SEQ ID NO: 70
DHD90	Heterodimer	a	SPLKELNNQLRLLRELVKVSKKIVDLSKTIIEVLKHTDLDPRLDLSLEKS QQELDKSQKELDKVVKELTKVNKKLQ SEQ ID NO: 71
DHD90	Heterodimer	b	GSPLEDLVRYKDELVKTYEKLVEEFKAVDKYKAVKKAPVSKAATDSL DL IRKVLELLDRNLKLIKENAKLIKELK SEQ ID NO: 72
DHD91	Heterodimer	a	SPTRENEKVIKENEKVISDNERVLEEVVQVETATDRKEIQDAVDEVKRSV DKL RDSVRKLEESVRTLD SEQ ID NO: 73
DHD91	Heterodimer	b	GSPKIDISKRLLEISKRLVEISDRIVELLQRIADSKDPNKDLQKEVKDVLE EYKRLVREYREVVKYEVKVS SEQ ID NO: 74
DHD92	Heterodimer	a	DEDEVKQLIKNADLLRKHAE LLKELVKLFQETASQIPDRVAKKVTDVVD RIDKILKQTEKLVRRTKQILDYSR SEQ ID NO: 75
DHD92	Heterodimer	b	GSNLEELVKLLKEVLEMHERLLRIHEDLVEAHKSNASDKESERLKKSKDK

			IKESLKKIKSIIDQVRYIQS SEQ ID NO:76
DHD93	Heterodimer	a	PVEDIIIEESLRLLEESLKLNLRLKLLLEDLSLRKLPRSEEWQRLEDEFRRKL EDWKEELERWIEDVRYKKT SEQ ID NO:77
DHD93	Heterodimer	b	GSDEYESREIIDEIRKLLDRSKKIVHRSQRLVERVKSTPLSEDQEDLI RR HEETINRHRELVKLEKVLLEDHERHIR SEQ ID NO:78
DHD94	Heterodimer	a	PEEDSRRVLERFVRVSREVLKVLLEEFLLRVSEELLREADRDRDRRLEEYERQ VDELREEIRRYKKEVDKFDKEVKYK SEQ ID NO:79
DHD94	Heterodimer	b	GSPEKDENRKLKLDKVRKLVKESRRLVEELRKLVDQSTKNGLIDEKALRKQQ EVLKRVVEVLEKQERVLRELEEISYRVI SEQ ID NO:80
DHD94_3 :214	Heterodimer	a	GSPERDENRKLKLDKVRKLVKESRRLVEELRKLVDQSTKN SEQ ID NO:81
DHD94_3 :214	Heterodimer	b	GSDEKALRKQQEVLKRVVEVLEKQERVLRELEEISYRVITRGEDHKAEDS RRVLERFVRVSREVLKVLLEEFLLRVSEELLREADRDRDRRLEEYERQVDEL EIRRYKKEVDKFDKEVKYK SEQ ID NO:82
DHD94_2 :143	Heterodimer	a	GSDRRLEEYERQVDELREEIRRYKKEVDKFDKEVKYK SEQ ID NO:83
DHD94_2 :143	Heterodimer	b	GSPERDENRKLKLDKVRKLVKESRRLVEELRKLVDQSTKNGLIDEKALRKQQ EVLKRVVEVLEKQERVLRELEEISYRVITRGEDHKAEDSRRVLERFVRVS REVLKVLLEEFLLRVSEELLREADR SEQ ID NO:84
DHD95	Heterodimer	a	DLSEESKFFVEKVKLEKESRELEKQVKKIEEDSRVENDVQKEFLELLKR LLDIQKVVVEVLREVVKVQYVDS SEQ ID NO:85
DHD95	Heterodimer	b	GSDSEYERQVRLDVTVLKDSHTVLEALRQVIRDSQDVVSKSDEESRRVI DDLEKVIQDSKVLDDIKRLIDKSKSIKS SEQ ID NO:86
DHD96	Heterodimer	a	NEDELLKLLTENLKLLENLKLRENLSLLRQANNITDKNRIREIVKQSK IVKQSRREILKQSKKEIVERIKYIVS SEQ ID NO:87
DHD96	Heterodimer	b	GSSLYELTQRYEKLQVQYEELVKDYRRLVKKLEKLRDNKPKDRLLKEIVD VIKKSVEIIDRSLKLEESIKILEETD SEQ ID NO:88
DHD97	Heterodimer	a	SQERSLEILKRIIDLKESLEILKESLSILRQLASRIKPNRKEIEILKES DKIIKESDKVLKEIEEVIRYSS SEQ ID NO:89
DHD97	Heterodimer	b	GSDIEYESKEILELIKELLKLSRELLKESRRALVRSRDDSIVEEVIQV HKKVLDIHKVLEKIVRKVVEVHRRVKS SEQ ID NO:90
DHD98	Heterodimer	a	SKKDESTKLERLAEKIDEITKRIEELVKDVKRSSEGVKDQQQKIDEVFQ KLLDLQREILEILDRIKLVQYILD SEQ ID NO:91
DHD98	Heterodimer	b	GSLEYLNRRLLQLIKTLIDLNRHLLKLDKLLKNSREGDEEKI KESKQ IQEQFKEIVERSKEIKQIKKIKRSQ SEQ ID NO:92
DHD99	Heterodimer	a	DFERSRRLEKVVEDLRRSSDRLREVIDELRKSADKDEDEDLRRARKEHR DLIEELKRALEKQEEIKHLQELVYRQL SEQ ID NO:93
DHD99	Heterodimer	b	GSESEEVKVVVERIKKISRELEEVKELDRVSKFDRHGETDEIVREHER IVEKLEEVKHTKIVEELAEIVYKQ SEQ ID NO:94
DHD100	Heterodimer	a	SDDSVRVLDEIVKILDESVKLLKESLKLDDFLRTKPDHLKEVVKESK VVEQSKKVLDRIKKIYESK SEQ ID NO:95
DHD100	Heterodimer	b	GSDLLYLSKELLKLVRELLKLSRELVELSRRLVNSTHKSPELVKKYDKLVK KYQDLLKKLADVADEYLRQRS SEQ ID NO:96
DHD101	Heterodimer	a	DEKDYHRRLEHLEDLVRREELIKRQKVVVEELERRGLDERLRVVDVDR RSSERWEEVIERFRQVVDKLRKSVE SEQ ID NO:97
DHD101	Heterodimer	b	GSDAYDLDRIVEKRRVVEEQRELVEELEKLVRRQEDHRVDDKESHEILER LERIIRSTRILTELEKLTDEFERRTR SEQ ID NO:98
DHD102	Heterodimer	a	DERYRAREHTRRVEEHTKRLRHILKRLREHEEKLRRELKPGDEITESVDRF KKIVDQFEESIKKFETVSEELRKS SEQ ID NO:99
DHD102	Heterodimer	b	GSDRQRIIDLRLDKILEKLDDILKLLKDILETLSKDDVSDRRHKDLVEKFRE LVDTHHKLVERYRELQYQNR SEQ ID NO:100
DHD102_ 1:243	Heterodimer	a	GSDEITESVDRFKKIVDQFEESIKKFETVSEELRKSIS SEQ ID NO:101
DHD102_ 1:243	Heterodimer	b	GSDPQRAADRLDKILEKLDDILKLLKDILETLSKDDVSDRRRAKDLVEKFRE LVDTHHKLVERYRELVYTATAGSDLARELIRRVEEHTKRLRHILKRLREHE EKLR SEQ ID NO:102
DHD103	Heterodimer	a	NADDQLATSIKKLEDSIDQLIKIVRKFEEVKKLQKHGVDQHHEVILRKIV

			EIFRQHIEKLLKKHLEKLRYS SEQ ID NO:103
DHD103	Heterodimer	b	GSDKEYLVTEHEKLVREHEKIVSEIEKLVKKHEAGVDESELEELKLVKEL LRKLEILEQLTQLLRKTE SEQ ID NO:104
DHD103_1:423	Heterodimer	a	GSDQHVVEILRKIVVEIFRQHIEKLLKKHLEKLRYS SEQ ID NO:105
DHD103_1:423	Heterodimer	b	GSDAEYLVTEHEKLVREHEKIVSEIEKLVKKHEAGVDESELEELKLVKEL LRKLEILEQLTQLLRKAEKHIDKHSKAADQLATSICKLEDSIDQLIKIVR KFESVKKLQKH SEQ ID NO:106
DHD104	Heterodimer	a	DEDDDIRRVLDSESRVLEHSRRVLRSEEVLEKASRKKEKDETEIEKHLKR LREHAKKLEKRRRELDDEFLYKEI SEQ ID NO:107
DHD104	Heterodimer	b	GSRDKYLLERLNDILKKLDEIVDKLSDILKRLKDVRRHDDLQELVERYKEI VKEYKRIVEEYKLVREFEEQQR SEQ ID NO:108
DHD105	Heterodimer	a	DRDYEDKEFKKIKKELEDVQEELKKLQEKIKRFSSELEEPNELLKEQLKVN EEQLEVNKKILKILRDQLKQNE SEQ ID NO:109
DHD105	Heterodimer	b	GSDAEYKRVESVRSKESVKSSEDVVDKLNKSVKLSSEGHSDAEKASRELV KLVREVVLSREVILKSEKVLRVIS SEQ ID NO:110
DHD106	Heterodimer	a	DLQYKQEKLI RHFDRVREWDKLVRFKSKVLEKQKHESKDKELEEASRRVD ELIKRLREQLKRSKEILRRLKELSRKSS SEQ ID NO:111
DHD106	Heterodimer	b	GSDWEELLRLEKVLQYEEIVKELIDLIERLIKVSEDKSKDASEYKLV ELEKLISKLEEISKKLEELVKEYEYKTE SEQ ID NO:112
DHD107	Heterodimer	a	DAKDELEKSLQEI EESLKEKLLLEELDKSRLRELTSSQGNKKLEEHKVKQ KFI ELVKKYIKAVQDYLKEVRYDNS SEQ ID NO:113
DHD107	Heterodimer	b	GSDKERAARATEEMVLTKKLLKAVEDLVRDVRRLKEGLISEKHARIAET ILEVFKKHAKIKKHVDIVKYDES SEQ ID NO:114
DHD108	Heterodimer	a	GSPLKERLLEIQRDLDRLVLEEVVERLLRIQERLDSVVERKPPDVHEEYKYI VDEIREIVERVVREYEEIVKRIDEVR SEQ ID NO:115
DHD108	Heterodimer	b	GSEEDERIRYDLRIRKDVRRKLEEI RQVRLEKLLRDAGHRRDEKELLR ELIETS KDILRLVEELLKIIDKSEDLRRTKTE SEQ ID NO:116
DHD109	Heterodimer	a	GSDEEDYINENVEKDVREI EDDVRRINERIRELLEKIRTEEVLQRVLEEH ELVERVLRKLVLEILRKHEENR SEQ ID NO:117
DHD109	Heterodimer	b	GSDEEYKELKHLKLLREIEELLKHYRELVRREELVKGELDKDAAHIL ERLSELLERIRRAVHTLRRLSEERR SEQ ID NO:118
DHD110	Heterodimer	a	GSDEDEISYDKRRVVEIVRQAREKSEKSRKDI EDVAEVLKRGDVSEKEVV DELVKVLEEQVKVLEAVERLEVLKQVDDVR SEQ ID NO:119
DHD110	Heterodimer	b	GSDIVELVDHLLKRSKLLLEELAEVLRRLLEKSTELKRRTEEHKEEVVEE SEYVRELEERLRRVDESEKLVDRADKHIR SEQ ID NO:120
DHD111	Heterodimer	a	GSKEKDIVKTLDVLLRENLETLERLIEEVVRLKENVDVDRDEGRDDKDSER ILRDIKRRIDEAAKESREI IERIEKEVEYRSR SEQ ID NO:121
DHD111	Heterodimer	b	GSPKLDKLELRLVREILKASEELLRLRLKLI DEALKLSEKRRKDSQYREVVD RVKLELRLLEDEYRKLVEELKEKLRDTR SEQ ID NO:122
DHD112	Heterodimer	a	GSDKRYESEKLRRLDEAVEKRVVVERVERESDRVLEEVRRRESKEVVD KVI EDNDKALEDVLRVDEVAKVVRDVVRENTR SEQ ID NO:123
DHD112	Heterodimer	b	GSPREYHSDILRVDEILERIRRHADRVKKSERLRENVDVNEHSDVK RVIRELLELVKELLRLAKKHSDDQOE SEQ ID NO:124
DHD113	Heterodimer	a	GSDEDEILYHSERLLQKLLKELDDLKEKSRELLLEELKKEDPDDRLIERIIR LHDEVLKDLDVLEKNI LEVHREVLRLR SEQ ID NO:125
DHD113	Heterodimer	b	DKLDRLLKIHAEALRAEELIKRLLDIHRRALDLARRGELDDYLLKESERE LREIIRRAEELKESRDRLEEISR SEQ ID NO:126
DHD114	Heterodimer	a	GSPKEELIRVLEEVKRLNEKLEIIRRAEELVCRANDELPETEKLEIDR ELEKLLKEIEDELRRIDKELDDALYEIED SEQ ID NO:127
DHD114	Heterodimer	b	GSPKLDKLELRLNLEKLEI EELVLRITNLERLREDIRDEVDLQEYE RLIRKAEEDLRRVLEKEDDLLKLVYELR SEQ ID NO:128
DHD115	Heterodimer	a	GSKEDSVKRAEIVRTLLKLLLEDSLREASLRDIKNGEDEHNLRRRISEK LEELSKRITETIERLLRELQYTSR SEQ ID NO:129
DHD115	Heterodimer	b	GSPNQELLDVRVKILEDDLRLNEELVRLNKEKLRALMRRKNRDSEEVLE RLAEYRKRLEEYRRELEKLEELLETIYRYKR SEQ ID NO:130
DHD116	Heterodimer	a	GSDESEEAQHEVEKVLDDIRRLSEHLQKRLVEEVLEEVYELRREGSDRTEVV

			ELLKEVIREIVRVNREALERLLRVVEEAVKRNE SEQ ID NO:131
DHD116	Heterodimer	b	GSDEEELVETVKRIQKEILDRLTELAKLLVEIQREIKKLDGEDDKELKR LSDELEEKVRQVVEEIKRSLDELEETVEYVSR SEQ ID NO:132
DHD117	Heterodimer	a	GSDEEEVVRRAEELVKEHEELIERVIRTHEELVYKLEDQGNADKKLVDVLK RVVEESERVAREIVKVSRELIRLLEEASR SEQ ID NO:133
DHD117	Heterodimer	b	GSSKEEILKELEDLQRRLEELKKLQERVVLEELIKRLRDRGRDDKHLK RLVKEVRRLLSEEVLRSIKEVSDRVRYQLR SEQ ID NO:134
DHD118	Heterodimer	a	GSDKEEESYLLRDLVRLLEKVKKEIEEVNREVEKLLKVKDGRDLDRREVL REILRLNRELAIEIKVVDRI RHVVERSER SEQ ID NO:135
DHD118	Heterodimer	b	GSDLHEVVYETKELKRIEVEELRKKSEDIIRKAERGEISEDELKRLQE EIAAREAKLLDEIKRVLERHLEQTL SEQ ID NO:136
DHD119	Heterodimer	a	GSPVEEIIKEVVKRVI EVQEKVLR IISHAVKRVVEVQKKYDPGSEESNRVV EEVKKTIEDAIRESDEVVDEVVKRIQYTVR SEQ ID NO:137
DHD119	Heterodimer	b	GSPEQEIADRLTEIRESQKELERLARKILKLLDESQEKAKRGRLLSEESD ELLERIKKELDELLERSKELKKIEYELR SEQ ID NO:138
DHD120	Heterodimer	a	GSDEDEKANRVLDEVLKTVRDLLETANEVLEKLVYRLKRTDDQEKVVRTLT EVLKEHLKLVEEIVRILDKVLKEHLETEK SEQ ID NO:139
DHD120	Heterodimer	b	GSPEDDVLRRLEEVSEKILRVAEDVARQLREVSEKITQGVDRKEWEEDIK RLKRELEELLREWKEEIERLTYELR SEQ ID NO:140
DHD121	Heterodimer	a	GSRRREVVKRIRELKRNKELIDRIRELLEENEYLDKDKARDKDLRRSVEL LEELVRILEESVELAKEI IKLREVVE SEQ ID NO:141
DHD121	Heterodimer	b	GSDEKEDNRRLQHKIERILEKNEDLQRKLEEILELLERGEADEEKIDRLRK AVEDYRRVVEEIKEDVKRHKYTVR SEQ ID NO:142
DHD122	Heterodimer	a	GSDEKEEAKKASEESVRTVERILEELLKASEESVELLRGEDAKDVVERSK EALKRVKELLEDVVKRSDEILKYIHN SEQ ID NO:143
DHD122	Heterodimer	b	GSDEKKLINEVVETQKRLIKEAAKRLSEVVRHQTELIRELREKNVDDKDV KLLKESLDLAEIIVRRIKELDESCKLVEYVSN SEQ ID NO:144
DHD123	Heterodimer	a	GSPDMDEVKRVLDLEIEIQEELREIKRVLEKLIKIQEDNGSEYESREVVR EIVEIARKLVRSRRVVKITETLQ SEQ ID NO:145
DHD123	Heterodimer	b	GSDEYATREIVERIERIAREILKRTEEIVREVREVLSDVDQEEVVRRLA DLLRESVELVQHLVRRVEELLQESVERKK SEQ ID NO:146
DHD124	Heterodimer	a	GSPEREALREVLEDLKRVTDRLELVERVLEELKVKVDHVDSERILRESRR VLKELKDIIEEILRESEKVLKLYTED SEQ ID NO:147
DHD124	Heterodimer	b	GSPAREILEEVVKKHLEVVEDAARI LEEI IREHEKAVREDRDKKELEEISR DLLRKAREALKVKDISDDLSREIEYVAS SEQ ID NO:148
DHD125	Heterodimer	a	GSPVEEAIKKVIDDLRDVQRKIRELVEELIRLLEEVQRDNKRESEYVVER VEEILRRI TETSREVVKAVEDLS SEQ ID NO:149
DHD125	Heterodimer	b	GSDSDEKAAYLLKEMERVRESDEVVKKILRDLEEVLERLRGEISEDVVT EILKELAEHRIRAI EELVRRLELLEHRKR SEQ ID NO:150
DHD126	Heterodimer	a	GSPVEEVLKELSEVNERVRDIAREI IERLSEVNEEVKETE DDEDELKKISKK VVDEVEDLLRKILEVSEEVRRVEYHDR SEQ ID NO:151
DHD126	Heterodimer	b	GSPKEDILREVLRRHKEIVREIVRLVREAVETHLELVKRNSDDRDAQDVIR KLEEDLERLVRHAQEVIEEIFYRLH SEQ ID NO:152
DHD127	Heterodimer	a	GSPRSYLLKELADLSQHLVRLLELRLVRESERVVEVLERGEVDEEELKRL LHRELEKAVREVRETHREIRERSR SEQ ID NO:153
DHD127	Heterodimer	b	GSDREYTIKIDILDSQEHLLRLIEELLEETQKELLEILKRRPDSVERVRELVR RSKEIADERRQSDRNVRLLEEVSK SEQ ID NO:154
DHD128	Heterodimer	a	GSDEKDEIRHVIESVERLIEDIKRLLKTLRELAHDDSDKKTVKEVLDRVKE MIERHRLEEBHRKELEPRAEYEVV SEQ ID NO:155
DHD128	Heterodimer	b	GSESEDRIKELLKRHIELVERHEELLHEIKKLI DLEEKDDKDREEAVKRID DAIKSEEMLEESKTEIEEIEYLN SEQ ID NO:156
DHD129	Heterodimer	a	GSSLEDSVRLNDEVVKVVERVRLNQEVRRLIKHATDVEDEETVKYVLERV REVLDRESREVLKRVHELLEESERRLE SEQ ID NO:157
DHD129	Heterodimer	b	GSHEKDIVYKVEDLVRKSDRIAERAREIVKRSRDIMREIRKDKDNKKLSDD LLKVTDRDLQRVVDELEELSRELLRVAEESRK SEQ ID NO:158
DHD130	Heterodimer	a	GSPLEDEVKKLIDELKKSVERLEESI REVKESI KKLKRGDIDAEENIKLLK ENIKIVRENIKIKEIDVVQYVLR SEQ ID NO:159

DHD130	Heterodimer	b	GSDEEEIEELLRELEKLLKKSEEALESKKLIDSEELLRRDRDLKKEKHAVR ASEEHVKLSEEHRLRSREIVKILEKAVYSTR SEQ ID NO:160
DHD131	Heterodimer	a	GSDESDRIKRIVEESDEIVKESRKLAEARELIKESDKRVSEERNERLLE ELLRI LDENAELLKRNLELLKVEVLYRTR SEQ ID NO:161
DHD131	Heterodimer	b	GSDEDELEERLLREYHRVLRREYKLEELLRRLYEYKRGVSEESDRI LR EIKEILDKSERLWDLSEEVWRTLLYQAE SEQ ID NO:162
DHD132	Heterodimer	a	GSDKKDASRRAIRVLEHFEVVRVSEEVLEVLKRSVESLKRDLVDDEKIKRTHDR IEEELRRWKRELEELIERLREWEYHQD SEQ ID NO:163
DHD132	Heterodimer	b	GSDEEEDKRLLEEVKRSLDTDERILEKLRHSLEQLDVKDEDSRRVLR ELDEITKRSREVVKRLRKLAYESK SEQ ID NO:164
DHD133	Heterodimer	a	GSDKEYKLDRI LRRLDELIKQLSRILEEIERLVDLELEREPLDDKEVQDVIE RIVELIDEHLELLKEYIKLLEEYIKTKK SEQ ID NO:165
DHD133	Heterodimer	b	GSPSKEYQKSAERQKELLHHEYKLVRLHRELVEKLRQRELDKKEEVLRLV EILERLKDHLKKEIDAHKNEEAHKENK SEQ ID NO:166
DHD134	Heterodimer	a	GSRDRKISEELIKALEDHIRMLEELIRAEHIIKLAERGVDEKELRESLEE LKKIVDELEKSLEELRKLAEYKYETR SEQ ID NO:167
DHD134	Heterodimer	b	GSPKEESVEELKRVIDKHEEILRELKRVLEEHERVSHDEDENELRRSLERL KHILDRRLHESLKEHELLKKNYEYTER SEQ ID NO:168
DHD135	Heterodimer	a	GSDEHYVVKIVERILRVMEKHAEIVKHLLEIVERVVREGESEDLRKLLKES LREIEESLRELKELLDDELSEKTR SEQ ID NO:169
DHD135	Heterodimer	b	GSDEEYVTRSQRRLKRLLEEYIKVVEEHARLVERNERDDKELKRSIDELDK LTKELELVKRYKELVDKTET SEQ ID NO:170
DHD136	Heterodimer	a	GSKDEEIVKLQDEVIKTLERHLDILRKHIDLLEKLDHLSEELKERVDRSI KKLEESIKRLERIEELQELAEYSL SEQ ID NO:171
DHD136	Heterodimer	b	GSREEELKESAEELERSVRELKKEADKYKEEVDRLHYRGKVDKDWVRVVEK LIKLVVEEHLELIREHLELLKEERR SEQ ID NO:172
DHD137	Heterodimer	a	GSDMEYELKKSAAELRKSLEELKRI LDELHKS LRELRRHGDEEYVQTVVE LRKELEEHAKKLEHLKELEERVAT SEQ ID NO:173
DHD137	Heterodimer	b	PEYELKKSVDLDRKRDVDRLVVEEVEVFELSKERLREDRKHLELVEEMVRLI EKHLELIKEHLKLADHVR SEQ ID NO:174
DHD138	Heterodimer	a	GSREKDESKELNDEYKLLLEEYERLLRRSEELVKRAKGPDEKELKRILEE NEDILRRTKEILERTKEISSEQKYRRR SEQ ID NO:175
DHD138	Heterodimer	b	GSDKDERQERLNEESDKSNEESERSNRESEELNRRARGPNDEKELQEILDR HLELLERNQRLLDENKEILRESQYLN SEQ ID NO:176
DHD139	Heterodimer	a	GSENKYILKEIKLKLRENKLHLHDILRLLDENLEELEKHGAKLDLDYRRKI EEIRKKVEDYREKIEIEIKKVERDR SEQ ID NO:177
DHD139	Heterodimer	b	GSESEYTOEIELELLKESIKLLREILRLLEESEELWRRENTKSERSEEI KE RAKEAIKRSEEILERVKRLSDHSR SEQ ID NO:178
DHD140	Heterodimer	a	GSDEEEANYVSDKAVKIAEDVQELLKELLESEVVRGEVDEDEYDRVLRK LQEVMKKEYEVLKEYEEVSRKHE SEQ ID NO:179
DHD140	Heterodimer	b	GSPKYLIKTOEELLRRHAETLEDLIRKVERQVDLRRKVDREDEDLKRELE RSLRELERLVRESSRLVEEIRELSKEIKR SEQ ID NO:180
DHD141	Heterodimer	a	GSDEEYELERISRESKELLERYKRLREYQELLKELRHVKDLDRAVKIIHE LMRVSKELVEISHRLELHERLVRRK SEQ ID NO:181
DHD141	Heterodimer	b	GSEKEYIEKLSRKIEEDIRREERAKDSERLVRRLEELAKRKRRLDDVLR VAENLEILEDNLRILEEILKEQDKSNR SEQ ID NO:182
DHD142	Heterodimer	a	GSPHEEVVELHERVMEISERAVELIQRIIDIRRIREDDKDIEKLVKTI RD LVREYEELHRELEEIDEEIYKKE SEQ ID NO:183
DHD142	Heterodimer	b	GSDHEDVRLHEDLVKQEDARRVLEETVRLAEIIVEIKKDEKDKDRVTR LVEEIEKLVVEYKVVDEMRKISDEIKYRSR SEQ ID NO:184
DHD143	Heterodimer	a	GSRAREVVKRAKRIIEEWQKILEEWRRILEEWRRLLEDERVDDRDNERIIR ENERVIRENEKIIDVIRLLEELLYERR SEQ ID NO:185
DHD143	Heterodimer	b	GSREDEELEEEDIRIQMVEEYELVKEYEELTEKYKQGVKDEESKIIIE KSERLLDLSQDAVRKVKIEIRRI LYTNR SEQ ID NO:186
DHD144	Heterodimer	a	GSPKEEIVKLHDESALHRRSVEVADEIKMHRSKDVDDERESRELSKEI ERLIREVEEVSKRIKRLSEEVEYLV SEQ ID NO:187
DHD144	Heterodimer	b	GSPLEELIKTQRRINKIQDDINKILHEILRMQEKLNRS SDKVEEESLRRI RELKRIKDLKSEIEDLSREVKYRTT SEQ ID NO:188

DHD145	Heterodimer	a	GSPEDEHVYVREIYEVLRHEAEVLEENREVIERLLEAKKRGDKSEELVKE LKKSIDDKLEISRKLEEVKLEKVSSEKLLK SEQ ID NO:189
DHD145	Heterodimer	b	GSDEDETSYRILELLREIVRASRELI RLSEELLEVARDDKDETVLETLIR EYKELLDYRRLIEELTRLVVEEYERSR SEQ ID NO:190
DHD146	Heterodimer	a	GSTQEEINRIQHEVLRIQEEI DEILRDIVEKPKAISRGELDHEVVKDVEDK VREALEKSEELLDKSRKVEYKSE SEQ ID NO:191
DHD146	Heterodimer	b	GSDEEELNREELLEKSKRLVDINRDII RTAQELI EMLKDSKGRVDEDTKRE LRDKLRKLEEKLEVRREELRKYEELLRYVQR SEQ ID NO:192
DHD147	Heterodimer	a	GSDEKDRVYEILKEVQRLVKEYRDISKEIEDLVKHYEHITDDEAQEVSKEL IDKSLRASEIVRELI RLKELLDLELE SEQ ID NO:193
DHD147	Heterodimer	b	GSDEEDVLYHLRELEELKRVSDDYERLVREIKETSERKDRDTKENKMDLD ELVKAHREQEKLLELRVLRLEELFERKR SEQ ID NO:194
DHD1	Heterodimer	a	PREQAIRISEEII RLSKKI IEILERTRS STAREAMKWAKDSIRLAEESKYL LDK SEQ ID NO:195
DHD1	Heterodimer	b	IEDDVKKIQDSTKKAQKETIEALERSTSS TARKQMEEQKEQIRLQKEAMYL LKK SEQ ID NO:196
DHD2	Heterodimer	a	SREEIAKLQEEVIKLRQRRVIELQKEVIELQRRAKELTSSYTKELI EQRR EEIQREIEEIQKRIEIEIQEEIQRRT SEQ ID NO:197
DHD2	Heterodimer	b	SDEEIKRLEEVIQLSRRVIKMSREAIKLSREVQKLTSPSYQKRIKEIADRS IELARESIEIAKRSEKIAEESQRRT SEQ ID NO:198
DHD3	Heterodimer	a	PAKDEALKMANESLELAKKSARLIQESSKEILERIEKIQRRIAELQDRIA YLIKK SEQ ID NO:199
DHD3	Heterodimer	b	PAKDEALRMIDESRELIKKSNELIQRSSSKEILERILEIQRKIAELQKRIQ YLLKS SEQ ID NO:200
DHD4	Heterodimer	a	TDEARYRSERIVKEAKRLLDEARRRSEKIVREAKQRSNSEDAKRIMEENLR ESEEAARRLREIIRRNLEESRETG SEQ ID NO:201
DHD4	Heterodimer	b	TREALEYQRKMAEEIEDLLREALRRQEEMVREAKQRSLSSEFKRIMERILE EQERVMRLAKEALERILEEQKRTG SEQ ID NO:202
DHD5	Heterodimer	a	SERTKREAKRSQEEILREAKEAMRRAKESQDHRQNRDGSNSEDLERLSQEQ KRELEEVERRLKELAREQKYKLEDS SEQ ID NO:203
DHD5	Heterodimer	b	SEDLKRIKKEITERELKLMQDLMEILKKITEDENNLDNNSSEDLKRSIEKA RRILDEALRKLEESARRAKYIQEDN SEQ ID NO:204
DHD6	Heterodimer	a	TEDEIRESLKLWDEVLQELREIARESNEVLERNRQKSRSDKLRREDIERYKK RMEEARKKLDDQLNKYKRM DENRS SEQ ID NO:205
DHD6	Heterodimer	b	TEEBLKESKFAEDLARSARRALKESKRVLEETISQASRSKKEEIVRRYKE QVQRWQDEWDERAREYRKRMMKENRS SEQ ID NO:206
DHD7	Heterodimer	a	TKTEEIERLAREIKKLSSEKVERLAQEI EELSRRVKEENSTDRELKEANREI ERAI REIEKANKRMEEALRRMKYNG SEQ ID NO:207
DHD7	Heterodimer	b	TKTEEHERLAREISKLADHRKLAKIIEELARRIKEENLTDDELREAIRKI EDALRKNKEALKIMKAAERNRYNT SEQ ID NO:208
DHD8	Heterodimer	a	TKKEESRELA RESEELARESEKLARKSLELARRAESGSEEEKRRIIDENR KIIERNREIERNKEIEYNKELIS SEQ ID NO:209
DHD8	Heterodimer	b	TKDEESLELNRESEELNRKSEELNRKSKELNDRAESSNSEEEKEILREHK EILREHLEILRRHKEILRRHKYLTSS SEQ ID NO:210
DHD16	Heterodimer	a	TREELLRENI ELAKEHI EIMREILELQKMEELLERQSS EDILEELRKIIE RRELLDRSRKIHERSEIEIAYKEE SEQ ID NO:211
DHD16	Heterodimer	b	SEDIAREIKELLRRLKEI IERNQRIAKEHEYIARERKKLDPSNEKERKLE RSRRLQEE SKRLLDEMAEIMRRIKKLLD SEQ ID NO:212
DHD18	Heterodimer	a	DRQKLI EENIKLLDKHIKILEEILRL LKKDIDLKKSSEEVLEELKKIHR RIDKLLDES KKIHKRSSEIVKKRS SEQ ID NO:213
DHD18	Heterodimer	b	DEQKLIETSQRLQEKSERLLEKFEQILREASDLYRKPDS EELLRRVEKLLR ELEKLI RENQDLARKHEKILRDQS SEQ ID NO:214
DHD19	Heterodimer	a	DRQELIRENIELLKKHIKIVKEIQKLIETFI ELLKKSSEELLRRLKKI LK RIEKLYRESQEIHKRSEIEIAKKRQ SEQ ID NO:215
DHD19	Heterodimer	b	DEERLIDKSRRELQKES EELLKELKIFKRIEELLEKPDSEELIREIKKLE TLSEIHKRNEKLARTHEEILRQQS SEQ ID NO:216
DHD22	Heterodimer	a	STRDVQREIAKAFKMAADVQKLAEEIKRHVKNVEKKNKDNDEYRKIATEL LKKATESQKLLKELLDRIKSDS SEQ ID NO:217

DHD22	Heterodimer	b	DKDDRSTSLKRVEKLIDESDRIIDKFTTLELSRNGKIDDDQYKKELKEI LELLKKYDKHVKEVEELLKRLNS SEQ ID NO:218
DHD23	Heterodimer	a	SKRKALEVSEVRRISEKVVRLDESDDLKKSYYDDSKFAELIDRHEEKI KKWKKLIKWELEIIQRHKS SEQ ID NO:219
DHD23	Heterodimer	b	SAEEFVKLSEEAVKRSKEILDIVRKQVKLVKAGVDKHEITDSLRKSEKLEI EHKELIKTHRDLRREN SEQ ID NO:220
DHD24	Heterodimer	a	SSTEILKRFKRALRESEKIVKHSRRVLKIIREVLKQKPTQAVHDLVRIIET QVKALEEQLVKLVKRIVEALERQS SEQ ID NO:221
DHD24	Heterodimer	b	DKQKEIKDILEKTRRIAEESRKIAEKFDEIIKRSTEGKIDESLTKELEELV KEVIKLSDDARTSDDLVRKES SEQ ID NO:222
DHD26	Heterodimer	a	DEDESIKLRTRKSIEETRKS LKIIKEVVELIREVLKHIKDLKKEIFERIDKI LDKYKKQVDTYDEILKEYEKKQR SEQ ID NO:223
DHD26	Heterodimer	b	SELDEQKELIKKQEKLEIEEQRLLSKIRRMFKERVKDQELLREIQVKLKR QEIIVETS KKIILDRSDKTTE SEQ ID NO:224
DHD28	Heterodimer	a	DQKEINTRIVEKLERIFKKSKEIVRQSERVISTIEKKTEDERELDLRRHV KIVREHLKLEELLKIIKEVQKES SEQ ID NO:225
DHD28	Heterodimer	b	DTEELVKRLNELLKELSKLVKEFIKI LETYRKDQTKDTSKISERVDRILKT YEDLLQKYKEILEKIEKQLS SEQ ID NO:226
DHD29	Heterodimer	a	DYARLIDQAVEVTRKVVVEVNVTVARVNDKFAHLGDEELRRVSEHLKEVSK DLQEVAKSKDAARQVK SEQ ID NO:227
DHD29	Heterodimer	b	DVSKVAEEYLQISKTLVDISRTLLEISERLVRVLRVTVADDRSEVKKAIEDS IEVLKTSEEVVRQIKRASDKLVKAIS SEQ ID NO:228
DHD31	Heterodimer	a	DAKEIQRRVVEIQTEVVKLQKKAVDIIRKII EAFNNSNIDQSLEAAKEIV KEIDKLEKLTESLLEESKLLKRSS SEQ ID NO:229
DHD31	Heterodimer	b	SAEEVVKLAKIFLELLRESIKLLKRSVDLLRKS SDPSLDKSEAEKVSREIE KVS DTS LKLSKALDVVKRALKVAS SEQ ID NO:230
DHD32	Heterodimer	a	DEKDAARKARKVSEEAKEASKKIEKALEESKRI LNTLQKQKDEQEVKVIKE HEDVLRQIEKIQKQVLEIQKEVAKLLES LD SEQ ID NO:231
DHD32	Heterodimer	b	SADDVARASEKVLRVARESAKAADKSLVVFKEVVKRGDKEAFQVVKINEE VVKINITVIRILIEVSKTAT SEQ ID NO:232
DHD38	Heterodimer	a	DEYVKTTLKQLREALASLREADKRITELVKEARKKPLSEAAKFAEAIVTH VKVVVEHVEVLRHVEVLVEAKKNGVIDKSIDNALRIENVI RLLSNVIR VVDEVLQDL SEQ ID NO:233
DHD38	Heterodimer	b	DASDVIRRIHELFEVHRLIEAVHRAIEDVAKAAQKKGLEDSEAVEILAE KELAKLSRRLAEISREIQVVTDPDDKEAVERLKEIKKIQDLDELDRDL RKLQDQLLYK SEQ ID NO:234
DHD60	Heterodimer	a	SEDKAHHDIVRVLEELIKIHDELMKISEEILKATSDSTATDETKEELKR KEAQKSDTLVKIVKELEKESRKAQS SEQ ID NO:235
DHD60	Heterodimer	b	DDEEKYRQIITREAQETSKTAKRILRDAQEISKRIRHQGVDRSEHQRLVDLL RELKIEHHLRRQEQEADTRND SEQ ID NO:236
DHD63	Heterodimer	a	DRKDKARKASEKLEVIQRWKT VADKWKMMVLDVSNGLKSQEEVARVTEEL LKIQTTELAKLLEEHAQVLQESAS SEQ ID NO:237
DHD63	Heterodimer	b	SDEESIKTQSELIKTSEELLKDVKRI DEELQKLRDDPTLDESELKRVKEW SDRVRKAKEISRKIQEIVKESKKRSS SEQ ID NO:238
DHD66	Heterodimer	a	DKDEELRKVIEKYREMVKEYRKYREYEEVIKSSKTIDKSSLISLSRKMVE LSQRVIDVSEVAKVLSRKQS SEQ ID NO:239
DHD66	Heterodimer	b	TDEERLKKQTKELKEQTKQLEKQKDLLEKISNGEISKDEIQEIIKESKIA KESQKALDSSRKALEEVS SEQ ID NO:240
DHD67	Heterodimer	a	DEKEVSKKIIKVLKDIKVVQKQVIEVSQRLASVLRADDNVVKRALEEYK ILEELRELNKEIEKLTDKYRKVTS SEQ ID NO:241
DHD67	Heterodimer	b	DSDEQTKLEKLTLEHKKRVEKLLKQTKESREVD SNKLWKS KDVKDKLS EKELQKLS DQDKAKDALESSRRKND SEQ ID NO:242
DHD69	Heterodimer	a	DAEQKLLTKLRLHQRLQLIKESLKLIEKIQSSQENQDEIRKRWREVT KKLRELIKTSEKLVRELEKSYKSS SEQ ID NO:243
DHD69	Heterodimer	b	SLRDVVRRYQELVRRYDELIKTLTEILKKYQKGAEDKDASTELVKAVRTS LKLKSKELLKLNSELLKEDS SEQ ID NO:244
DHD71	Heterodimer	a	SKEELKRKLDLKKRSDTLKELSKLKEISERNPDDKSVHRTIIRIHRE FVKNHKEIVRVIEEIVSDKS SEQ ID NO:245

DHD71	Heterodimer	b	SKQDEHDRLLKIHDKLVKQHDELLKLLTKLSRAGDSVTKKKLEELRKLQE VSKQLEESLKDADKVS KDIN SEQ ID NO:246
DHD72	Heterodimer	a	TVQSLLEQHVKIVKRSIEILERHTQILQDIARSQGVSKELVEDVERQVKEYR KEVKKLEEDLRQLSRNSK SEQ ID NO:247
DHD72	Heterodimer	b	SDSDRIEKLIRESTELLKEQQKLAKRSRELAETVESLPLTEEYLYKQOREHQ KKIEKLLKDKSEKHLLEELKRLVKSEK SEQ ID NO:248
DHD73	Heterodimer	a	DSEKRIEDILRTDLELAKRDAELVKEHIKLVKRIDLSEELKKQVEDVEKES KKLEDSSEKLVQKVRKRSS SEQ ID NO:249
DHD73	Heterodimer	b	DEEERAKDLRKYLEEQTQYYRTVTEHLRNLEKVVVEELERRGKPSSELQQIL ERSQRIYKETTEIYDTSKKLIEELDKHHR SEQ ID NO:250
DHD148	Heterodimer	a	PLEDILKRHLDKVRELVRVLSSEVNKLAKVLDLKDKRVDKELDKVLKEL EKVVEEYERAVKESRDLLRELRTRR SEQ ID NO:251
DHD148	Heterodimer	b	DKERLLEIHERIQKLLDRNLEIERLLRLLREARDIKDDDKLDKVIKRLKE LSEESKDLDKIKELKSEKELT SEQ ID NO:252
DHD149	Heterodimer	a	PEDEVIRVIEELLRIAEEVDEVHRRNVEVQEEASRVTDRELRERLNRESEE LIKRSRELIEEQKLIERLERLAT SEQ ID NO:253
DHD149	Heterodimer	b	DLEELIKEYAEVVRHHKAVRDLERLVRELANAKHASEEELKRIATEILRI VKELIRVQERLIKLSSEDSNEESR SEQ ID NO:254
DHD150	Heterodimer	a	PTDEVIEVLKELLRIHRENLRVNEEIVEVNERASRVTDREELERLLRRSNE LIKRSRELNEESKKLIEKLERLAT SEQ ID NO:255
DHD150	Heterodimer	b	DNEEIIKEARRVVEEYKAVDRLEELVRAENAKHASEKELKDIVREILRI SKELNKVSERLIELWERSQERAR SEQ ID NO:256
DHD151	Heterodimer	a	PKEDIDRVSRRELVRVHKELLEVLRRKSTEIVEAVARNEKDERTIEEVLEEQE RAVRKLEEVSKKHKEAVKRLK SEQ ID NO:257
DHD151	Heterodimer	b	ELERLSEEIQKLSDRLEELIRRHSKVLEEIVRLKHKNDEREVRRLLKLL RDLTRRYEEVLRKVEEIVKRQEDSR SEQ ID NO:258
DHD152	Heterodimer	a	PEEDILRLLRKLVEVDKELLEVVRESTEVVRLVARNEKDVETVERVLRKQE EVVRKYERVSRELEAVRRLK SEQ ID NO:259
DHD152	Heterodimer	b	ELKDLVEEIVKLSKENLKLWEDHSRVLEEIVRLKHKNDEREVRRLLKLL EDLTRRAEETSRRIEIVKEAEDRAR SEQ ID NO:260
DHD153	Heterodimer	a	DEERELREVLRRKHHRVREWTQVVEELKRVVELLRGETSEEDLRLVLKKL LEMCKRILEVNRVLRVLEKRLT SEQ ID NO:261
DHD153	Heterodimer	b	SLEETIEELVELVRRSVEIAKESDEVARRIVESDKKKELIDTLRDLHREW QEVTKRAEELVREAEKEVR SEQ ID NO:262
DHD154	Heterodimer	a	TAEELLEHVHKSDRVTKHEHLRVSEELKVVVEVLRGEVSSVLRVLRKLE ELTDKLRRTVEEQRRVVEKLN SEQ ID NO:263
DHD154	Heterodimer	b	DLEDLRLRLRVLDEQRRLVEELERVSRLEKAVRDNEDERELARLSREHS DIQDKHDKLAREILEVLKRLLELTE SEQ ID NO:264
DHD155	Heterodimer	a	PEDDVVRIKEDLESNREVLREQEIHRIELVTRGEVSEEAIDRVLRKQE DLLKKQKESTDKARKVVEERR SEQ ID NO:265
DHD155	Heterodimer	b	DEVRLITWLKLSSESTRLLKELVELTRLLRNNVPNVEEILREHERISREL ERLSRRLKDLADKLERTRR SEQ ID NO:266
DHD156	Heterodimer	a	DEDEVVKVHEEHVKSHEEIHRSHEEVRAAEEDKRDRELRTLMEEHRKLL EENEKSIEEVKKIHERVKR SEQ ID NO:267
DHD156	Heterodimer	b	KKEELIDI SKEVLDDDEINKISKEILELIKLLRLKEGREDKDKAREVK RRIRELHRRIQELNKRRLRELHQRVQETKR SEQ ID NO:268
DHD157	Heterodimer	a	PEEDIARRVEDLLRSEELIKESKILKESKRLLDRNDSDKRVLETNLRLLI DKHTKLLERNLELLEELKLAEDVAK SEQ ID NO:269
DHD157	Heterodimer	b	RFKDLRSREYIEVVKRLELSREALEVLRKIDTDKTDKRIKELIDRLRKL IEEYKRIIDRLRKLKDLLEEHR SEQ ID NO:270
DHD158	Heterodimer	a	DEEELVKILKELQRLSEESLEINKRLEILRLLRRGEVPEEVEKKLREIK KEQEKLDREHEKIKKRIEETK SEQ ID NO:271
DHD158	Heterodimer	b	SLKEKILEIERNMKLVLSNRVETVARIKGEKDDEETLERLLREWDKI TRDYEEIIESRKLKVELEEEAK SEQ ID NO:272
DHD159	Heterodimer	a	SKTEILRKALEIHKQIDIVRKLIELSEEVKLVESKEKNLEKLRIDEE TDRLLERLDELHKLRELAERLK SEQ ID NO:273
DHD159	Heterodimer	b	SDDEARKQLEEMKRRLEVEKKSkrVEERVRELERLVRENREDEDRVLKTL EDLLRENEKLVRTIERHVREQRELSKEVK SEQ ID NO:274

DHD160	Heterodimer	a	SEEELEKKADELRLKLSSEWRKLQEEDKRLSEMVEKGELDLQEVDEHSRLVLERATEVHRTVVDKVIIEILRTTN SEQ ID NO:275
DHD160	Heterodimer	b	SEKERHRESQETQEEIRRTHEEIRKLEELRRAKAGELPEETLDRLRIMERLKELSERLDDLVRKLRDDHRREQ SEQ ID NO:276
DHD161	Heterodimer	a	SEKEILEELKRIKRVKDISDRLEELDKRTEETARREPTKELVDELVKIHRDWLRLHEEILKLVDDALKKVEDATK SEQ ID NO:277
DHD161	Heterodimer	b	DLRELELQREASRLHRELVKLLTELVKKLELIAKGEDIREEDLKRIKERLEEIKKRSKRIKEESDEIDKTK SEQ ID NO:278
DHD162	Heterodimer	a	SERELQRELNKIVRRILEIHREVSSELHQRAVKLIRENDNSEEELEISRRIEELSKLEKLVREHDEIVKTIE SEQ ID NO:279
DHD162	Heterodimer	b	SEREKLDNRNDEELKEINKRVEEIKERSDRITEAIEKNERSEEEIRRSREQNEALQRLLELHKLVKHLHRELEDTR SEQ ID NO:280
DHD163	Heterodimer	a	DKEDVIRVHDEQHKLIIEQLELTRRIAELVREIAKNTASEEEIKEMLKEIKRLDDRSREIQDRLQKLEEIIRKTK SEQ ID NO:281
DHD163	Heterodimer	b	TEEEIVELNKDIQRKSKEHIDLQNELVKKIERAIRENNITEELLEELERLLRESEKIVEEIRRIDTKIRKDAK SEQ ID NO:282
DHD164	Heterodimer	a	SEKEILERLLRSKEQNEISEEIHRLTERLVELKRRKDDDERLKRILDRQKRLVERAREISKEYEDLLRKE SEQ ID NO:283
DHD164	Heterodimer	b	SMEELLRNARLSRQQLKIIDEHLELSTKLTRGEAGDETLEEIERRSREMLEEQRRVDEESKRIREKLK SEQ ID NO:284
DHD165	Heterodimer	a	SEEEIRDIVEKLLRTHHEVLKEIKKLLDDSERVRRRELDKDLDRIQEQRDIQENKEKAKRFDELVKELKKA SEQ ID NO:285
DHD165	Heterodimer	b	SEEHRRTMKEKVEVRDIKRRSEEVKKVKANTLSEEDLVRLRLVEDHKRLQDLSQEIIEERDEKATK SEQ ID NO:286
DHD166	Heterodimer	a	DEDELAKEIEDVQRNKESQEEHDKSVKKLEAAERGEIDEDSLLRVLEEDIKVLEKDIEVLERSIEVIEKAE SEQ ID NO:287
DHD166	Heterodimer	b	SEKELIRRLLEQQRQHLRLSERLIELSRRLVEVVRKGDNRDLLRELKLS EEHKKHSDDEKVEIREK SEQ ID NO:288
DHS 17	Heterodimer	a	DRKDLLKRNKIKLLDRHLKILDTILKLEKLSSELLKSSSEEVVKEYKKILD EIRKLLSEESKEIHKESKEILERES SEQ ID NO:289
DHD17	Heterodimer	b	DEEKLIERSKRLQEESQLEKFEQILRELTELLEKPDSEELARKIKKLHD ELRKIKRNQELIREHEEILRKRD SEQ ID NO:290

In one aspect, the monomer A polypeptide comprises a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an odd-
5 numbered SEQ ID NO selected from the group consisting of selected from the group SEQ ID NOS: 1-290; wherein GlySer at amino acids 1 and 2 of SEQ ID NO: 1, 55, 81, 83, 101, 105, 115, 117, 119, 121, 123, 125, 127, 131, 133, 135, 137, 139, 141, 143, 145, 147, 149, 151, 153, 155, 157, 159, 161, 163, 165, 167, 169, 171, 173, 175, 177, 179, 181, 183, 185, 187, 189, 191, or 193 are optional, e.g., GlySer at amino acids 1 and 2 of
10 SEQ ID NO: 1, 55, 81, 83, 101, 105, 115, 117, 119, 121, 123, 125, 127, 131, 133, 135, 137, 139, 141, 143, 145, 147, 149, 151, 153, 155, 157, 159, 161, 163, 165, 167, 169, 171, 173, 175, 177, 179, 181, 183, 185, 187, 189, 191, or 193 are not present, and

wherein the odd-numbered SEQ ID NO ("chain a") is the binding partner of the SEQ ID NO. ("chain b") in Tables 1A.

In another aspect, the monomer B polypeptide comprises a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an even-numbered SEQ ID NO selected from the group consisting of selected from the group SEQ ID NOS: 1-290, wherein GlySer at amino acids 1 and 2 of SEQ ID NO: 6, 8, 14, 16, 26, 30, 32, 34, 36, 38, 40, 42, 46, 48, 54, 56, 58, 60, 62, 66, 68, 70, 72, 74, 76, 78, 80, 82, 84, 86, 88, 90, 92, 94, 96, 98, 100, 102, 104, 106, 108, 110, 112, 114, 116, 118, 120, 122, 124, 128, 130, 132, 134, 136, 138, 140, 142, 144, 146, 148, 150, 152, 154, 156, 158, 160, 162, 164, 166, 168, 170, 172, 176, 178, 180, 182, 184, 186, 188, 190, 192, or 194 are optional, e.g., GlySer at amino acids 1 and 2 of SEQ ID NO: 6, 8, 14, 16, 26, 30, 32, 34, 36, 38, 40, 42, 46, 48, 54, 56, 58, 60, 62, 66, 68, 70, 72, 74, 76, 78, 80, 82, 84, 86, 88, 90, 92, 94, 96, 98, 100, 102, 104, 106, 108, 110, 112, 114, 116, 118, 120, 122, 124, 128, 130, 132, 134, 136, 138, 140, 142, 144, 146, 148, 150, 152, 154, 156, 158, 160, 162, 164, 166, 168, 170, 172, 176, 178, 180, 182, 184, 186, 188, 190, 192, or 194 are not present, wherein the even-numbered SEQ ID NO ("chain b") is the binding partner of the SEQ ID NO. ("chain a") in Table 1A.

In another embodiment of any of the above embodiments,

(i) monomer A comprises a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an odd-numbered SEQ ID NO selected from the group consisting of selected from the group SEQ ID NOS: 1-290, 331, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494; and

(ii) monomer B comprises a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an even-numbered SEQ ID NO selected from the group consisting of selected from the group SEQ ID NOS: 1-290, 331, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484,

486, 488, 490, 493, and 494, wherein the even-numbered SEQ ID NO is the binding partner of the odd-numbered SEQ ID NO. in step (i).

The amino acid sequences of SEQ ID NOS: 1-290, 331, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 5 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494 are provided in Table 1B. The “binding partners” have similar design names as shown in Table 1B. For example, SEQ ID NO:1 (DHD9 A) and SEQ ID NO:2 (DHD9 B) are binding partners, and For example, SEQ ID NO:331 (DHD9 A) and SEQ ID NO:2 (DHD9 B) are binding partners, so that if monomer A comprises the polypeptide having 10 at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of SEQ ID NO:1 or SEQ ID NO:331, then monomer B comprises the polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of SEQ ID 15 NO:2. Similarly, SEQ ID NOS:3-4 are binding partners, SEQ ID NO:5-6 and 5-332 are binding partners, etc. Those of skill in the art will clearly understand what is meant by binding partner based on the teachings herein.

Table 1B

Design name	Oligomerization State	Chain	Design sequence
DHD9	Heterodimer	a	GSPKEEARELIRKQKELIKEQKKLIKEAKQKSDSRDAERIWKRSREINRES KKINKRIKELIKS SEQ ID NO:1 PKEEARELIRKQKELIKEQKKLIKEAKQKSDSRDAERIWKRSREINRESKK INKRIKELIKS SEQ ID NO:331
DHD9	Heterodimer	b	PKKEAEELAESEELHDRSEKLHERAEQSSNSEARKILEDIERISERIEE ISDRIERLLRS SEQ ID NO:2
DHD13_X AAA	Heterodimer	a	GTKEDILERQRKIIERAQEIHRQQEILEELERIIRKPGSSEEAMKRMLKL LEESLRLKELLELEESAQLLYEQR SEQ ID NO:3
DHD13_X AAA	Heterodimer	b	GTEKRLLLEEAREHREQKEIKKAQELHRRLEEVQRSGSSEEAKKEAKKI LEEIRELKRSLELLREILYLSQEQQKSLVPR SEQ ID NO:4
DHD13_X AXA	Heterodimer	a	TKEDILERQRKIIERAQEIHRQQEILEELERIIRKPGSSEEAMKRMLKLL EESLRLKELLELEESAQLLYEQR SEQ ID NO:5
DHD13_X AXA	Heterodimer	b	GSTEKRLLLEEAREHREAKEIKKAQELHRRLEEVQRSGSSEEAKKEAKK ILEEIRELKRLLELLREILYLSQEQQ SEQ ID NO:6 TEKRLLLEEAREHREAKEIKKAQELHRRLEEVQRSGSSEEAKKEAKKIL EEIRELKRLLELLREILYLSQEQQ SEQ ID NO:332
DHD13_X AAX	Heterodimer	a	TKEDILERARKIIERAQEIHRQQEILEELERIIRKPGSSEEAMKRMLKLL EESLRLKELLELESELAQLLYEQR SEQ ID NO:7
DHD13_X AAX	Heterodimer	b	GSTEKRLLEEAREIREQKEIKKAQELHRRLEEVQRSGSSEEAKKEAKK ILEEIRELKRSLELLREILYLLQEQQ SEQ ID NO:8 TEKRLLEEAREIREQKEIKKAQELHRRLEEVQRSGSSEEAKKEAKKIL EEIRELKRSLELLREILYLLQEQQ SEQ ID NO:334
DHD13_2	Heterodimer	a	TKEDILERQRKIIERAQEIHRQQEILEELEYIIR SEQ ID NO:9

:341			
DHD13_2 :341	Heterodimer	b	MSEAMKRMKLKLEESLRLKELLESEESAQLLYEQRKANNGETEKRLLEEAEARAHREQKEIIKKAQELHRRLEEVQRSGSSEAEAKKEAKKILEEIRELSKRSLELLREILYLSQEQQ SEQ ID NO:10
DHD13_A AAA	Heterodimer	a	MTKEDILERQRKIIERAQEIHRQEQEILKEQEKIIRKPGSSEAMKRSLKLEESLRLKELLESEESAQLLYEQR SEQ ID NO:11
DHD13_A AAA	Heterodimer	b	GTEKRLLEEAERAHREQKEIIKKAQELHKELTQIQQSGSSEAEAKKRALKSQEIRELSKRSLELLREILYLSQEQQ SEQ ID NO:12
DHD13_B AAA	Heterodimer	a	TKEDILERQRKIIERAQEIHRQEQEILKRSEEIIRKPGSSEAELETRELQEEESLRLKELLESEESAQLLYEQR SEQ ID NO:13
DHD13_B AAA	Heterodimer	b	GSTEKRLLEEAERAHREQKEIIKKAQELHRRTEEIIHQSGSSEAEAKDELRRIQEEIRELSKRSLELLREILYLSQEQQ SEQ ID NO:14 TEKRLLEEAERAHREQKEIIKKAQELHRRTEEIIHQSGSSEAEAKDELRRIQEEIRELSKRSLELLREILYLSQEQQ SEQ ID NO:336
DHD13_4 :123	Heterodimer	a	TTKRYLEEAERAHREQKEIIKKAQELHRRLEEVQR SEQ ID NO:15
DHD13_4 :123	Heterodimer	b	GSSEAEAKKEAKKILEEIRELSKRSLELLREILYLSQQVNDVDEKALERQRKIIRAQEIHRQEQEILEELERIIRKPGSSEAMKRMKLKLEESLRLKELLESEESAQLLYEAR SEQ ID NO:16 SEAEAKKEAKKILEEIRELSKRSLELLREILYLSQQVNDVDEKALERQRKIIRAQEIHRQEQEILEELERIIRKPGSSEAMKRMKLKLEESLRLKELLESEESAQLLYEAR SEQ ID NO:338
DHD13_1 :234	Heterodimer	a	EAMKRMKLKLEESLRLKELLESEESAQLLYEAR SEQ ID NO:17
DHD13_1 :234	Heterodimer	b	TTKRYLEEAERAHREQKEIIKKAQELHRRLEEVQRSGSSEAEAKKEAKKILEEIRELSKRSLELLREILYLSQQVNDVDEKALERQRKIIRAQEIHRQEQEILEELERIIRKPGS SEQ ID NO:18
DHD15	Heterodimer	a	TREELLRENIELAKEHIEIMREILELQKMEELLEKARGADEDVAKTIKELLRRLKEIIEIRNRIAKEHEIYARERS SEQ ID NO:19
DHD15	Heterodimer	b	GTERKLLERSRRLQEESKRLLEDMAEIMRRIKLLKARGADEKVLDELRIIERIRELDRSRKIHESSEIAYKEE SEQ ID NO:20
DHD20	Heterodimer	a	GDRQELIRNIELKEHIKILEEISQLIEELSDKSSSEEVVKRYKKILEERYKQLLRKSQEIHKESSEIAKES SEQ ID NO:21
DHD20	Heterodimer	b	GDEQKLIERSQRMQKESLELLKEIKILDITIEKLLDKPDSSELLDTIKKLHDTLKKIHDRNKKLKEHEEILRQRSGSLVPR SEQ ID NO:22
DHD21	Heterodimer	a	DKEEYKRLLEDEIKELKESKEVLKDSKRVLEDIKRVPDDDLVKLLEKHVRLLEEYKLLLEQLIREAEKSSK SEQ ID NO:23
DHD21	Heterodimer	b	QGSSEAEELKKEIKESEKKIRDSLKIKIIEIKSRKEGVDDKQLDLIRKVVESHRLDLRLHRDLRLREETS SEQ ID NO:24
DHD25	Heterodimer	a	DIDESIKEVEKLEVEQSLQKLDLSDKLLKLEKVNQDPDSDSVRKIVKRHVEILKRHEEVLKRLIEVVKEHTKTVK SEQ ID NO:25
DHD25	Heterodimer	b	GSDREEVHKEIVKLIREIIEIKHKKILKIEHEKIKNGEIDPSEILKLSEIIEKLTDTIIEIKIIEDEQLTRDLRR SEQ ID NO:26 DREEVHKEIVKLIREIIEIKHKKILKIEHEKIKNGEIDPSEILKLSEIIEKLTDTIIEIKIIEDEQLTRDLRR SEQ ID NO: 340
DHD27	Heterodimer	a	DRKEIVKRHQKVVLELKESSKLLRESSKLLQRLLDKTDGDNLQKAVDDQDKAIKQETAIRKSQEASKKLD SEQ ID NO:27
DHD27	Heterodimer	b	DNSEIIEKVAKTSREVAEYSERVAKENDKVVKTLEEGKIDSEELRLLEESIKIFDTALKLHEEAYKLHQDLVRKVS SEQ ID NO:28
DHD30	Heterodimer	a	DESEASVAIESVQILVESVKLEESVRILLDAVKKNGVEDLLRVAQRWEKLVDEWLKVVKRWLDNVRDIQR SEQ ID NO:29
DHD30	Heterodimer	b	GSDKAEVEKSVRKIEESIKKIRKSIKKAEDAVQLLKEGKIDAKDFLRIVREDELVVVEDVEIVKEDVENVREFSS SEQ ID NO:30 DKAEVEKSVRKIEESIKKIRKSIKKAEDAVQLLKEGKIDAKDFLRIVREDELVVVEDVEIVKEDVENVREFSS SEQ ID NO: 342
DHD33	Heterodimer	a	SDKEVSDKLLKASKKLLKVVSEELLEVVRLLKALKDDELIIKKIADLLRKIIDKDKKFIRTSEIIVKESR SEQ ID NO:31

DHD33	Heterodimer	b	GSDLKEVLKTVVEEAVKEI IKSSEELLQISRKILEISRQVDEHEYISAIRE YKALEKHIQILKKFIEILKELIRAVS SEQ ID NO:32 DLKEVLKTVVEEAVKEI IKSSEELLQISRKILEISRQVDEHEYISAIREYL KALEKHIQILKKFIEILKELIRAVS SEQ ID NO:344
DHD34_X AAXA	Heterodimer	a	SKEEIDKIVKHHKKKI EEHKKKVDLKKLVEEHDKRVSQDKDDKVKKLSEE VKKI I KRLEEVSKRLEEVSKLLKVISDKR SEQ ID NO:33
DHD34_X AAXA	Heterodimer	b	GSNDEELKKI LETLDRI LKKLDKILTRLIEVLKKS EDPNLDDKDYTELVKQ FIELIKKYEVEVKEYEEVVRQLIRLFS SEQ ID NO:34 NDEELKKI LETLDRI LKKLDKILTRLIEVLKKS EDPNLDDKDYTELVKQFI ELIKKYEVEVKEYEEVVRQLIRLFS SEQ ID NO:346
DHD34_X AXXA	Heterodimer	a	SKEEIDKIVKHHKKKI EELKKLVDELKKLVEEHDKRVSQDKDDKVKKLSEE VKKI I KRVEEVAKRLEEVSKLLKVISDKR SEQ ID NO:35
DHD34_X AXXA	Heterodimer	b	GSNDEELKKI LETLDRI LKKLEKILTRLIEVLKKS EDPNLDDKDYTELVKQ FIELIKKFEEVIKEYEEVVRQLIRLFS SEQ ID NO:36 NDEELKKI LETLDRI LKKLEKILTRLIEVLKKS EDPNLDDKDYTELVKQFI ELIKKFEEVIKEYEEVVRQLIRLFS SEQ ID NO:348
DHD34_X AAAA	Heterodimer	a	SKEEIDKIVKHHKKKI EEHKKKVDLKKLVEEHDKRVSQDKDDKVKKLSEE LKKI SKRLEEVSKRLEEVSKLLKVISDKR SEQ ID NO:37
DHD34_X AAAA	Heterodimer	b	GSNDEELKKI LETLDRI LKKLDKILTRLDEVLKKS EDPNLDDKDYTELVKQ YIELVKKYEVEVKEYEEVVRQLIRLFS SEQ ID NO:38 NDEELKKI LETLDRI LKKLDKILTRLDEVLKKS EDPNLDDKDYTELVKQYI ELVKKYEVEVKEYEEVVRQLIRLFS SEQ ID NO:418
DHD36	Heterodimer	a	DHSRKLKEILDRLRKHVKRLKEHLDELRLDVRQVPEDKLEHVVKLSDKIL QISERAVREFTKSVDKDS SEQ ID NO:39
DHD36	Heterodimer	b	GSDKKDELERILDEIRRLIERLDEILSRNLKLELLKHGVPNAKEVVKDYI RLLKEYLELVKEFLKLVKRHADLVS SEQ ID NO:40 DKKDELERILDEIRRLIERLDEILSRNLKLELLKHGVPNAKEVVKDYIRL LKEYLELVKEFLKLVKRHADLVS SEQ ID NO:350
DHD37_A BxB	Heterodimer	a	DSDEHLKKLKTFLENLRRHLDRDLKHIKQLRDI LSENPEDEVRKDVIDLSE RSVRIVKTVIKIFEDSVRKE SEQ ID NO:41
DHD37_A BxB	Heterodimer	b	GSDDKELDKLLDTLEKILQATKIIDANKLLEKLRRSERKDPKVVETYVE LLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO:42 DDKELDKLLDTLEKILQATKIIDANKLLEKLRRSERKDPKVVETYVELL KRHEKAVKELLEIAKTHAKKVE SEQ ID NO:352
DHD37_B BBB	Heterodimer	a	MDEEDHLKKLKTHLEKLERHLKLELHAKKLEDILKERPEDSAVKESIDEL RRSIELVRESIEIFRQSVVEEE SEQ ID NO:43
DHD37_B BBB	Heterodimer	b	GDVKELTKILDITLTKI LETATKVIKDATKLEEHKRSKDPDRLIETHKKL VEEHETLVRQHKELAEHLKRTR SEQ ID NO:44
DHD37_X BxB	Heterodimer	a	DSDEHLKKLKTFLENLRRHLDRDLKHLKELRDI LSENPEDEVRKDVIDELE RVIRIVKTVIKIFEDSVRKE SEQ ID NO:45
DHD37_X BxB	Heterodimer	b	GSDDKELDKLLDTLEKILQATKIIDLNKVEKLRRSERKDPKVIETVVE LLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO:46 DDKELDKLLDTLEKILQATKIIDLNKVEKLRRSERKDPKVIETVVELL KRHEKAVKELLEIAKTHAKKVE SEQ ID NO:354
DHD37_A XXB	Heterodimer	a	DSDEHLKKLKTFLENLRRLEDLLDKHIKQLRDI LSENPEDEVRKDVIDLSE RVVRTVKTIVIKIFEDSVRKE SEQ ID NO:47
DHD37_A XXB	Heterodimer	b	GSDDKELDKLLDTLEKILQATKVVDDANKLLEKLRRSERKDPKVVETYVE LLKRLEKLIKELLEIAKTHAKKVE SEQ ID NO:48 DDKELDKLLDTLEKILQATKVVDDANKLLEKLRRSERKDPKVVETYVELL KRLEKLIKELLEIAKTHAKKVE SEQ ID NO:356
DHD37_3 :124	Heterodimer	a	DSDEHLKKLKTFLENLRRHLDRDLKHIKQLRDI LSEN SEQ ID NO:49
DHD37_3 :124	Heterodimer	b	EDERVKDVIDLSESRVIRIVKTVIKIFEDSVRKEKTKPDSKAKELDKLLD TLEKILQATKIIDANKLLEKLRRSERKDPKVVETYVELLKRHEKAVKEL LEIAKTHAKKVE SEQ ID NO:50
DHD37_1 :234	Heterodimer	a	DSDEHLYKLTFLLENLRRHLDRDLKHIKQLRDI LSENPEDEVRKDAIDLSE RSVRIVKTVIKIFEDSVRKEKRPIDKRDDKELDKLLDTLEKILQATKII DDANKLLEYLRR SEQ ID NO:51

DHD37_1:234	Heterodimer	b	GDPKVVETYVELLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 52
DHD37_A XBB	Heterodimer	a	DSDEHLDRLDKHLKLLKTFLENLRRHIKQLRDLSENPEDEVRKDVLDLSK TVIKI FEDSVRKKERSVRIVE SEQ ID NO: 53
DHD37_A XBB	Heterodimer	b	GSDDKEATKI IDDLKLLDTLEKILQ TANKLLEKLRRSERKDPKVVETVYK AVKELLEIAKTHAELLKRHEKKVE SEQ ID NO: 54 DDKEATKI IDDLKLLDTLEKILQ TANKLLEKLRRSERKDPKVVETVYKAV KELLEIAKTHAELLKRHEKKVE SEQ ID NO: 358
DHD37_X BBA	Heterodimer	a	DSDEHIKQLRDLDRLDKHLKLLKTFLENLRRILSENPEDEVRKTVIKIFE DSVRKKERSVRIVKDVIDLSE SEQ ID NO: 55
DHD37_X BBA	Heterodimer	b	GSDDKEANKLLEKATKI IDDLKLLDTLEKILQ TLRRSERKDPKAVKELLE IAKTHAELLKRHEKVVETVYVKKVE SEQ ID NO: 56 DDKEANKLLEKATKI IDDLKLLDTLEKILQ TLRRSERKDPKAVKELLEIA KTHAELLKRHEKVVETVYVKKVE SEQ ID NO: 360
DHD39	Heterodimer	a	DHSRKLEELDRLRKHKVRLLEHLRELLSLVKENPEDKDLVEVLELSLAIL RRSLEAVEAFLKSVTKKDPDEDLRRKADEIRKEVEEIKKSLAEVEKEIYK LK SEQ ID NO: 57
DHD39	Heterodimer	b	GSSADDVLEDILKIRELIEILDQILSLNQLLKLRRHGVNPAKVVVEKYK EILEYLQVLVSLFLKIVKTHADAVSGKIDKKAEEEEIKKEEIKI KEKLRQAK DILKQLQEEIDKTR SEQ ID NO: 58 SADDVLEDILKIRELIEILDQILSLNQLLKLRRHGVNPAKVVVEKYKEI LELYLQVLVSLFLKIVKTHADAVSGKIDKKAEEEEIKKEEIKI KEKLRQAKDI LKQLQEEIDKTR SEQ ID NO: 362
DHD40	Heterodimer	a	DRDAHLYKLLTFLEQLVRHLDRLVKHITQLRDI VVKDPEDERAVDVIRQSV RSLIEIVITVLKIFVDSVSDAARSKEAEKIVRKIRKEIDEIRQKLEIDKEV KKTTS SEQ ID NO: 59
DHD40	Heterodimer	b	GSNDKVLDKILDILDRILRLATRVIDLANKLLQVKKKSTHKDPRIVETYKE LLKIHETAVRLLLELADLHRLKSKDEEANKRVETELDRI RKKVKDIEDKV RKLEDKVRKTAS SEQ ID NO: 60 NDKVLDKILDILDRILRLATRVIDLANKLLQVKKKSTHKDPRIVETYKELL KIHETAVRLLLELADLHRLKSKDEEANKRVETELDRI RKKVKDIEDKVRK LEDKVRKTAS SEQ ID NO: 364
DHD43	Heterodimer	a	NDSLKVELKLEKSV EELLRRVQKSVKAEQKRGLLSDELVDRHLKILNQLV KRHLELLQEVIKRSDKK SEQ ID NO: 61
DHD43	Heterodimer	b	GSDEAVKRVEKSLKILDEVIKKS LDILRELI ELQIRHAKDDES VIRASKS ALKDAIEALKKSLDEIKKALKR SADEG SEQ ID NO: 62 DEAVKRVEKSLKILDEVIKKS LDILRELI ELQIRHAKDDES VIRASKSAL KDAIEALKKSLDEIKKALKR SADEG SEQ ID NO: 366
DHD65	Heterodimer	a	SSEEVVKEVHEKVKLHKEILELLKIIKIHETAARDPDKDSIKKLSDEIK KIVKRIEDISDQAKRESSDAQRKQS SEQ ID NO: 63
DHD65	Heterodimer	b	DKEESKELKLLKKEIKLRSEELLEESKELLKLAKNGEIDSE LADADRKL NKKHEKLVQDIQDLLREHERQDR SEQ ID NO: 64
DHD70	Heterodimer	a	DEKKIDKIVKETEDLLQKSEKLLQQSKEAVKRIRSQVKENEIVDRLLRIS EELKISRRLVEISRRIASTLS SEQ ID NO: 65
DHD70	Heterodimer	b	GSSKEVIRLLKENVRLIKENLELLTRNLKLITDLVRGSGNSEEKIKTLKE LLKEYRELLKRYRKLVEDYKRLVDKHD SEQ ID NO: 66 SKEVIRLLKENVRLIKENLELLTRNLKLITDLVRGSGNSEEKIKTLKELL KEYRELLKRYRKLVEDYKRLVDKHD SEQ ID NO: 368
DHD88	Heterodimer	a	EIQELIKSRRRIEESKELIKES EEVLRRIKEILDRI RINGVDNQEDLLREI LKLLTKNLKIIQRNLKLLQDNAEILKRLVS SEQ ID NO: 67
DHD88	Heterodimer	b	GSYIEDVIKILDVSR ELIKLSRTIIKISEEINKQLQQGRDTKDLVKKYDE I IKKYTRIVQHYTEL IKELQKLLS SEQ ID NO: 68 YIEDVIKILDVSR ELIKLSRTIIKISEEINKQLQQGRDTKDLVKKYDEI I KKYTRIVQHYTEL IKELQKLLS SEQ ID NO: 370
DHD89	Heterodimer	a	SPTEEAIQLSQRVIELSKRVIELSKEILKLLKRVLDLLPDLKNEEKRLDD YDKELKEYDKELKKYKRLKDLAS SEQ ID NO: 69
DHD89	Heterodimer	b	GSEEEELKIQKELRIQSEILDKQKKILD TLRNNGAVTEEVRSILEKVER LSEEAKELSKEAKELTKEVSKLIS SEQ ID NO: 70

			EEEEILKIQKELLRIQSEI LDKQKKI LDTLRSNGAVTEEVRSILEKVERLS EEAKELSKEAKELTKEVSKLIS SEQ ID NO: 372
DHD90	Heterodimer	a	SPLKELNNQLRLRLRELVKVSKKIVDLSKTIIEVLKHTDLDPRLLDSLEKS QQELDKSQKELDKVVKELTKVNKKLQ SEQ ID NO: 71
DHD90	Heterodimer	b	GSPLEDLVRRKYDELVKTYEKLVEEFKAVDKYDKAVKKAPVSKAATDSL DLIRKVELELDRNLKLIKENAKLIKELK SEQ ID NO: 72 PLEDLVRKYDELVKTYEKLVEEFKAVDKYDKAVKKAPVSKAATDSL DLIRKVELELDRNLKLIKENAKLIKELK SEQ ID NO: 374
DHD91	Heterodimer	a	SPTRENEKVIKENEKVISDNERVLEEVVKVETATDRKEIQDAVDEVRKSV DKLRDSVRKLEESVRTLD SEQ ID NO: 73
DHD91	Heterodimer	b	GSPKIDISKRLLEISKRLVEISDRIVELLQRIADSKDPNKDLQKEVKD VLE EYKRLVREYREVVKYEYKVVV SEQ ID NO: 74 PIKDISKRLLEISKRLVEISDRIVELLQRIADSKDPNKDLQKEVKD VLE EYKRLVREYREVVKYEYKVVV SEQ ID NO: 376
DHD92	Heterodimer	a	DEDEHVQKLIKNADLLRKHAEELLKELVKLFQEIASQIPDDRVAKVT DQVVD RIDKILKQTEKLVRRTKQILDYSR SEQ ID NO: 75
DHD92	Heterodimer	b	GSNLEELVKLLKEVLEMHERLLRIHEDLVEAHKSNASDKESERK LKKSDKD IKESLKKIKSIIDQVRYIQS SEQ ID NO: 76 NLEELVKLLKEVLEMHERLLRIHEDLVEAHKSNASDKESERK LKKSDKDIK ESLLKKIKSIIDQVRYIQS SEQ ID NO: 378
DHD93	Heterodimer	a	PVEDITIEESRLRLEESLKLNRILKLLKLEDSLRLKLPSEEW RQRLDEFRRKLE DWKEELERWIEDVRYKKT SEQ ID NO: 77
DHD93	Heterodimer	b	GSDEYDYESREIIDEIRKLLDRSKKIVHRSQRLVERVKSTPL SEDQEDLIRR HEETINRHRELVKLEKVEDHERHIR SEQ ID NO: 78 DEDYDYESREIIDEIRKLLDRSKKIVHRSQRLVERVKSTPL SEDQEDLIRRHE ETINRHRELVKLEKVEDHERHIR SEQ ID NO: 380
DHD94	Heterodimer	a	PEEDSRVLERFVRSREVLKVLVEEFLRVSEELLREADRDRDRR LEEYERQVDELREEIRRYKEEVDKFDKEVKYK SEQ ID NO: 79
DHD94	Heterodimer	b	GSPEKDNRKLLDKVRKLVKESRRLVEELRKLVDQSTKNGLI DEKALRKQQ EVLRKVEEVLEKQERVLRELEEISYRVI SEQ ID NO: 80 PEKDNRKLLDKVRKLVKESRRLVEELRKLVDQSTKNGLI DEKALRKQQEV LRKVEEVLEKQERVLRELEEISYRVI SEQ ID NO: 382
DHD94_3 :214	Heterodimer	a	GSPERDENRKLDDKVRKLVKESRRLVEELRKLVDQSTKN SEQ ID NO: 81 PERDENRKLDDKVRKLVKESRRLVEELRKLVDQSTKN SEQ ID NO: 337
DHD94_3 :214	Heterodimer	b	GSDEKALRKQQEVLRLKVEEVLEKQERVLRELEEISYRVI TRGEDHKAEDSRRVLERFVRSREVLKVLVEEFLRVSEELL READRDRDRRLEEYERQVDELREEIRRYKEEVDKFDKEVKYK K SEQ ID NO: 82 DEKALRKQQEVLRLKVEEVLEKQERVLRELEEISYRVI TRGEDHKAEDSRRVLERFVRSREVLKVLVEEFLRVSEELL READRDRDRRLEEYERQVDELREEIRRYKEEVDKFDKEVKYK K SEQ ID NO: 384
DHD94_2 :143	Heterodimer	a	GSDRRLEEYERQVDELREEIRRYKEEVDKFDKEVKYK SEQ ID NO: 83 DRRLEEYERQVDELREEIRRYKEEVDKFDKEVKYK SEQ ID NO: 339
DHD94_2 :143	Heterodimer	b	GSPERDENRKLDDKVRKLVKESRRLVEELRKLVDQSTKNGLI DEKALRKQQ EVLRKVEEVLEKQERVLRELEEISYRVI TRGEDHKAEDSRRVLERFVRSREVLKVLVEEFLRVSEELL READR SEQ ID NO: 84 PERDENRKLDDKVRKLVKESRRLVEELRKLVDQSTKNGLI DEKALRKQQEV LRKVEEVLEKQERVLRELEEISYRVI TRGEDHKAEDSRRVLERFVRSREVLKVLVEEFLRVSEELL READR SEQ ID NO: 386
DHD95	Heterodimer	a	DLSEESKFFVEKVKLEKESRELEKQVKKIEEDSRVENDVQK EFLLELLKRLLDIQKKVVEVLREVVKVQQYVDS SEQ ID NO: 85
DHD95	Heterodimer	b	GSDSEYERQVRLRELDTVLKDSTVLEALRQVIRDSQDVVSKS DEESRRVIDDLEKVIQDSKKVLDLIDKSKSIK SEQ ID NO: 86 DSEYERQVRLRELDTVLKDSTVLEALRQVIRDSQDVVSKS DEESRRVIDDLEKVIQDSKKVLDLIDKSKSIK SEQ ID NO: 388

DHD96	Heterodimer	a	NEDELLKLLFTENLKLLEDENLKLLENLSLLRQANNITDKNRIREIVKQSKE IVKQSREILKQSKEIVERIKYIVS SEQ ID NO: 87
DHD96	Heterodimer	b	GSSLYELTQRYEKLQVQYEELVKDYRRLVKKLEKLRDNKPKDRLLKEIVD VIKKSVEIIDRSLKLLLEESIKILEETD SEQ ID NO: 88 SLYELTQRYEKLQVQYEELVKDYRRLVKKLEKLRDNKPKDRLLKEIVDVI KKSVEIIDRSLKLLLEESIKILEETD SEQ ID NO: 390
DHD97	Heterodimer	a	SQRSLEILKRILDVLKESLEILKESLSILRQLASRIKKNPNRKIEEILKES DKIKESDKVLKEIEEVIRYSS SEQ ID NO: 89
DHD97	Heterodimer	b	GSDIEYESKEILELIKELLKLSRELLKESRRALELVRKSRDSDIVEEVIQV HKKVLDIHKEVLKIVRKVVEVHRRVKS SEQ ID NO: 90 DIEYESKEILELIKELLKLSRELLKESRRALELVRKSRDSDIVEEVIQVHK KVLDIHKEVLKIVRKVVEVHRRVKS SEQ ID NO: 392
DHD98	Heterodimer	a	SKKDESTKLERLAEKIDEITKRIEELVKDVKRKSSSEGVDKQQQKIDEVFQ KLLDLQREILEILDRIKVVQYILD SEQ ID NO: 91
DHD98	Heterodimer	b	GSDLEYLNRRLLQLIKTLIDLNRHLLKLDKLLKLSREGDEEIKESKQ IQEQFKEIVERSKEIKQIKKIKRSQ SEQ ID NO: 92 DLEYLNRRLLQLIKTLIDLNRHLLKLDKLLKLSREGDEEIKESKQIQ EQFKEIVERSKEIKQIKKIKRSQ SEQ ID NO: 394
DHD99	Heterodimer	a	DFERSRRLEKVVEDLRRSSDRLREVIDELRKSADKEDDEDLRRARKEHR DLIEELKRALEKQEEIKHLQELVYRQL SEQ ID NO: 93
DHD99	Heterodimer	b	GSESEEVKVVVERIKKISRELEEVVKELDRVSKFDRHGETDEIVREHER IVEKLEEVKHTKIVEELAEIVYKQQ SEQ ID NO: 94 ESEEVKVVVERIKKISRELEEVVKELDRVSKFDRHGETDEIVREHERIV EKLEEVKHTKIVEELAEIVYKQQ SEQ ID NO: 396
DHD100	Heterodimer	a	SDDSVRVLDEIVKILDESVKLLKESLKLDDFLRTPKDDHLKEVVKESKK VVEQSKKVLDRIKKIYESK SEQ ID NO: 95
DHD100	Heterodimer	b	GSDLLYLSKELLKLVRELLKLSRELVLSRRLVNSTHKSPELVKYYDKLVK KYQDLKLLKLVADVADEYLRQRS SEQ ID NO: 96 DLYLSKELLKLVRELLKLSRELVLSRRLVNSTHKSPELVKYYDKLVKYY QDLKLLKLVADVADEYLRQRS SEQ ID NO: 398
DHD101	Heterodimer	a	DEKDYHRRLEIHELDLVRHEELIKRQKVVVEELERRGLDERLRRVVDVFR RSSERWEEVIERFRQVVDKLRKSVE SEQ ID NO: 97
DHD101	Heterodimer	b	GSDAYDLDRIVKEHRRLVVEEQRELVEELEKLVRRQEDHRVDDKESHEILER LERIIRSTRILTELEKLTDEFERRTR SEQ ID NO: 98 DAYDLDRIVKEHRRLVVEEQRELVEELEKLVRRQEDHRVDDKESHEILERLE RIIRSTRILTELEKLTDEFERRTR SEQ ID NO: 400
DHD102	Heterodimer	a	DERYRAREHTRVEEHTKRLRHILKRLREHEEKLRRELKPGDEITESVDRF KKIVDQFEESIKKFETVSEELRKSIS SEQ ID NO: 99
DHD102	Heterodimer	b	GSDRQRLDRDLKILEKLDDILKLLKDILETLSKDDVSDRRHKDLVEKFRE LVDTHHKLVERYRELQVQNR SEQ ID NO: 100 DRQRLDRDLKILEKLDDILKLLKDILETLSKDDVSDRRHKDLVEKFRELV DTHHKLVERYRELQVQNR SEQ ID NO: 402
DHD102_ 1:243	Heterodimer	a	GSDITESVDRFKKIVDQFEESIKKFETVSEELRKSIS SEQ ID NO: 101 DEITESVDRFKKIVDQFEESIKKFETVSEELRKSIS SEQ ID NO: 341
DHD102_ 1:243	Heterodimer	b	GSDPQRAADRDLKILEKLDDILKLLKDILETLSKDDVSDRRAKDLVEKFRE LVDTHHKLVERYRELQVQNR SEQ ID NO: 102 DPQRAADRDLKILEKLDDILKLLKDILETLSKDDVSDRRAKDLVEKFRELV DTHHKLVERYRELQVQNR SEQ ID NO: 404
DHD103	Heterodimer	a	NADDQLATSIKKLEDSIDQLIKIVRKFEESVKKLQKHGVDQHHVEILRKIV EIFRQHIEKLLKHEKLRYSSTSS SEQ ID NO: 103
DHD103	Heterodimer	b	GSDKEYLVTEHEKLVREHEKIVSEIEKLVKKHEAGVDESELEEI LKKVEKL LRKLDLDEILEQLTQLLRKTE SEQ ID NO: 104 DKEYLVTEHEKLVREHEKIVSEIEKLVKKHEAGVDESELEEI LKKVEKLLR KLDEILEQLTQLLRKTE SEQ ID NO: 406
DHD103_ 1:243	Heterodimer	a	GSDQHVVVEILRKIVEIFRQHIEKLLKHEKLRYSSTSS SEQ ID NO: 105

1:423			DQHVV EILRKIVEIFRQHIEK LKKHLEK LRYTSS SEQ ID NO: 343
DHD103_1:423	Heterodimer	b	GSDAEYLVTEHEKLVREHEKIVSEIEKLVKKHEKGVDESELEEILKKVEKLRKLEI LEQLTQLLRKA EKHI DKH SKAADQLATS I K K L E D S I D Q L I K I V R K F E E S V K K L Q K H SEQ ID NO:106 DAEYLVTEHEKLVREHEKIVSEIEKLVKKHEKGVDESELEEILKKVEKLLRKLDEI LEQLTQLLRKA EKHI DKH SKAADQLATS I K K L E D S I D Q L I K I V R K F E E S V K K L Q K H SEQ ID NO: 408
DHD104	Heterodimer	a	DEDDDIRRVLDESRRVLEHSRRVLRSEEVLEKASRKKEKDTEETIEKHLKRLREHAKKLEKHPRELD DFLYKEI SEQ ID NO:107
DHD104	Heterodimer	b	GSRDKYLLERLNDILK K L D E I V D K L S D I L K R L K D V R H D D R L Q E L V E R Y K E I V K E Y K R I V E E Y E K L V R E F E E Q Q R SEQ ID NO:108 RDKYLLERLNDILK K L D E I V D K L S D I L K R L K D V R H D D R L Q E L V E R Y K E I V K E Y K R I V E E Y E K L V R E F E E Q Q R SEQ ID NO: 410
DHD105	Heterodimer	a	DRDYEDKEFKKI KELEEDVQEELK K L Q E K I K R F S S E L E E P N E L L K E Q L K V N E E Q L E V N K K I L K I L R D Q L K Q N E SEQ ID NO:109
DHD105	Heterodimer	b	GSDAEYKVRRESVKRSKESVKHSE DVVDKLNKSVKLSSESGHSDAEKASRELVKLVREVV ELSREVIK LSEKVL R V I S SEQ ID NO:110 DAEYKVRRESVKRSKESVKHSE DVVDKLNKSVKLSSESGHSDAEKASRELVKLVREVV ELSREVIK LSEKVL R V I S SEQ ID NO: 412
DHD106	Heterodimer	a	DLQYKQEKLI RHFDVVREWDKLVRFKSKVLEKQKHESKDKELEEASRRVD ELIKRLREQLKRSKEILRRLKELSRKSS SEQ ID NO:111
DHD106	Heterodimer	b	GSDWEELLRRLEKVLQ EYEEIVKELI DLIERLIKVSEDKSKDASEYK KLVTELEK L I S K L E E I S K K L E E L V K E Y E Y K T E SEQ ID NO:112 DWEELLRRLEKVLQ EYEEIVKELI DLIERLIKVSEDKSKDASEYK KLVTELEK L I S K L E E I S K K L E E L V K E Y E Y K T E SEQ ID NO: 414
DHD107	Heterodimer	a	DAKDELEKSLQEIEESL K E L K K L L E E L D K S L R E L T S Q G R N K K L E E H I K K V Q K F I E L V K K Y I K A V Q D Y L K E V R Y D N S SEQ ID NO:113
DHD107	Heterodimer	b	GSDKERAARATEEMVKLT K K L L K A V E D L V R D V R R L L K E G L I S E K H A R I A E T I L E V F K K H A K I I K K H V D I V K Y D E S SEQ ID NO:114 DKERAARATEEMVKLT K K L L K A V E D L V R D V R R L L K E G L I S E K H A R I A E T I L E V F K K H A K I I K K H V D I V K Y D E S SEQ ID NO: 416
DHD108	Heterodimer	a	GSP L K E R L L E I Q R D L D R V L E E V V E R L L R I Q E R L D S V V E R K P P D V H E E Y K Y I V D E I R E I V E R V V R E Y E E I V K R I D E E V R SEQ ID NO:115 PLKERLLEIQ RDLDRVLEEVVERLLRIQERLDSVVERKPPDVHEEYKYIVD E I R E I V E R V V R E Y E E I V K R I D E E V R SEQ ID NO: 459
DHD108	Heterodimer	b	GSEEDERIRYDLDRIRKDVRRKLEEI RQ RVRELEK KLRDAGHRRDEKELLR ELIETS KDILRLVEELLK K I I D K S E D L L R K T E SEQ ID NO:116 EEDERIRYDLDRIRKDVRRKLEEI RQ RVRELEK KLRDAGHRRDEKELLR ELIETS KDILRLVEELLK K I I D K S E D L L R K T E SEQ ID NO: 420
DHD109	Heterodimer	a	GSDEEDYINENVEKDV R D I E D D V R R I N E R I R E L L E K I R T E E V L Q R V L E E H H E L V E R V L R K L V E I L R K H E E E N R SEQ ID NO:117 DEEDYINENVEKDV R D I E D D V R R I N E R I R E L L E K I R T E E V L Q R V L E E H H E L V E R V L R K L V E I L R K H E E E N R SEQ ID NO: 345
DHD109	Heterodimer	b	GSDEEYK E K L H K L L R E I E E L L K H Y R E L V R R L E E L V K R G E L D K D T A A H I L E R L S E L L E R I I R R V A H T L R R L S E E R R SEQ ID NO:118 DEEYK E K L H K L L R E I E E L L K H Y R E L V R R L E E L V K R G E L D K D T A A H I L E R L S E L L E R I I R R V A H T L R R L S E E R R SEQ ID NO: 422
DHD110	Heterodimer	a	GSDEDEISYDSKRRVEEIVRQAREKSEKSRKDI EDVAEVL R K G D V S E K E V V D E L V K V L E E Q V K V L R E A V E R L R E V L K K Q V D D V R SEQ ID NO:119 DEDEISYDSKRRVEEIVRQAREKSEKSRKDI EDVAEVL R K G D V S E K E V V D E L V K V L E E Q V K V L R E A V E R L R E V L K K Q V D D V R SEQ ID NO: 347
DHD110	Heterodimer	b	GSDIVELVDHLLKRS L K L L E E L A E L V R R L L E K S T E L L K R R T E E H K E E V V E E S E Y M V R E L E E R L R R V D E S E K L V R D A D K H I R SEQ ID NO:120 DIVELVDHLLKRS L K L L E E L A E L V R R L L E K S T E L L K R R T E E H K E E V V E E S E Y M V R E L E E R L R R V D E S E K L V R D A D K H I R SEQ ID NO: 424
DHD111	Heterodimer	a	GSKEKDIVKTLVDLLRENLETLERLIEEVVRLKENVDVRDEGRDDKDSERILRDIKRRIDEAAKESREI I E R I E K E V E Y R S R SEQ ID NO:121 KEKDIVKTLVDLLRENLETLERLIEEVVRLKENVDVRDEGRDDKDSERIL

			RDIKRRIDEAAKESREIIERIEKEVEYRSR SEQ ID NO: 349
DHD111	Heterodimer	b	GSPEVDLRRIVREILKASEELLRLRLKLIIDEALKLSERKRDSQEYREVVD RVKKELERLLDEYRKLVEELKEKLRDTR SEQ ID NO:122 PEVDVLRRIVREILKASEELLRLRLKLIIDEALKLSERKRDSQEYREVVD KKELEERLLDEYRKLVEELKEKLRDTR SEQ ID NO: 426
DHD112	Heterodimer	a	GSDKRYESEKLRRLDEAVEKVVREVERVERESDRVLEEVRRRESKEVVD KVIEDNDKALEDVLRVDEVAKVVRDVVRENTR SEQ ID NO:123 DKRYESEKLRRLDEAVEKVVREVERVERESDRVLEEVRRRESKEVVDKV IEDNDKALEDVLRVDEVAKVVRDVVRENTR SEQ ID NO: 351
DHD112	Heterodimer	b	GSPREYHSDILRKVDEILERIRRHADRVKKKSERLKRENVVNEHSKDVK RVIRELLELVKELLRLAKKHSDDQQE SEQ ID NO:124 PREYHSDILRKVDEILERIRRHADRVKKKSERLKRENVVNEHSKDVKRV IRELLELVKELLRLAKKHSDDQQE SEQ ID NO: 428
DHD113	Heterodimer	a	GSDEDEILYHSERLLQKLLKELDDLKEKSRELLEELKKEDPDRRIERIIR LHDEVKLDLDEVLNILEVHREVLERLR SEQ ID NO:125 DEDEILYHSERLLQKLLKELDDLKEKSRELLEELKKEDPDRRIERIIRLH DEVKLDLDEVLNILEVHREVLERLR SEQ ID NO: 353
DHD113	Heterodimer	b	DKLDRLKKIHEEALRRAEELIKRLLDIHRRALDLARRGELDDYLLKESERE LREIIRRAEELKESRDRLEEISR SEQ ID NO:126
DHD114	Heterodimer	a	GSPKEELIRRVLEEVKRLNEKLEIIRRAELVKRANDELPETEKLEIREDR ELEKLLKEIEDELRRIDKELDDALYEIED SEQ ID NO:127 PKEELIRRVLEEVKRLNEKLEIIRRAELVKRANDELPETEKLEIREDREL EKLLKEIEDELRRIDKELDDALYEIED SEQ ID NO: 355
DHD114	Heterodimer	b	GSPKLDKRELLERNLEKLEIREEVVKILRTNLERVREDIRDEDVLQEYE RLIRKAEEDLRRVLKEYDDLKLVYELR SEQ ID NO:128 PKLDKRELLERNLEKLEIREEVVKILRTNLERVREDIRDEDVLQEYERL IRKAEEDLRRVLKEYDDLKLVYELR SEQ ID NO: 430
DHD115	Heterodimer	a	GSKEDSVKRAEEIVRTLKLLLEDLSLREASRLRDIKNGEDEHNLRRRISEK LEELSKRITETIERLLRELQYTSR SEQ ID NO:129 KEDESVKRAEEIVRTLKLLLEDLSLREASRLRDIKNGEDEHNLRRRISEKLE ELSKRITETIERLLRELQYTSR SEQ ID NO: 357
DHD115	Heterodimer	b	GSPNQELLDVRVKILEDLRLNEELVRLNKEKLLKRALEMRRKNRDSSEVLE RLAEYRKRLEEYRRELEKLELEETIYRYKR SEQ ID NO:130 PNQELLDVRVKILEDLRLNEELVRLNKEKLLKRALEMRRKNRDSSEVLERL AEYRKRLEEYRRELEKLELEETIYRYKR SEQ ID NO: 432
DHD116	Heterodimer	a	GSDEEEAQHEVEKVLDDIRRLSEHLQKRLEEVLEEVYELRREGSDRTEVV ELLKEVIREIVRVNREALERLLRVVEAVKRNE SEQ ID NO:131 DESEEAQHEVEKVLDDIRRLSEHLQKRLEEVLEEVYELRREGSDRTEVV LKEVIREIVRVNREALERLLRVVEAVKRNE SEQ ID NO: 359
DHD116	Heterodimer	b	GSDEEELVETVKRIQKEILDRLTELAKLLVEIQREIKKLDGEDDKELKR LSDELEEKVRQVVEIKRSLDELEETVEYVSR SEQ ID NO:132 DEEELVETVKRIQKEILDRLTELAKLLVEIQREIKKLDGEDDKELKRSL SDELEEKVRQVVEIKRSLDELEETVEYVSR SEQ ID NO: 434
DHD117	Heterodimer	a	GSDEEEVVRRAEELVKEHEELIERVIRTHEELVYKLEDQGADKKLVVVLK RVVEESERVAREIVKVSRELIRLLEEASR SEQ ID NO:133 DEEEVVRRAEELVKEHEELIERVIRTHEELVYKLEDQGADKKLVVVLKRV VEESERVAREIVKVSRELIRLLEEASR SEQ ID NO: 361
DHD117	Heterodimer	b	GSSKEELKELEDLQRRLEELKQLQERVVLEELIKRLRDRGRDDKHLK RLVKEVRRSSEVLRSIKEVSDRVRYQLR SEQ ID NO:134 SKEELKELEDLQRRLEELKQLQERVVLEELIKRLRDRGRDDKHLKRL VKEVRRSSEVLRSIKEVSDRVRYQLR SEQ ID NO: 436
DHD118	Heterodimer	a	GSDKEESEYLLRDLVRLLEKVKKEIEEVNREVEKLLKVKDGRDLDRREVL REILRLNRELAIEIKVVDRIHVVERSER SEQ ID NO:135 DKEESEYLLRDLVRLLEKVKKEIEEVNREVEKLLKVKDGRDLDRREVLRE ILRLNRELAIEIKVVDRIHVVERSER SEQ ID NO: 363
DHD118	Heterodimer	b	GSDLHEVVYETKELKRIEEVVEELRKKSEDIIRKAERGEISEDELKRLQE EIAEAKKLLDEIKRVLERHLEQTL SEQ ID NO:136 DLHEVVYETKELKRIEEVVEELRKKSEDIIRKAERGEISEDELKRLQEEI

			AREAKKLLDEIKRVLERHLEQTL SEQ ID NO: 438
DHD119	Heterodimer	a	GSPVEEIIKEVVKRVI EVQEKVLRII SHAVKRVVEVQKKYDPGSEESNRVV EEVKKTIEDAIRESDEVVDEVVKRIQYTVR SEQ ID NO:137 PVEEIIKEVVKRVI EVQEKVLRII SHAVKRVVEVQKKYDPGSEESNRVVEE VKKTIEDAIRESDEVVDEVVKRIQYTVR SEQ ID NO: 365
DHD119	Heterodimer	b	GSPEQEIADRILTEIRESQKELERLARKILKLLDESQEKAKRGRLLSEESD ELLERIKKELDELLERSKELLKKIEYELR SEQ ID NO:138 PEQEIADRILTEIRESQKELERLARKILKLLDESQEKAKRGRLLSEESDEL LERIKKELDELLERSKELLKKIEYELR SEQ ID NO: 440
DHD120	Heterodimer	a	GSDEKKEANRVLDEVLKTVRDLLETANEVLEKLYRLKRTDDQEKVVRTL EVLKEHLKLVVEEIVRILDKVLKEHLETEK SEQ ID NO:139 DEDKKEANRVLDEVLKTVRDLLETANEVLEKLYRLKRTDDQEKVVRTL LKEHLKLVVEEIVRILDKVLKEHLETEK SEQ ID NO: 367
DHD120	Heterodimer	b	GSPEDDVLRRLEEVSEKILRVAEDVARQLREVSEKITQGKVDKREWEEDIK RLKRELEELLREWKEEIERLTYELR SEQ ID NO:140 PEDDVLRRLEEVSEKILRVAEDVARQLREVSEKITQGKVDKREWEEDIKRL KRELEELLREWKEEIERLTYELR SEQ ID NO: 442
DHD121	Heterodimer	a	GSRRREVVKRIRELLKRNKELIDRIRELEENEYLDKDKDARDKDLRRSVEL LEELVRILEESVELAKEI IKLLREVVE SEQ ID NO:141 RREVVVKRIRELLKRNKELIDRIRELEENEYLDKDKDARDKDLRRSVELLE ELVRILEESVELAKEI IKLLREVVE SEQ ID NO: 369
DHD121	Heterodimer	b	GSDEKEDNRRLOHQKIERILEKNEDLQRKLEEILELLERGEADEEKIDRLRK AVEDYRRVVEEIKEDVKRHKYTVR SEQ ID NO:142 DEKEDNRRLOHQKIERILEKNEDLQRKLEEILELLERGEADEEKIDRLRKAV EDYRRVVEEIKEDVKRHKYTVR SEQ ID NO: 444
DHD122	Heterodimer	a	GSDEKEEAKKASEESVRTVERILEELLKASEESVELLRGEGADKDVVERSK EALKRVKELDEVVVKRSDEILKYIHN SEQ ID NO:143 DEKEEAKKASEESVRTVERILEELLKASEESVELLRGEGADKDVVERSK LKRKVELDEVVVKRSDEILKYIHN SEQ ID NO: 371
DHD122	Heterodimer	b	GSDEKKLINEVVETQKRLIKEAAKRLSEVVRHQTELI RELREKNVDDKDVE KLLKESLDLAEIIVRRIKELLDKESKLVVEYVSN SEQ ID NO:144 DEKKLINEVVETQKRLIKEAAKRLSEVVRHQTELI RELREKNVDDKDVEKL LKESLDLAEIIVRRIKELLDKESKLVVEYVSN SEQ ID NO: 446
DHD123	Heterodimer	a	GSPDMDEVKRVLDLIEIQEELREIKRVLEKLIKIQEDNGSEYESREVV EIVEIARKLVRSRRVVKKITETLQ SEQ ID NO:145 PDMDEVKRVLDLIEIQEELREIKRVLEKLIKIQEDNGSEYESREVVREI VEIARKLVRSRRVVKKITETLQ SEQ ID NO: 373
DHD123	Heterodimer	b	GSDERYATREIVERIERIAREILKRTTEEIVREVREVLSDVDQEEVVRRLA DLLRESVELVQHLVRRVEELLQESVERKK SEQ ID NO:146 DERYATREIVERIERIAREILKRTTEEIVREVREVLSDVDQEEVVRRLADL LRESVELVQHLVRRVEELLQESVERKK SEQ ID NO: 448
DHD124	Heterodimer	a	GSPEREALREVLEDLKRVTDRLELVERVLEELKKVTDHVDSERILRESRR VLKELKDIIEEILRESEKVLKLYTED SEQ ID NO:147 PEREALREVLEDLKRVTDRLELVERVLEELKKVTDHVDSERILRESRRVL KELKDIIEEILRESEKVLKLYTED SEQ ID NO: 375
DHD124	Heterodimer	b	GSPAREILEEVVKKHLEVVEDAARILEEIIREHEKAVREDRDKKELEEISR DLLRKAREALKKVKDISDDLSREIEYVAS SEQ ID NO:148 PAREILEEVVKKHLEVVEDAARILEEIIREHEKAVREDRDKKELEEISRD LRKAREALKKVKDISDDLSREIEYVAS SEQ ID NO: 450
DHD125	Heterodimer	a	GSPVEEAIKKVIDDLRDVQRKIRELVEELIRLLEEVQRDNDKRESEYVVER VEEILRRITETSREVVRAVEDLS SEQ ID NO:149 PVEEAIKKVIDDLRDVQRKIRELVEELIRLLEEVQRDNDKRESEYVVERVE EILRRITETSREVVRAVEDLS SEQ ID NO: 377
DHD125	Heterodimer	b	GSDSDEKAEYLLKEMERVVRESDEVVKKILRDLEEVLERLRRGEISEDVT EILKELAERHIRAIEELVRRRLRELLERHKKR SEQ ID NO:150 DSDEKAEYLLKEMERVVRESDEVVKKILRDLEEVLERLRRGEISEDVT LKELAERHIRAIEELVRRRLRELLERHKKR SEQ ID NO: 452
DHD126	Heterodimer	a	GSPVEEVLKELSEVNERVRDIAREI IERLSEVNEEVKETDDEDELKISKK

			VVDEVEDLLRKILEVSEEVVRRVEYHDR SEQ ID NO:151 PVBEVLKELSEVNERVRDIAREIIEERLSEVNNEEVKETDDEDELKKISKVV DEVEDLLRKILEVSEEVVRRVEYHDR SEQ ID NO: 379
DHD126	Heterodimer	b	GSPKEDILREVLRRHKEIVREIVRLVREAVETHLELVKRNDDRDAQDVIR KLEEDLERLVRHAQEVIEEIFYRLH SEQ ID NO:152 PKEDILREVLRRHKEIVREIVRLVREAVETHLELVKRNDDRDAQDVIRKL EEDLERLVRHAQEVIEEIFYRLH SEQ ID NO: 454
DHD127	Heterodimer	a	GSPRSYLLKELADLSQHLVRLLELRLVRESERVVEVLERGEVDEEELKRLED LHRELEKAVREVRETHREIRERSR SEQ ID NO:153 PRSYLLKELADLSQHLVRLLELRLVRESERVVEVLERGEVDEEELKRLEDLH RELEKAVREVRETHREIRERSR SEQ ID NO: 381
DHD127	Heterodimer	b	GSREYI IKDILDSQEHLRLIEELLETTQKELLEILKRRPDSVERVRELVR RSKEIADEIRRQSDRNVRLLLEVSK SEQ ID NO:154 DREYI IKDILDSQEHLRLIEELLETTQKELLEILKRRPDSVERVRELVRRS KEIADEIRRQSDRNVRLLLEVSK SEQ ID NO: 456
DHD128	Heterodimer	a	GSDEKDEIRHVIESVERLIEDIKRLLKTLRELAHDDSDKKTVKEVLDRVKE MIERHRRELEEHRKELEAEYEVV SEQ ID NO:155 DEKDEIRHVIESVERLIEDIKRLLKTLRELAHDDSDKKTVKEVLDRVKEMI ERHRRELEEHRKELEAEYEVV SEQ ID NO: 383
DHD128	Heterodimer	b	GSESEDRIKELLKRHI ELVERHEELLHEIKKLI DLEEKDDKDREEAVKRID DAIKESEEMLEESKEILEEIEYLN SEQ ID NO:156 ESEDRIKELLKRHI ELVERHEELLHEIKKLI DLEEKDDKDREEAVKRIDDA IKESEEMLEESKEILEEIEYLN SEQ ID NO: 458
DHD129	Heterodimer	a	GSSLEDSVRLNDEVVKVVERVRLNQEVRVRLIKHATDVEDEETVKYVLERV REVLDESREVLKRVHELLEESERLE SEQ ID NO:157 SLEDSVRLNDEVVKVVERVRLNQEVRVRLIKHATDVEDEETVKYVLERVRE VLDESREVLKRVHELLEESERLE SEQ ID NO: 385
DHD129	Heterodimer	b	GSHEKDIVYKVEDLVKSDRIAERAREIVKRSRDIMREIRKDKDNKKSDD LLKVTRDLQRVVDELEELSRELLRVAEESRK SEQ ID NO:158 HEKDIVYKVEDLVKSDRIAERAREIVKRSRDIMREIRKDKDNKKSDDLL KVTRDLQRVVDELEELSRELLRVAEESRK SEQ ID NO: 460
DHD130	Heterodimer	a	GSPDELVEVKKLIDELKKSVERLEESI REVKESIKKLRKGDIDAEENIKLLK ENIKIVRENIKI KEIIDVVQYVLR SEQ ID NO:159 PELDEVKKLIDELKKSVERLEESI REVKESIKKLRKGDIDAEENIKLLKEN IKIVRENIKI KEIIDVVQYVLR SEQ ID NO: 387
DHD130	Heterodimer	b	GSDEEEIEELLRELEKLLKKEEAELESKKLIDESEELLRRDRDLKKEHVR ASEEHVKLSEEHRLISREIVKILEKAVYSTR SEQ ID NO:160 DEEEIEELLRELEKLLKKEEAELESKKLIDESEELLRRDRDLKKEHVRAS EEHVKLSEEHRLISREIVKILEKAVYSTR SEQ ID NO: 462
DHD131	Heterodimer	a	GSDESDRIRKIVEESDEIVKESRKLAEARELI KESEDKRVSEERNERLLE ELLRLIDENAELLKRNLELLKEVLYRTR SEQ ID NO:161 DESDRIRKIVEESDEIVKESRKLAEARELI KESEDKRVSEERNERLLEEL LRLIDENAELLKRNLELLKEVLYRTR SEQ ID NO: 389
DHD131	Heterodimer	b	GSDEDELERLREYHRVREYKLEELRRLYEYKRGVSEESDRI LR EIKEILDKSERLWDLSEEVWRTLLYQAE SEQ ID NO:162 DEDELERLREYHRVREYKLEELRRLYEYKRGVSEESDRI LREI EIKEILDKSERLWDLSEEVWRTLLYQAE SEQ ID NO: 464
DHD132	Heterodimer	a	GSDKSDASRRAIRVLEHEFVRVSEEVLEVLKRSVESLKRDLVDDEKIKRTHDR IEEELRRWKRELEELIERLREWEYHQD SEQ ID NO:163 DKSDASRRAIRVLEHEFVRVSEEVLEVLKRSVESLKRDLVDDEKIKRTHDR IEEELRRWKRELEELIERLREWEYHQD SEQ ID NO: 391
DHD132	Heterodimer	b	GSDDEEDKRLLLEEVKRSRLTDERILEKLRHSLEQLLEDVDKDEDSRRVLR ELDEITKRSREVVKRLKLAYESK SEQ ID NO:164 DDEEDKRLLLEEVKRSRLTDERILEKLRHSLEQLLEDVDKDEDSRRVLR ELDEITKRSREVVKRLKLAYESK SEQ ID NO: 466
DHD133	Heterodimer	a	GSDKEYKLDRI LRRLELIKQLSRILEEIERLVDELEREFLDDKEVQDVIE RIVELIDHELELLKEYIKLLEEYIKTKT SEQ ID NO:165 DKEYKLDRI LRRLELIKQLSRILEEIERLVDELEREFLDDKEVQDVIERI

			VELIDEHLELLKEYIKLLEEYIKTTK SEQ ID NO: 393
DHD133	Heterodimer	b	GSPSKEYQEKSAERQKELLHEYEKLVRLHRELVEKLRRELDKKEEVLRRLLV EILERLKDHLHKKIEDAHRKNEEAHKENK SEQ ID NO:166 PSKEYQEKSAERQKELLHEYEKLVRLHRELVEKLRRELDKKEEVLRRLLVEI LERLKDHLHKKIEDAHRKNEEAHKENK SEQ ID NO: 468
DHD134	Heterodimer	a	GSRDRKISEELIKALEDHIRMLEELIRAIIEHIKLAERGVDEKELRESLEE LKKIVDELEKSLEELRKLAEYKYETR SEQ ID NO:167 RDRKISEELIKALEDHIRMLEELIRAIIEHIKLAERGVDEKELRESLEE KIVDELEKSLEELRKLAEYKYETR SEQ ID NO: 395
DHD134	Heterodimer	b	GSPKEESVEELKRVIDKHEEILRELKRVLEEHERVSHDEDENELRRSLERL KHILDRLHESLKLHELHLLKNEYTER SEQ ID NO:168 PKEESVEELKRVIDKHEEILRELKRVLEEHERVSHDEDENELRRSLERLKH ILDRLHESLKLHELHLLKNEYTER SEQ ID NO: 470
DHD135	Heterodimer	a	GSDHEYVVKIVERILRVMEKHAETVKKHLEIVERVVRGEPSEDLRRLKES LREIEESLRELKELDELDELSEKTR SEQ ID NO:169 DHEYVVKIVERILRVMEKHAETVKKHLEIVERVVRGEPSEDLRRLKESLR EIEESLRELKELDELDELSEKTR SEQ ID NO: 397
DHD135	Heterodimer	b	GSDEEYVTRSQRRLKRLLEEYIKVVEEHARLVNERDDKELKRSIDELDK LTKELELVKRYKELVDKTET SEQ ID NO:170 DEEYVTRSQRRLKRLLEEYIKVVEEHARLVNERDDKELKRSIDELDKLT KELELVKRYKELVDKTET SEQ ID NO: 472
DHD136	Heterodimer	a	GSDKEEIVKLQDEVIKTLERHLDILRKHIDLLEKLDHLSEELKERVDRSI KKLEESIKRLERIEELQELAEYSL SEQ ID NO:171 DKEEIVKLQDEVIKTLERHLDILRKHIDLLEKLDHLSEELKERVDRSIKK LLEESIKRLERIEELQELAEYSL SEQ ID NO: 399
DHD136	Heterodimer	b	GSREEELKESAEELERSVRELKKEADKYKEEVDRLHYRGKVDKDWVRVVEK LIKLVVEEHLELIREHLELLKEERR SEQ ID NO:172 REELKESAEELERSVRELKKEADKYKEEVDRLHYRGKVDKDWVRVVEKLI KLVVEEHLELIREHLELLKEERR SEQ ID NO: 474
DHD137	Heterodimer	a	GSDMEYELKKSAAEELRKSLEELKRILDELHKSLELRRHGDEEYVQTVVEE LRKELEEHAKKLEELHKELEERVAT SEQ ID NO:173 DMEYELKKSAAEELRKSLEELKRILDELHKSLELRRHGDEEYVQTVVEELR KELEEHAKKLEELHKELEERVAT SEQ ID NO: 401
DHD137	Heterodimer	b	PEYELKKSVDLKRVDRLVEEVEEFELSKERLREDRKHLELVEEMVRLI EKHLELIKEHLKADDDHVR SEQ ID NO:174
DHD138	Heterodimer	a	GSREKDESKELNDEYKLLLEEYERLLRRSEELVKRAKGRDEKELKRILEE NEDILRRTKEILERTKEISSEQKYRRR SEQ ID NO:175 REKDESKELNDEYKLLLEEYERLLRRSEELVKRAKGRDEKELKRILEENE DILRRTKEILERTKEISSEQKYRRR SEQ ID NO: 403
DHD138	Heterodimer	b	GSDKDERQERLNEESDKSNEESERSNRESEELNRRARGPNDEKELQEILDR HLELLERNQRLLDENKEILRESQYLND SEQ ID NO:176 DKDERQERLNEESDKSNEESERSNRESEELNRRARGPNDEKELQEILDRHL ELLEERNQRLLDENKEILRESQYLND SEQ ID NO: 476
DHD139	Heterodimer	a	GSENYILKEIKLKLRENKLLHDLRLLDENLEELEKHGAKDLDDYRRKI EEIRKKVEDYREKIEEIEKKVERDR SEQ ID NO:177 ENKYILKEIKLKLRENKLLHDLRLLDENLEELEKHGAKDLDDYRRKIEE IRKKVEDYREKIEEIEKKVERDR SEQ ID NO: 405
DHD139	Heterodimer	b	GSESEYQEEILELLKESIKLLREILRLLEESEELWRRENTKSERSEEIKE RAKAIKRSEEILERVKRLSDHSR SEQ ID NO:178 ESEYQEEILELLKESIKLLREILRLLEESEELWRRENTKSERSEEIKERA KAIKRSEEILERVKRLSDHSR SEQ ID NO: 478
DHD140	Heterodimer	a	GSDEEANYVSDKAVKIAEDVQELLKELLESEVVRGVEDEYDVRVLRK LQEVMEKEYEVLKEYEEVSRKHE SEQ ID NO:179 DEEANYVSDKAVKIAEDVQELLKELLESEVVRGVEDEYDVRVLRKLQ EVMKEYEVLKEYEEVSRKHE SEQ ID NO: 407
DHD140	Heterodimer	b	GSPEKYLIKQEEELRRHAEILEDLIRKVERQVDLRRKVDERDEDLKRELE RSLRELERLVRESSRLVEEIRELSKEIKR SEQ ID NO:180 PEKYLIKQEEELRRHAEILEDLIRKVERQVDLRRKVDERDEDLKRELEERS

			LRELERLVRESSRLVEEIRELSKEIKR SEQ ID NO: 480
DHD141	Heterodimer	a	GSDEEYELERISRESKELLERYKRLLEQYQELLKELRHVKDLDRAVKIIHE LMRVSKELVEISHRLELHERLVRRRK SEQ ID NO:181 DEEYELERISRESKELLERYKRLLEQYQELLKELRHVKDLDRAVKIIHELM RVSKELVEISHRLELHERLVRRRK SEQ ID NO: 409
DHD141	Heterodimer	b	GSEKEYIEKLSRKIEEDIRRSEERAKDSERLVRRLEELAKRKRDLDDVLR VAENLEILEDNLRILEEILKEQDKSNR SEQ ID NO:182 EKEYIEKLSRKIEEDIRRSEERAKDSERLVRRLEELAKRKRDLDDVLRVA EENLEILEDNLRILEEILKEQDKSNR SEQ ID NO: 482
DHD142	Heterodimer	a	GSPHEEVVELHERVMEISERAVELIQRIIDIIRRIREDDKDIEKLVKTIIRD LVREYEELHRELEEIDEEIYKKE SEQ ID NO:183 PHEEVVELHERVMEISERAVELIQRIIDIIRRIREDDKDIEKLVKTIIRD REYEELHRELEEIDEEIYKKE SEQ ID NO: 411
DHD142	Heterodimer	b	GSDHEDVVRLEDLVRKQEDARRVLEEVRLAEIIVEVIKKDEKDKRVTR LVEEIEKLVVEYKVKVDEMRSISDEIKYRSR SEQ ID NO:184 DHEDVVRLEDLVRKQEDARRVLEEVRLAEIIVEVIKKDEKDKRVTR LVEEIEKLVVEYKVKVDEMRSISDEIKYRSR SEQ ID NO: 484
DHD143	Heterodimer	a	GSRAREVVKRAKRIIEEWQKILEEWRRILEEWRRLLEDERVDDRDNERIIR ENERVIRENEKIIRDVIRLLEELLYERR SEQ ID NO:185 RAREVVKRAKRIIEEWQKILEEWRRILEEWRRLLEDERVDDRDNERIIR ERVIRENEKIIRDVIRLLEELLYERR SEQ ID NO: 413
DHD143	Heterodimer	b	GSREDEEELVEEIDRIRQMVVEEYELVKEYEELTEKYKQKGVDKKEESKIIIE KSERLLDLSQDAVRKVKIIRRIILYTNR SEQ ID NO:186 REDEEELVEEIDRIRQMVVEEYELVKEYEELTEKYKQKGVDKKEESKIIIE KSERLLDLSQDAVRKVKIIRRIILYTNR SEQ ID NO: 486
DHD144	Heterodimer	a	GSPKEEIVKLHDESAELHRRSVEVADEILKMHRSKDVDDERESRELSKEI ERLIREVEEVSKRIKRLSEEVEYLVR SEQ ID NO:187 PKEEIVKLHDESAELHRRSVEVADEILKMHRSKDVDDERESRELSKEI ERLIREVEEVSKRIKRLSEEVEYLVR SEQ ID NO: 415
DHD144	Heterodimer	b	GSPLEEILKIQRINKIQDDINKILHEILRMQEKLNRSDDKDEVEESLRRI RELKRIKDLKSKIEEDLSREVKYRRTT SEQ ID NO:188 PLEEILKIQRINKIQDDINKILHEILRMQEKLNRSDDKDEVEESLRRI RELKRIKDLKSKIEEDLSREVKYRRTT SEQ ID NO: 488
DHD145	Heterodimer	a	GSPDEHVYVREIYEVLRHAEVLEENREVIERLLEAKRGDKSEELVKE LKKSIDKLEKISRKLEEVKELEKVSEKLV SEQ ID NO:189 PEDEHVYVREIYEVLRHAEVLEENREVIERLLEAKRGDKSEELVKE LKKSIDKLEKISRKLEEVKELEKVSEKLV SEQ ID NO: 417
DHD145	Heterodimer	b	GSDEDETSYRIELREIVRASRELIRLSEELLEVARDDKDETLETILIR EYKELLDYRRLIEELTRLVEEYEERSR SEQ ID NO:190 DEDETSYRIELREIVRASRELIRLSEELLEVARDDKDETLETILIR EYKELLDYRRLIEELTRLVEEYEERSR SEQ ID NO: 490
DHD146	Heterodimer	a	GSTQEEINRIQHEVLRIQEEIDEILRDIVEKKAISRGELDHEVVKDVEDK VREALEKSEELLDKSRKVEYKSE SEQ ID NO:191 TQEEINRIQHEVLRIQEEIDEILRDIVEKKAISRGELDHEVVKDVEDK VREALEKSEELLDKSRKVEYKSE SEQ ID NO: 419
DHD146	Heterodimer	b	GSDEEELNRELEKSKRLVDINRDIIRTAQELIEMLKDSKGRVDEDTKRE LRDKLRKLEEKLERVREELRKYEELLRVQOR SEQ ID NO:192 DEEELNRELEKSKRLVDINRDIIRTAQELIEMLKDSKGRVDEDTKRE LRDKLRKLEEKLERVREELRKYEELLRVQOR SEQ ID NO: 492
DHD147	Heterodimer	a	GSDEKDRVYEILKEVQRLVKEYRDISKEIEDLVKHYEHITDDEAQEVSKEL IDKSLRASEIVRELIRLIKELDELE SEQ ID NO:193 DEKDRVYEILKEVQRLVKEYRDISKEIEDLVKHYEHITDDEAQEVSKEL IDKSLRASEIVRELIRLIKELDELE SEQ ID NO: 421
DHD147	Heterodimer	b	GSDEEDVLYHLRELEELKRVSDDYERLVREIKETSERKDRDTKENKDMLD ELVKAHQEKLLERLVRLLEELFERKR SEQ ID NO:194 DEEDVLYHLRELEELKRVSDDYERLVREIKETSERKDRDTKENKDMLD ELVKAHQEKLLERLVRLLEELFERKR SEQ ID NO: 494
DHD1	Heterodimer	a	PREQAIRISEEIRISKIIEILERTRSSTAREAMKWAKDSIRLAESKYL

			LDK SEQ ID NO:195
DHD1	Heterodimer	b	IEDDVKKIQDSTKKAQKETIEALERSTSSSTARKQMEEQEQIIRLQKEAMYL LKK SEQ ID NO:196
DHD2	Heterodimer	a	SREETAKLQEEVIKLRVIELQKEVIELQRRAKELTSSYTKELIQRRI EEIQREIEEIQKRIEIQEIIQRRT SEQ ID NO:197
DHD2	Heterodimer	b	SDEEIKRLSEEVIIQLSRRVIKMSREAIKLSREVQKLTPSYQKRIKEIADRS IELARESIEIAKRSEKIAEESQRRT SEQ ID NO:198
DHD3	Heterodimer	a	PAKDEALKMANESLELAKKSARLIQESSKEILERIEKIQRRIAELQDRIA YLIKK SEQ ID NO:199
DHD3	Heterodimer	b	PAKDEALRMI DESRELIKKSNELIQRSSSKEILERILEIQRKIAELQKRIQ YLLKS SEQ ID NO:200
DHD4	Heterodimer	a	TDEARYRSERIVKEAKRLLDEARRRSEKIVREAKQRSNSED AKRIMEENLR ESEEARRLREIIRRNLEESRETG SEQ ID NO:201
DHD4	Heterodimer	b	TREALEYQRKMAEEIEDLLREALRRQEEVREAKQRSLSSEEFKRIMERILE EQERVMLAKEALERILEEQKRTG SEQ ID NO:202
DHD5	Heterodimer	a	SERTKREAKRSQEEILREAKEAMRRAKESQDHRQNRDGSNSEDLERLSQEQ KRELEEVERLRLKELAREQKYKLEDS SEQ ID NO:203
DHD5	Heterodimer	b	SEDLKRIKLEITERELKLMQDLMEILKKITEDENNLDNNSSEDLKRSIEKA RRILDEALRKLEESARRAKYIQEDN SEQ ID NO:204
DHD6	Heterodimer	a	TEDEIRESLKLWDEVLQELREIARESNEVLERNRQKSRSDKLRREDIERYKK RMEEARKKLDQNLKYYKRM DENRS SEQ ID NO:205
DHD6	Heterodimer	b	TEELKESKFFAEDLARSARRALKESKRVLEEISQASRSKKLEEVRRYKE QVKRWQDEWDERAREYRKRMMKENRS SEQ ID NO:206
DHD7	Heterodimer	a	TKTEEIERLAREIKKLSEKVERLAQEIIEELSRVKEENSTDRELKEANREI ERAIREIEKANKRMEALRRMKYNG SEQ ID NO:207
DHD7	Heterodimer	b	TKTEEHERLAREISKLADHRKLAKIIEELARRIKEENLTDDELREAIRKI EDALRKNKEALKIMKEAERNRYNT SEQ ID NO:208
DHD8	Heterodimer	a	TKKEESRELAARESELEARESEKLARKSLELARRAESGSEEEKRRIIDENR KIIERNREIERNKEIIEYNKELIS SEQ ID NO:209
DHD8	Heterodimer	b	TKDEESLELNRESEELNRKSEELNRKSKELNDRAESSNSEEEKEILREHK EILREHLEILRRHKEILRRHKYLTSS SEQ ID NO:210
DHD16	Heterodimer	a	TRELLRENI ELAKEHIEIMREILELQKMEELLERQSSSEDI LEELRKIE RIRELLDRSRKIHESSEEIAYKEE SEQ ID NO:211
DHD16	Heterodimer	b	SEDIAREIKELLRRLKEIERNQRIAKEHEIYIARERKKLDPSNEKERKLE RSRRLQEE SKRLLDEMAEIMRRIKLLD SEQ ID NO:212
DHD18	Heterodimer	a	DRQKLI EENIKLLDKHIKILEEILRLLKDDIDLKKSSEEVLEELKKIHR RIDKLLDES KKHRSSEIVKKRS SEQ ID NO:213
DHD18	Heterodimer	b	DEQKLIETSQRLQEKSERLLEKFEQILREASDLYRKPDS EELLRRVEKLLR ELEKLI RENQDLARKHEKILRDQS SEQ ID NO:214
DHD19	Heterodimer	a	DRQELIRENIELLKKHIKIVKEIQKLIETFI ELLKKSSEELRRLKLIK RIEKLYRESQEIHKRSEEI AKKRQ SEQ ID NO:215
DHD19	Heterodimer	b	DEERLIDKSRELQKES EELLKELLKIFKRIEELLEKPDSEELIREIKKLE TLSEIHKRNEKLARTHEEILRQSS SEQ ID NO:216
DHD22	Heterodimer	a	STRDVQREIAKAFKMKADVQKLAEEIKRHVKVVEKKNKDNDEYRKIATEL LKKATESQKLLKELDRIRKSDS SEQ ID NO:217
DHD22	Heterodimer	b	DKDDRSTSLLRVEKLIDESDRIIDKFTTIELSRNGKIDDDQYKKEKLEI LELLKKYDKHVKEVEELLKRLNS SEQ ID NO:218
DHD23	Heterodimer	a	SKRKALEVSEVRVRISEKVVRLDES DLLKKS YDSDSKFAELIDRHEEKI KKWKKLIKEWLEIIQRHKS SEQ ID NO:219
DHD23	Heterodimer	b	SAEEFVKLSEAVKRSKEILDIVRKQVKLVKAGVDKHEITDSLRKSEKLEI EHKELIKTHRDLRREN SEQ ID NO:220
DHD24	Heterodimer	a	SSTEILKRFKRALRESEKIVKHSRRVLKIIREVLKQKPTQAVHDLVRIIET QVKALEEQLVKLRIVEALERQS SEQ ID NO:221
DHD24	Heterodimer	b	DKQKEIKDILEKTRRIAEESRKIAEKFDEIIRKSTEGKIDESLTKELEELV KEVIKLS EDDARTSDDLVRKES SEQ ID NO:222
DHD26	Heterodimer	a	DEDESIKLTRKSIETRKS LKIIKEVV ELIREVLKHIKDL DKEIFERIDKI LDKYKQVDTYDEILKEYEKKQR SEQ ID NO:223

DHD26	Heterodimer	b	SELDEQKELIKKQEKLIIEQQRLLSKIRRMFKERVVDQELLREIQVVKRS QEIVETSKKILDRSCKTTE SEQ ID NO:224
DHD28	Heterodimer	a	DQKEINTRIVEKLERIFKKSKEIVRQSERVISTIEKKTEDERELDLRRHV KIVREHLKLEELLKIIKEVQKQES SEQ ID NO:225
DHD28	Heterodimer	b	DTEELVKRLNELLKELSKLVKEFIKILETYRKDQTKDTSKISERVDRILKT YEDLQKYKEILEKIEKQLS SEQ ID NO:226
DHD29	Heterodimer	a	DYARLIDQAVEVTRKVVVEVNVTVARVNDKFAKHLGDEELRRVSEHLKEVSK DLQEVAKKSKDAARQVK SEQ ID NO:227
DHD29	Heterodimer	b	DVSKVAEEYLQISKTLVDISRTLLEISERLVRLVRTVADDRSEVKKAIEDS IEVLKTSSEVVRQIKRASDKLVKAIS SEQ ID NO:228
DHD31	Heterodimer	a	DAKEIQRRVVEIQTEVVKLQKKAVDIRKII EAFNNSNIDQSLLAAKEIV KEIDKLEKTESLLEESKLLKRSS SEQ ID NO:229
DHD31	Heterodimer	b	SAEEVVKLAKIFLELLRESIKLLKRSVDLLRKS SDPSLDKSEAEKVSREIE KVS DTS LKLSK KALD VVKRALKVAS SEQ ID NO:230
DHD32	Heterodimer	a	DEKDAARKARKVSEEAKEASKKIEKALEESKRILNTLQKQKDEQEVKVIKE HEDVLRQIEKIQKQVLEIQKEVAKLLES LD SEQ ID NO:231
DHD32	Heterodimer	b	SADDVARASEKVLRVARESAKAADKS LEVFKVVKRGDKEAFLQVVKINEE VVKINITVIRILIEVSKTAT SEQ ID NO:232
DHD38	Heterodimer	a	DEYKETLQQLREALASLREADKRITELVKEARKKPLSEAAKFAEAIVTH VKVVVEHVVELRHVEVLVEAKKNGVIDKSIDNALRIENVI RLLSNVIR VVDEVLQDL SEQ ID NO:233
DHD38	Heterodimer	b	DASDVIRRIHELFEVHRLIEAVHRAIEDVAKAAQKGLDESAREVILAEIS KELAKLSRRLAEISREIQKVVTPDDKKAVERLKEIKKIQDLDELDRDL RKLQDLLYKLS SEQ ID NO:234
DHD60	Heterodimer	a	SEDKAHHDIVRVLEELIKIHDELMKISEEILKATSDSTATDETKEELKRSS KEAQKKS DTLVKIVKELEKESRKAQS SEQ ID NO:235
DHD60	Heterodimer	b	DDEEKYRQIIREAQEISKTAKRILRDAQEISKRIRHQGVDRSEHQRLVDLL RELIEKHHKLLRRQEQEADTRND SEQ ID NO:236
DHD63	Heterodimer	a	DRDKARKASEKLEEVQRWKT VADKWKMMVDLVSNGKLSQEEVARVTEEL LKIQTTELAKLLEEHAKVLQESAS SEQ ID NO:237
DHD63	Heterodimer	b	SDESIKTQSELIKTSEELLKDVKRI DEELQKLRDPTLDESELKRVKEW SDRVKAKEISRKIQEIVKESKRSS SEQ ID NO:238
DHD66	Heterodimer	a	DKDEELRQVIKEYREMVKYRKVIREYEEVVKSSKTIDKSSLSISLRKMVE LSQRVIDVSDDEVAKVLSRKQS SEQ ID NO:239
DHD66	Heterodimer	b	TDEERLKKQTKELKEQTKQLEKQKDLLEKISNGEISKDEIQEIKESKKA KESQKALDSSRKALEEVS SEQ ID NO:240
DHD67	Heterodimer	a	DEKEVSKKIKVLKDIKQKQVIEVSRQLASVLRADDNVVKRALEEYK ILEELRELNKEIEKLTDKYRKVTS SEQ ID NO:241
DHD67	Heterodimer	b	DSDEQTKLEKLELTKRHRVEKLLKQTKESREVDNKLWKSVDKDKLSES EKELQKLS DQDKAKDALESSRRKND SEQ ID NO:242
DHD69	Heterodimer	a	DAEQKLLTKLRLRQQLQLIKESLKLIEKIQSSQENQDEIRKWEVT KKLRELIKTSEKLVRELEKSYKSS SEQ ID NO:243
DHD69	Heterodimer	b	SLRDVVRRYQELVRRYDELIKTLTEILKKYQKGAEDKDASTELVKAVRTS LKLSKELLKLNSELLKEDS SEQ ID NO:244
DHD71	Heterodimer	a	SKEELKRLDELKRS DTLKELSKLKEISERNPDDKSVHRTIIRIHRETV KNHKEIVRVEEIVSDKS SEQ ID NO:245
DHD71	Heterodimer	b	SKQDEHDLRLKIHDKLVKQHDPELLKLLTKLSRAGDSVTKKLEELLRKLQE VSKQLEESLKDADKVS KDIN SEQ ID NO:246
DHD72	Heterodimer	a	TVQSLLEQHVIVKRSIEILERHTQILQDIARSQGVSKLEDVERQVKEYR KEVKKLEEDLRQLSRNSK SEQ ID NO:247
DHD72	Heterodimer	b	SDSDRIEKLI RESTELLKEQQKLAKRSRELAETVESLPLTEEYKQQREHQ KKIEKLLKSEKHLLELKRVLKSEK SEQ ID NO:248
DHD73	Heterodimer	a	DSEKRIEDILRTDLELAKRDAELVKEHIKLVKRIDLSEELKQVEDVEKES KKLEDSSEKLVQKVRKRSS SEQ ID NO:249
DHD73	Heterodimer	b	DEERAKDLRKYLEEQTQYYRTVTEHLRNLKVVVEELERRGKPSSELQQL ERSQRIYKETTEIYDTSKKLIEELDKHHR SEQ ID NO:250
DHD148	Heterodimer	a	PLEDILKRRHLDKVRELVRLESEVNKLAKEVLDILKDKRVDEKELDKVLKEL EKVVEYERAVKESRDLRELRETRR SEQ ID NO:251

DHD148	Heterodimer	b	DKERLLEIHERIQKLLDRNLEIIERLLRLLREARDIKDDDKLKVIRLKE LSEESKLDIKIKELKSEKELT SEQ ID NO:252
DHD149	Heterodimer	a	PEDEVIRVIEELLRIAAAEVDEVHRRNVEVQEEASRVTDRELERLNRESEE LIKRSRELIEEQKLIERLERLAT SEQ ID NO:253
DHD149	Heterodimer	b	DLEELIKEYAEVRRHHKAVRDLERLVRELANAKHASEELKRIATEILRI VKELIRVQERLIKLSSEDSNEESR SEQ ID NO:254
DHD150	Heterodimer	a	PTDEVIEVLKELLRIHRENLRVNEEIVEVNERASRVTDREELERLLRRSNE LIKRSRELNEESKKLIEKLERLAT SEQ ID NO:255
DHD150	Heterodimer	b	DNEEIIKEARRVVEEYKAVDRLEELVRAENAKHASEKELKDIVREILRI SKELNKVSERLIELWERSQERAR SEQ ID NO:256
DHD151	Heterodimer	a	PKEDIDRVSRRELVRVHKELLEVLKSTEIVEAVARNEKDERTIEEVLEEQE RAVRKLEEVSKKHKEAVKRLK SEQ ID NO:257
DHD151	Heterodimer	b	ELERLSEEIQKLSDRLELIRRHSKVLEEIVRLKHKNDEREVRRLKLL RDLTRRYEEVLRKVEEIVKRQEDES SEQ ID NO:258
DHD152	Heterodimer	a	PEEDILRLLRKLVEVDKELLEVVRESTEVVRLVARNEKDVETVERVLRKQE EVVRKYERVSRELEAVRRLK SEQ ID NO:259
DHD152	Heterodimer	b	ELKDLVEEIVKLSKENLKLWEDHSRVLEEIVRLKHKNDEREVRRLKLL EDLTPRAEETSRRIEEIVKEAEDRAR SEQ ID NO:260
DHD153	Heterodimer	a	DEERELREVLKHHRVVREWTKVVEELKRVVELLKRGETSEEDLRLVKKL LEM DKRILEVNRVLRVLEKRLT SEQ ID NO:261
DHD153	Heterodimer	b	SLEEIEELVELVRRSVEIAKESDEVARRIVESEDKKELIDTLRDLHREW QEVTKRAEELVREAEKEVR SEQ ID NO:262
DHD154	Heterodimer	a	TAEELLEVHKSDRVTKHELRVSEELKVVVEVLRGEVSEVLKRVLRKLE ELTDKLRVVEEQRRVVEKLN SEQ ID NO:263
DHD154	Heterodimer	b	DLEDLRLRLRVDEQRRLVEELERVSRLEKAVRDNEDERELARLSREHS DIQDKHDKLAREILEVLKRLLELTE SEQ ID NO:264
DHD155	Heterodimer	a	PEDDVVRIKEDLESNREVLREQEIHRIELVTRGEVSEEAIDRVLRKQE DLLKKQKESTDKARKVVEERR SEQ ID NO:265
DHD155	Heterodimer	b	DEVRLITWKLKSEESTRLLKELVELTRLLRNNVPNVEEILREHERISREL ERLSRRLKDLADKLERTRR SEQ ID NO:266
DHD156	Heterodimer	a	DEDEVVKVHEEHVKSHEEIHRSHEEVVRAAEEDKRSRELRTLMEEHRKLL EENEKSIEEVKKIHERVVR SEQ ID NO:267
DHD156	Heterodimer	b	KKEELIDISKEVLDLDEINKISKEILELIKLLRLKEEGREDKDKAREVK RRI RELHRRIQELNKRRLRELHQRVQETKR SEQ ID NO:268
DHD157	Heterodimer	a	PEEDARRVEDLKRSEELIKESKILKESKRLDRNDSKRVLETNLRILI DKHTKLEENLELELLKLAEDVAK SEQ ID NO:269
DHD157	Heterodimer	b	RFKDLRSREYIEVVKRLELSREALEVLRKIDTDKTDKKRIKELIDRLRKL IEYKRIIDRLRKLKDLDEEHR SEQ ID NO:270
DHD158	Heterodimer	a	DEEELVKILKELQRLSEESLEINKRLVEILRLLRRGEVPEEVEKKLREIK KEQEKLDREHEKIKKRIEETK SEQ ID NO:271
DHD158	Heterodimer	b	SLKEKILEIERNMKLVELSNRSVEIVARILKGEKDDEETLERLLREWDKI TRDYEEIIESRKLKVELEEEAK SEQ ID NO:272
DHD159	Heterodimer	a	SKTEILRKALEIHKQIDIVRKLIELSEEVKLKVEESKEKNLEKLKRIDEE TDRLLERLDELHKRLTELAERLK SEQ ID NO:273
DHD159	Heterodimer	b	SDDEARKQLEEMKRRLEVEKSKRVEERVRELERLVRENREDEDVLRKTL EDLLRENEKLVRTIERHVREQRELSKEVK SEQ ID NO:274
DHD160	Heterodimer	a	SEEELEKKADELKRLSEEWRLQEEDKRLSEMVEKGELDLQEVDEHSRLVL ERATEVHRTVDKVIIEILRTTN SEQ ID NO:275
DHD160	Heterodimer	b	SEKERHRESQETQEETRRTHEEIRKLEELRRAKAGELPEETLDRLRIM ERLSELRLDDLVKLRDDHREQK SEQ ID NO:276
DHD161	Heterodimer	a	SEKEILEELKRIKRVKDISDRLEELDKRTEETARREPTKELVDELVKIHR DWLRLHEEILKLVDDALKKVEDATK SEQ ID NO:277
DHD161	Heterodimer	b	DLRELELQREASRLHRELVKLLTELKKELEIAKGEDIREEDLKRIKERL EEIKKRSKRIKESDEIDKTK SEQ ID NO:278
DHD162	Heterodimer	a	SERELQRELNKIVRRI LEIHREVSSELHQRAVKLI RENDNSEEELEISRIE ELSKLEKLVREHDEIVKTI E SEQ ID NO:279
DHD162	Heterodimer	b	SEREKLDNRNDEELKEINKRVEEIKERSDRITEATEKNERSEEEIRRLSREQ NEALQRLLELHKKLVKLHRELEEDTR SEQ ID NO:280

DHD163	Heterodimer	a	DKEDVIRVHDEQHKLIIEEQLELTRRIAELVREIAKNTASEEEIKEMLKEIK RLDDRSREIQDRLQKLLLEEIRKTK SEQ ID NO:281
DHD163	Heterodimer	b	TEEEIVELNKDIQRKSKEHIDLQNELVKKIERAIRENNITEELLEELERLL RESEKIVEEIRRIDTKIRKDAK SEQ ID NO:282
DHD164	Heterodimer	a	SEKEILERLLRLSKEQNEISEEIHRLTERLVELKRRKDDDERLKRILDRQK RLVERAREISKEYEDLLRKE SEQ ID NO:283
DHD164	Heterodimer	b	SMEELLRKNARLSRQKQKIIDEHLELSTKLTRGEAGDETLEEIERRSREML EEQRRVDEESKRIREKLK SEQ ID NO:284
DHD165	Heterodimer	a	SEEEIRDIVEKLLRTHHEVLKEIKKLLDDSERVRRRELDKDLDRIQKEQR DIQEENKEKAKRFDELVKELKKAAS SEQ ID NO:285
DHD165	Heterodimer	b	SEEHRTMEKVEKEVRDIKRRSEEVKKVKANTLSEEDLVRLLELRLVEDH KRLQDLSQEIIERDEKATK SEQ ID NO:286
DHD166	Heterodimer	a	DEDELAKEIEDVQRRNKESQEEHDKSVKLEAAERGEI DEDSLLRVLEEDI KVLEKDIEVLERSIEVIEKAE SEQ ID NO:287
DHD166	Heterodimer	b	SEKELIRRLLEQQRQHLRLSERLIELSRRLVEVVRKGDNRDLLRELKCLS EEHKKHSDHEKVREREREK SEQ ID NO:288
DHS 17	Heterodimer	a	DRKDLLKRNIKLLDRHLKILDTILKLEKLSSELKKSSEEVVKEYKKILD EIRKLLLESKEIHKESKEILERES SEQ ID NO:289
DHD17	Heterodimer	b	DEEKLIERSKRLQEESEQLLEKFEQILRELTELLEKPDSEELARKIKKLHD ELRKIIKRNQELIREHBEILRKR SEQ ID NO:290

In some aspects, non-limiting examples of the monomer A polypeptide and monomer B polypeptide pairs are shown in FIG. 16.

In some aspects, the amino acid sequence of SEQ ID NOs: 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494 useful for the monomer A polypeptide or the monomer B polypeptide is not linked to GlySer at the N terminus of the sequence or does not comprise GlySer at the N terminus. In some aspects, the monomer A polypeptide and/or the monomer B polypeptide comprises at least one amino acid, at least two amino acids, at least three amino acids, at least four amino acids, at least five amino acids, at least six amino acids, at least seven amino acids, at least eight amino acids, at least nine amino acids, or at least ten amino acids at the N terminus or the C terminus of the amino acid sequence. In some aspects, the additional amino acids are not GlySer at the N terminus.

In some aspects, the protein of the present disclosure comprises a heterodimer comprising a monomer A polypeptide and a monomer B polypeptide, wherein the monomer A polypeptide comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence as set forth in SEQ ID NO: 331, 5, 7, 13, 15, 25, 29, 31, 33, 35, 37, 39, 41, 45, 47, 53, 55, 57, 59, 61, 65, 67, 69, 71, 73, 75, 77, 79, 337, 339, 85, 87, 89, 91, 93,

95, 97, 99, 341, 103, 343, 107, 109, 111, 113, 459, 345, 347, 349, 351, 353, 355, 357, 359, 361, 363, 365, 367, 369, 371, 373, 375, 377, 379, 381, 383, 385, 387, 389, 391, 393, 395, 397, 399, 401, 403, 405, 407, 409, 411, 413, 415, 417, 419, or 421 and the monomer B polypeptide comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 5 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence as set forth in SEQ ID NO: 2, 332, 334, 336, 338, 340, 342, 344, 346, 348, 418, 350, 352, 354, 356, 358, 360, 362, 364, 366, 368, 370, 372, 374, 376, 378, 380, 382, 384, 386, 388, 390, 392, 394, 396, 398, 400, 402, 404, 406, 408, 410, 412, 414, 416, 420, 422, 424, 426, 428, 126, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 10 452, 454, 456, 458, 460, 462, 464, 466, 468, 470, 472, 474, 174, 476, 478, 480, 482, 484, 486, 488, 490, 492, or 494, respectively.

In some aspects, the protein of the present disclosure comprises a heterodimer comprising a monomer A polypeptide and a monomer B polypeptide, wherein the monomer A polypeptide comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 15 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of SEQ ID NO: 331, 5, 7, 13, 15, 25, 29, 31, 33, 35, 37, 39, 41, 45, 47, 53, 55, 57, 59, 61, 65, 67, 69, 71, 73, 75, 77, 79, 337, 339, 85, 87, 89, 91, 93, 95, 97, 99, 341, 103, 343, 107, 109, 111, 113, 459, 345, 347, 349, 351, 353, 355, 357, 359, 361, 363, 365, 367, 369, 371, 373, 375, 377, 379, 381, 383, 385, 387, 389, 391, 393, 395, 397, 399, 401, 403, 405, 407, 409, 20 411, 413, 415, 417, 419, or 421, wherein the amino acid sequence of the monomer A polypeptide does not comprise GlySer at the N terminus, and the monomer B polypeptide comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of SEQ ID NO: 2, 332, 334, 336, 338, 340, 342, 344, 346, 348, 418, 350, 352, 354, 356, 358, 360, 362, 25 364, 366, 368, 370, 372, 374, 376, 378, 380, 382, 384, 386, 388, 390, 392, 394, 396, 398, 400, 402, 404, 406, 408, 410, 412, 414, 416, 420, 422, 424, 426, 428, 126, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458, 460, 462, 464, 466, 468, 470, 472, 474, 174, 476, 478, 480, 482, 484, 486, 488, 490, 492, or 494, respectively, wherein the amino acid sequence of the monomer B polypeptide does not comprise GlySer at the N 30 terminus.

In some aspects, the protein of the present disclosure comprises a heterodimer comprising a monomer A polypeptide and a monomer B polypeptide, wherein the monomer A polypeptide consists of an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of

the amino acid sequence as set forth in SEQ ID NO: 331, 5, 7, 13, 15, 25, 29, 31, 33, 35, 37, 39, 41, 45, 47, 53, 55, 57, 59, 61, 65, 67, 69, 71, 73, 75, 77, 79, 337, 339, 85, 87, 89, 91, 93, 95, 97, 99, 341, 103, 343, 107, 109, 111, 113, 459, 345, 347, 349, 351, 353, 355, 357, 359, 361, 363, 365, 367, 369, 371, 373, 375, 377, 379, 381, 383, 385, 387, 389, 391, 393, 395, 5 397, 399, 401, 403, 405, 407, 409, 411, 413, 415, 417, 419, or 421 and the monomer B polypeptide consists of an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence as set forth in SEQ ID NO: 2, 332, 334, 336, 338, 340, 342, 344, 346, 348, 418, 350, 352, 354, 356, 358, 360, 362, 364, 366, 368, 370, 372, 374, 376, 378, 10 380, 382, 384, 386, 388, 390, 392, 394, 396, 398, 400, 402, 404, 406, 408, 410, 412, 414, 416, 420, 422, 424, 426, 428, 126, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458, 460, 462, 464, 466, 468, 470, 472, 474, 174, 476, 478, 480, 482, 484, 486, 488, 490, 492, or 494, respectively.

In one embodiment of any of the above embodiments, amino acid changes from the reference amino acid sequence are conservative amino acid substitutions. As used herein, 15 “conservative amino acid substitution” means an amino acid substitution that does not alter or substantially alter polypeptide function or other characteristics. A given amino acid can be replaced by a residue having similar physiochemical characteristics, e.g., substituting one aliphatic residue for another (such as Ile, Val, Leu, or Ala for one another), or substitution of 20 one polar residue for another (such as between Lys and Arg; Glu and Asp; or Gln and Asn). Other such conservative substitutions, e.g., substitutions of entire regions having similar hydrophobicity characteristics, are well known. Polypeptides comprising conservative amino acid substitutions can be tested in any one of the assays described herein to confirm that a desired activity, e.g. antigen-binding activity and specificity of a native or reference 25 polypeptide is retained.

Amino acids can be grouped according to similarities in the properties of their side chains (in A. L. Lehninger, in *Biochemistry*, second ed., pp. 73-75, Worth Publishers, New York (1975)): (1) non-polar: Ala (A), Val (V), Leu (L), Ile (I), Pro (P), Phe (F), Trp (W), Met (M); (2) uncharged polar: Gly (G), Ser (S), Thr (T), Cys (C), Tyr (Y), Asn (N), Gln (Q); (3) 30 acidic: Asp (D), Glu (E); (4) basic: Lys (K), Arg (R), His (H). Alternatively, naturally occurring residues can be divided into groups based on common side-chain properties: (1) hydrophobic: Norleucine, Met, Ala, Val, Leu, Ile; (2) neutral hydrophilic: Cys, Ser, Thr, Asn, Gln; (3) acidic: Asp, Glu; (4) basic: His, Lys, Arg; (5) residues that influence chain orientation: Gly, Pro; (6) aromatic: Trp, Tyr, Phe. Non-conservative substitutions will entail

exchanging a member of one of these classes for another class. Particular conservative substitutions include, for example; Ala into Gly or into Ser; Arg into Lys; Asn into Gln or into His; Asp into Glu; Cys into Ser; Gln into Asn; Glu into Asp; Gly into Ala or into Pro; His into Asn or into Gln; Ile into Leu or into Val; Leu into Ile or into Val; Lys into Arg, into
 5 Gln or into Glu; Met into Leu, into Tyr or into Ile; Phe into Met, into Leu or into Tyr; Ser into Thr; Thr into Ser; Trp into Tyr; Tyr into Trp; and/or Phe into Val, into Ile or into Leu.

In another embodiment of any of the above embodiments, amino acid residues at 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of defined interface
 10 positions are invariant compared to the reference amino acid sequence. Table 2 below provides the residue numbers within each A and B monomer that are present at the interface in the heterodimer. The position of interface residues are the same for A-B binding partners. Table 2 is organized by heterodimer design name (see the left-hand column in Tables 1A and 1B). Note that for purpose of defining the position of
 15 interface residues for each polypeptide in Table 1A and 1B, the "GS" residues at the amino terminus, if present, are not included.

Table 2. Interface residues by position number across both chains 'a' and 'b'

20	DHD_1 [5, 6, 8, 9, 12, 13, 16, 19, 20, 22, 23, 31, 34, 35, 38, 41, 42, 45, 48, 52, 55, 59, 63, 66, 70, 73, 74, 77, 80, 81, 85, 88, 89, 92, 95, 96, 99, 102, 103, 106]
25	DHD_2 [5, 6, 9, 12, 13, 16, 19, 20, 23, 26, 27, 30, 33, 36, 37, 38, 41, 44, 45, 48, 51, 55, 58, 62, 65, 69, 72, 73, 76, 81, 85, 88, 89, 92, 95, 96, 99, 102, 103, 106, 109, 110, 112, 114, 117, 120, 123, 124, 127, 128, 131, 134, 135, 137, 138, 141, 144, 145, 148, 149, 152]
30	DHD_3 [6, 7, 10, 11, 13, 14, 17, 20, 21, 24, 25, 28, 33, 36, 40, 43, 44, 47, 50, 51, 54, 62, 63, 66, 69, 73, 76, 77, 80, 81, 84, 89, 92, 93, 96, 99, 100, 103, 106, 107, 110, 112]
35	DHD_4 [1, 8, 11, 12, 15, 19, 22, 26, 29, 30, 33, 38, 39, 42, 45, 46, 49, 50, 53, 56, 57, 60, 63, 64, 67, 68, 71, 75, 76, 80, 83, 87, 90, 94, 98, 101, 105, 108, 113, 114, 117, 121, 124, 125, 128, 132, 135, 139, 142, 143, 146, 150]
40	DHD_5 [4, 8, 12, 15, 16, 19, 22, 23, 26, 30, 34, 38, 39, 44, 48, 51, 55, 58, 62, 66, 69, 73, 76, 80, 84, 87, 88, 92, 95, 96, 98, 99, 102, 105, 106, 110, 114, 115, 117, 120, 124, 127, 131, 135, 138, 142, 145, 149, 152]
	DHD_6 [1, 5, 8, 9, 11, 12, 15, 16, 19, 22, 23, 26, 27, 30, 33, 35, 37, 42, 46, 49, 53, 56, 60, 64, 67, 71, 74, 77, 81, 84, 87,

88, 91, 92, 95, 99, 102, 105, 106, 109, 110, 111, 112, 113, 118, 122, 125, 129, 132, 133, 136, 140, 143, 147, 150
]

DHD_7

5 [3, 6, 10, 13, 17, 20, 24, 27, 31, 34, 44, 47, 48, 51, 55, 58, 61, 62, 65, 69, 72, 75, 76, 79, 82, 86, 89, 90, 93, 96, 100, 103, 107, 110, 115, 120, 124, 127, 130, 131, 134, 138, 141, 145, 148, 151, 152]

DHD_8

10 [6, 10, 13, 17, 20, 24, 27, 28, 31, 34, 47, 50, 54, 57, 61, 64, 68, 71, 75, 76, 82, 83, 86, 89, 93, 96, 100, 103, 107, 110, 123, 126, 130, 133, 134, 137, 140, 144, 147, 150, 151, 152]

DHD_9

15 [12, 16, 19, 23, 26, 30, 33, 38, 39, 42, 46, 49, 52, 53, 56, 59, 60, 63, 67, 69, 88, 92, 95, 99, 102, 106, 109, 115, 122, 125, 129, 132, 136, 139, 143, 145]

DHD_16

[6, 9, 10, 13, 16, 17, 20, 23, 24, 27, 28, 30, 34, 43, 46, 50, 53, 57, 60, 64, 67, 71, 72, 5, 8, 12, 15, 19, 22, 23, 26, 29, 33, 41, 50, 57, 60, 64, 67, 68, 71, 74, 78]

20 DHD_18

[3, 6, 9, 10, 13, 16, 17, 20, 23, 24, 27, 31, 34, 38, 43, 46, 50, 53, 57, 60, 64, 67, 68, 71, 75, 78, 81, 85, 88, 91, 95, 98, 102, 105, 106, 109, 118, 121, 125, 128, 132, 135, 136, 139, 142, 146, 149, 150]

DHD_19

25 [3, 6, 9, 10, 13, 16, 17, 20, 23, 24, 27, 30, 31, 34, 38, 43, 46, 50, 53, 57, 60, 61, 64, 67, 71, 75, 81, 88, 91, 95, 98, 99, 102, 105, 109, 118, 121, 125, 128, 129, 132, 135, 139, 142, 146, 149, 150]

DHD_22

30 [1, 2, 5, 6, 9, 10, 12, 13, 16, 19, 20, 23, 27, 30, 31, 33, 34, 38, 41, 44, 47, 48, 51, 55, 58, 62, 65, 69, 72, 80, 81, 84, 87, 91, 94, 97, 98, 101, 102, 105, 108, 118, 122, 125, 126, 129, 132, 136, 139, 143, 146, 147, 148]

DHD_23

35 [1, 5, 6, 8, 9, 12, 15, 16, 19, 22, 23, 26, 29, 30, 33, 34, 40, 43, 47, 51, 54, 57, 58, 61, 64, 65, 68, 72, 75, 76, 79, 82, 83, 86, 89, 90, 93, 96, 97, 100, 104, 109, 110, 113, 116, 120, 123, 126, 127, 130, 134, 138]

DHD_24

[1, 5, 9, 12, 16, 19, 22, 26, 29, 30, 33, 40, 41, 42, 45, 48, 49, 51, 52, 55, 56, 59, 62, 63, 66, 69, 70, 73, 77, 80, 83, 84, 87, 91, 94, 98, 101, 105, 108, 109, 113, 118, 121, 125, 128, 129, 132, 136, 139, 142, 143, 147]

40 DHD_26

[5, 8, 9, 12, 16, 19, 22, 23, 26, 29, 30, 33, 37, 40, 44, 45, 48, 51, 52, 55, 58, 59, 62, 65, 66, 69, 73, 77, 80, 84, 87, 91, 94, 95, 98, 101, 105, 109, 112, 114, 115, 118, 119, 122, 125, 126, 129, 132, 136, 139, 143]

DHD_28

45 [2, 5, 6, 9, 13, 16, 20, 23, 26, 27, 30, 31, 33, 34, 46, 47, 50, 53, 54, 57, 60, 61, 64, 67, 68, 71, 75, 78, 82, 85, 86, 89, 92, 93, 96, 99, 100, 103, 106, 110, 114, 115, 118, 121, 125, 128, 132, 135, 139, 142, 146, 147]

DHD_29

50 [2, 5, 6, 8, 9, 10, 12, 13, 16, 19, 20, 22, 23, 26, 27, 30, 33, 34, 39, 42, 45, 46, 49, 53, 56, 60, 63, 66, 67, 70, 71, 74, 77, 78, 81, 84, 85, 88, 91, 92, 95, 98, 99, 102, 106, 112, 115, 116, 119, 120, 122, 123, 126, 129, 130, 133, 137, 140, 141, 144, 145]

- 5 DHD_31
[2, 5, 6, 9, 10, 12, 13, 16, 17, 19, 20, 23, 24, 26, 27, 30, 31, 33, 34, 37, 38, 39, 43, 46, 50, 54, 57, 60, 61, 64, 65, 68, 71, 75, 78, 81, 82, 85, 88, 89, 92, 95, 96, 99, 102, 103, 106, 109, 110, 119, 123, 126, 130, 133, 134, 137, 140, 141, 144, 147, 148, 151, 152]
- 10 DHD_32
[5, 6, 9, 12, 16, 19, 23, 26, 27, 30, 33, 34, 37, 44, 48, 49, 52, 55, 56, 58, 59, 62, 63, 65, 66, 69, 70, 73, 76, 77, 79, 80, 82, 83, 86, 87, 90, 93, 94, 97, 100, 101, 104, 107, 108, 111, 114, 115, 118, 123, 124, 127, 130, 133, 134, 137, 138, 140, 141, 144, 145, 147, 148, 151, 152]
- 15 DHD_38
[42, 43, 46, 49, 50, 52, 53, 56, 57, 60, 63, 64, 67, 70, 71, 74, 77, 79, 80, 84, 87, 88, 91, 94, 95, 98, 101, 102, 105, 109, 116, 117, 120, 123, 124, 127, 131, 134, 137, 138, 141, 144, 145, 149, 153, 158, 161, 162, 165, 168, 169, 172, 175, 176, 179, 182, 183, 185, 186, 187, 189]
- 20 DHD_39
[2, 6, 9, 13, 16, 20, 23, 27, 29, 30, 34, 40, 43, 46, 47, 50, 51, 54, 57, 58, 60, 61, 64, 110, 114, 117, 121, 124, 128, 131, 132, 135, 138, 145, 149, 152, 156, 159, 160, 163, 164, 166, 167, 170, 173, 174, 176, 177, 178]
- 25 DHD_40
[5, 9, 11, 12, 15, 16, 19, 23, 26, 29, 30, 33, 43, 46, 49, 50, 53, 56, 57, 59, 60, 63, 64, 67, 71, 108, 112, 116, 119, 123, 127, 130, 133, 134, 137, 138, 148, 151, 154, 158, 161, 165, 168, 169, 172, 175]
- 30 DHD_43
[3, 4, 7, 11, 14, 18, 22, 25, 29, 34, 35, 39, 43, 46, 47, 50, 51, 54, 57, 58, 61, 62, 65, 71, 72, 75, 76, 79, 82, 83, 86, 87, 90, 93, 94, 97, 98, 100, 101, 104, 111, 114, 115, 117, 118, 122, 125, 126, 129, 133, 136, 137, 140, 144]
- 35 DHD_60
[6, 7, 9, 10, 13, 16, 17, 20, 23, 24, 27, 30, 31, 34, 35, 51, 54, 55, 58, 61, 62, 65, 68, 72, 75, 76, 77, 83, 87, 90, 91, 93, 94, 97, 100, 101, 104, 105, 108, 111, 115, 116, 121, 122, 124, 125, 128, 131, 132, 135, 136, 138, 139, 142, 143, 149]
- 40 DHD_63
[6, 10, 13, 17, 20, 24, 27, 30, 31, 34, 41, 44, 45, 48, 51, 52, 55, 56, 58, 59, 62, 65, 66, 68, 69, 70, 72, 73, 74, 80, 83, 84, 87, 90, 93, 94, 97, 104, 105, 118, 122, 125, 126, 129, 132, 136, 139, 140, 143, 146, 150]
- 45 DHD_65
[2, 5, 6, 8, 9, 12, 13, 16, 19, 20, 23, 26, 27, 29, 30, 33, 34, 37, 43, 47, 50, 54, 57, 61, 64, 68, 71, 72, 75, 82, 85, 86, 89, 92, 93, 96, 99, 100, 103, 106, 107, 110, 118, 120, 121, 123, 127, 128, 131, 134, 135, 138, 139, 141, 142, 145, 148]
- 50 DHD_66
[6, 10, 13, 16, 17, 20, 24, 27, 31, 34, 42, 43, 46, 49, 50, 53, 54, 56, 57, 60, 63, 64, 67, 68, 71, 78, 81, 82, 85, 88, 89, 92, 95, 99, 102, 103, 107, 112, 113, 115, 116, 119, 123, 126, 127, 130, 133, 137, 140, 141]
- DHD_67
[6, 9, 10, 13, 16, 17, 20, 21, 23, 24, 27, 28, 30, 31, 34, 36, 42, 46, 49, 52, 53, 56, 60, 63, 67, 70, 74, 77, 80, 81, 84, 87, 88, 91, 94, 95, 98, 101, 102, 105, 110, 114, 123, 130, 131, 133, 134, 140, 144, 147, 151]
- DHD_69
[2, 5, 6, 9, 10, 12, 13, 16, 17, 19, 20, 23, 26, 27, 30, 33, 35, 44, 47, 51, 54, 58, 61, 65, 68, 72, 75, 76, 78, 82, 85,

86, 89, 92, 96, 99, 100, 102, 103, 106, 107, 111, 117, 118, 120, 121, 124, 127, 128, 131, 134, 135, 138, 139, 141, 142, 146]

DHD_70

5 [6, 9, 10, 13, 16, 17, 20, 23, 24, 26, 27, 30, 31, 34, 37, 38, 41, 43, 44, 47, 50, 51, 54, 57, 58, 61, 64, 65, 68, 71, 72, 73, 74, 78, 81, 82, 85, 88, 89, 92, 95, 96, 99, 102, 103, 106, 107, 109, 110, 111, 119, 120, 123, 124, 127, 130, 131, 134, 137, 138, 141, 144, 145, 148]

DHD_71

10 [9, 16, 19, 23, 30, 34, 40, 43, 44, 46, 47, 50, 51, 54, 57, 58, 61, 64, 65, 69, 72, 75, 78, 79, 82, 85, 86, 89, 92, 93, 96, 97, 99, 100, 103, 106, 111, 115, 118, 119, 121, 122, 125, 128, 129, 132, 136, 139, 140]

DHD_72

15 [1, 2, 5, 6, 8, 9, 12, 16, 19, 23, 26, 27, 30, 33, 34, 40, 43, 46, 47, 50, 54, 57, 63, 64, 67, 68, 75, 78, 79, 82, 85, 86, 89, 92, 93, 96, 99, 102, 103, 106, 108, 112, 115, 116, 119, 120, 123, 126, 130, 133, 137, 140, 143]

DHD_73

20 [2, 6, 9, 10, 12, 16, 17, 23, 24, 27, 30, 31, 34, 36, 37, 40, 43, 44, 47, 51, 54, 57, 58, 61, 65, 69, 79, 82, 86, 89, 92, 93, 96, 99, 100, 103, 104, 107, 117, 120, 124, 127, 131, 134, 137, 138, 141, 142, 145, 148, 149]

DHD_88

25 [2, 3, 6, 9, 13, 16, 20, 23, 27, 30, 34, 37, 48, 51, 52, 55, 56, 58, 59, 62, 63, 65, 66, 69, 70, 72, 73, 76, 79, 80, 81, 83, 87, 90, 91, 94, 97, 98, 101, 104, 105, 108, 111, 112, 115, 116, 121, 125, 128, 132, 135, 136, 139, 140, 142, 143, 146, 149, 150, 153, 154]

DHD_89

30 [1, 2, 5, 8, 9, 12, 15, 16, 19, 22, 23, 26, 29, 30, 33, 34, 36, 37, 38, 48, 51, 55, 58, 62, 65, 69, 72, 79, 82, 83, 86, 87, 89, 90, 93, 94, 97, 100, 101, 103, 104, 107, 108, 109, 110, 111, 114, 117, 118, 121, 124, 128, 131, 135, 138, 139, 142, 145, 146]

DHD_90

35 [1, 2, 5, 6, 8, 9, 12, 13, 16, 19, 20, 23, 26, 27, 29, 30, 33, 34, 37, 39, 43, 46, 47, 50, 54, 57, 61, 64, 68, 71, 75, 77, 78, 81, 82, 85, 88, 89, 91, 92, 95, 96, 99, 102, 103, 106, 109, 110, 113, 116, 119, 122, 125, 126, 129, 132, 133, 136, 139, 140, 143, 146, 147, 150, 151]

DHD_91

40 [2, 5, 9, 12, 16, 17, 19, 23, 26, 27, 30, 33, 39, 40, 43, 46, 50, 53, 57, 60, 64, 67, 70, 74, 77, 78, 81, 84, 85, 88, 89, 92, 95, 96, 98, 99, 101, 109, 112, 116, 119, 123, 126, 130, 133, 137, 138]

DHD_92

45 [5, 6, 8, 9, 12, 13, 15, 16, 19, 20, 22, 23, 26, 27, 29, 30, 33, 36, 37, 38, 42, 46, 49, 50, 53, 56, 57, 59, 60, 63, 67, 70, 71, 74, 77, 80, 81, 84, 87, 88, 91, 94, 95, 98, 101, 102, 105, 109, 110, 114, 118, 121, 125, 128, 129, 132, 135, 136, 139, 142, 143]

DHD_93

50 [2, 5, 6, 9, 10, 12, 13, 16, 19, 20, 23, 26, 27, 30, 34, 35, 37, 40, 44, 47, 51, 54, 58, 61, 65, 76, 79, 83, 86, 87, 90, 93, 94, 97, 100, 104, 107, 109, 113, 116, 117, 120, 123, 124, 127, 130, 131, 134, 137, 138, 141, 144, 145]

DHD_94

[1, 5, 8, 9, 12, 15, 16, 19, 22, 23, 26, 29, 30, 33, 37, 45, 48, 51, 52, 55, 59, 62, 66, 69, 73, 75, 76, 84, 87, 88, 91, 94, 98, 101, 102, 105, 108, 109, 111, 112, 115, 117, 118, 122, 126, 129, 130, 133, 136, 140, 143, 147, 150, 151, 154]

- 5 DHD_95
[2, 6, 9, 13, 16, 20, 23, 26, 27, 30, 34, 37, 39, 41, 42, 45, 48, 49, 52, 55, 56, 59, 62, 63, 66, 69, 70, 72, 73, 75, 81, 85, 88, 91, 92, 95, 96, 98, 99, 102, 105, 106, 109, 110, 113, 114, 120, 123, 124, 127, 130, 131, 134, 137, 138, 141, 145, 148, 151]
- 10 DHD_96
[1, 5, 6, 9, 10, 12, 13, 16, 19, 20, 23, 26, 27, 30, 33, 34, 42, 45, 46, 49, 52, 53, 56, 59, 60, 63, 66, 67, 70, 74, 75, 77, 81, 84, 88, 91, 95, 98, 102, 105, 114, 118, 119, 122, 123, 126, 129, 130, 132, 133, 136, 137, 139, 140, 143, 144, 146, 147, 150]
- 15 DHD_97
[1, 2, 5, 6, 8, 9, 12, 15, 16, 19, 20, 22, 23, 26, 27, 29, 30, 32, 33, 37, 39, 41, 44, 47, 51, 54, 58, 61, 65, 68, 71, 72, 79, 82, 83, 86, 89, 90, 93, 96, 97, 100, 103, 104, 107, 110, 115, 116, 119, 120, 121, 123, 126, 127, 130, 133, 134, 136, 137, 140, 141, 144, 147]
- 20 DHD_98
[6, 7, 9, 13, 16, 20, 23, 27, 30, 34, 35, 38, 43, 44, 50, 51, 53, 54, 57, 60, 61, 64, 67, 68, 71, 72, 74, 75, 78, 82, 85, 86, 89, 92, 93, 96, 99, 100, 103, 106, 110, 119, 123, 127, 130, 133, 134, 137, 141, 144, 148, 151, 152]
- 25 DHD_99
[2, 5, 6, 9, 12, 16, 19, 20, 23, 26, 30, 33, 34, 46, 50, 53, 57, 60, 64, 67, 70, 71, 74, 78, 82, 85, 89, 92, 96, 99, 103, 106, 110, 113, 123, 126, 130, 133, 137, 140, 141, 144, 147, 148, 151, 154, 155]
- 30 DHD_100
[1, 5, 8, 9, 12, 15, 16, 19, 22, 23, 26, 29, 30, 33, 38, 41, 42, 45, 49, 52, 55, 56, 59, 63, 66, 70, 73, 74, 77, 80, 81, 84, 87, 88, 91, 94, 95, 98, 101, 102, 104, 105, 106, 108, 109, 112, 115, 119, 122, 123, 126, 129, 130, 133, 136, 137]
- 35 DHD_101
[5, 6, 9, 12, 13, 16, 20, 23, 27, 30, 34, 39, 43, 46, 47, 50, 53, 54, 57, 60, 64, 67, 68, 71, 74, 75, 78, 79, 81, 84, 85, 88, 92, 95, 99, 102, 106, 109, 119, 120, 123, 126, 130, 133, 134, 137, 138, 140, 144, 147, 151]
- 40 DHD_102
[6, 9, 13, 16, 20, 22, 23, 27, 30, 34, 38, 44, 47, 51, 54, 55, 57, 58, 61, 65, 68, 69, 72, 75, 80, 83, 86, 90, 93, 97, 100, 104, 107, 108, 112, 113, 117, 121, 124, 128, 131, 132, 135, 138, 142, 145]
- 45 DHD_103
[1, 2, 5, 6, 8, 9, 13, 16, 17, 19, 20, 23, 27, 30, 34, 39, 43, 46, 47, 50, 53, 54, 57, 61, 64, 68, 70, 71, 79, 80, 82, 86, 89, 93, 94, 96, 100, 103, 106, 112, 116, 119, 123, 126, 130, 133, 134, 137, 140]
- 50 DHD_104
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- 55 DHD_105
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- 60 DHD_106
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- 5 DHD_107
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- 10 DHD_108
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- 15 DHD_109
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- 20 DHD_110
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- 25 DHD_111
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- 30 DHD_112
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- 35 DHD_113
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- 40 DHD_114
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- 45 DHD_115
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- 50 DHD_116
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- DHD_117
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- DHD_118
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- 5 DHD_119
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- 10 DHD_120
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- 15 DHD_121
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- DHD_122
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- 20 DHD_123
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- 25 DHD_124
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- 30 DHD_125
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- 35 DHD_126
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- 40 DHD_127
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- 45 DHD_128
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- DHD_129
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- 50 DHD_130
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- DHD_131

[3, 6, 9, 10, 13, 16, 17, 20, 24, 27, 31, 34, 44, 48, 51, 52, 54, 55, 58, 59, 61, 62, 65, 66, 69, 72, 73, 76, 83, 87, 90, 91, 94, 97, 101, 104, 108, 111, 121, 124, 125, 128, 131, 132, 135, 138, 139, 142, 145, 146, 149, 150, 152, 153]

DHD_132

5 [5, 6, 9, 12, 13, 16, 19, 20, 23, 26, 27, 30, 33, 34, 37, 39, 43, 46, 47, 50, 54, 57, 64, 68, 71, 74, 75, 85, 86, 89, 92, 93, 95, 99, 100, 103, 105, 106, 107, 110, 114, 120, 123, 127, 130, 134, 137, 141, 144, 148]

DHD_133

10 [6, 10, 13, 17, 20, 21, 24, 27, 31, 34, 39, 44, 45, 48, 51, 52, 55, 58, 59, 62, 65, 66, 69, 72, 73, 76, 83, 87, 90, 94, 97, 101, 104, 108, 111, 112, 121, 122, 125, 126, 129, 132, 136, 139, 143, 146, 149, 150, 153]

DHD_134

15 [6, 9, 10, 13, 16, 17, 20, 23, 24, 27, 30, 31, 34, 38, 43, 47, 50, 54, 57, 61, 64, 68, 71, 75, 82, 85, 89, 92, 96, 99, 103, 106, 110, 111, 118, 122, 125, 129, 132, 133, 136, 139, 140, 143, 146]

DHD_135

[2, 5, 6, 9, 12, 13, 16, 19, 20, 23, 26, 27, 30, 33, 34, 37, 38, 46, 50, 53, 57, 60, 64, 67, 71, 80, 81, 84, 87, 91, 94, 95, 98, 101, 102, 105, 108, 115, 119, 122, 126, 129, 130, 133, 136, 140, 143]

20 DHD_136

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DHD_137

25 [2, 6, 9, 13, 16, 17, 20, 23, 24, 27, 30, 31, 34, 43, 46, 47, 50, 54, 57, 61, 64, 68, 71, 72, 74, 78, 81, 85, 89, 92, 96, 99, 102, 103, 107, 113, 116, 117, 120, 123, 124, 127, 130, 131, 134, 137, 138, 141]

DHD_138

30 [6, 10, 13, 17, 20, 24, 27, 31, 34, 43, 47, 50, 54, 57, 61, 64, 68, 71, 82, 86, 89, 93, 96, 100, 103, 107, 110, 119, 120, 123, 126, 127, 130, 133, 134, 137, 140, 143, 144, 147, 148, 150, 151]

DHD_139

35 [2, 5, 6, 9, 12, 13, 16, 19, 20, 23, 26, 27, 30, 34, 37, 38, 39, 45, 49, 52, 56, 59, 63, 66, 76, 79, 80, 83, 86, 87, 90, 93, 94, 97, 100, 101, 104, 107, 108, 113, 115, 118, 121, 125, 128, 132, 135, 139, 142, 143, 145, 146]

DHD_140

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40 DHD_141

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DHD_142

45 [1, 2, 5, 6, 9, 12, 13, 15, 16, 19, 20, 23, 24, 26, 27, 30, 33, 40, 44, 47, 51, 54, 58, 61, 68, 69, 72, 75, 78, 79, 82, 85, 86, 89, 92, 95, 96, 99, 100, 103, 106, 107, 110, 120, 121, 124, 127, 131, 134, 138, 141, 145, 148, 152]

DHD_143

50 [2, 6, 9, 13, 16, 17, 20, 23, 27, 30, 34, 39, 44, 48, 51, 55, 58, 62, 65, 66, 69, 72, 73, 83, 87, 90, 94, 97, 101, 104, 108, 111, 121, 124, 125, 128, 131, 132, 135, 136, 138, 139, 142, 145, 146, 149, 150, 152, 153]

DHD_144

- [1, 5, 6, 9, 12, 13, 16, 19, 20, 23, 26, 27, 30, 33, 42, 46, 49, 53, 56, 60, 63, 67, 70, 73, 74, 76, 77, 80, 81, 83, 84, 87, 88, 90, 91, 94, 95, 98, 99, 101, 102, 105, 108, 109, 117, 120, 121, 124, 128, 131, 135, 138, 142, 145, 149, 150]
- 5 DHD_145
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- DHD_146
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- DHD_147
15 [6, 7, 10, 13, 14, 17, 20, 24, 27, 31, 34, 37, 38, 42, 43, 46, 49, 50, 53, 54, 56, 57, 60, 63, 64, 67, 70, 71, 74, 81, 84, 88, 91, 95, 98, 102, 105, 109, 116, 119, 123, 126, 127, 130, 133, 137, 140, 141, 144, 147, 148]
- DHD_149
20 [1, 5, 6, 9, 12, 13, 16, 17, 19, 23, 26, 27, 30, 33, 34, 42, 46, 49, 53, 56, 60, 63, 67, 70, 74, 75, 77, 81, 84, 85, 88, 91, 92, 95, 98, 102, 105, 106, 116, 120, 123, 124, 127, 130, 131, 134, 137, 138, 141, 144, 145, 148]
- DHD_150
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- 25 DHD_151
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- DHD_152
30 [5, 6, 9, 12, 13, 19, 20, 23, 26, 27, 30, 33, 34, 36, 40, 43, 46, 47, 50, 54, 57, 61, 64, 68, 71, 74, 78, 81, 82, 85, 88, 89, 92, 95, 96, 99, 102, 103, 106, 111, 116, 120, 123, 127, 134, 137, 141, 144, 148]
- DHD_153
35 [6, 10, 13, 14, 17, 20, 21, 24, 27, 31, 34, 44, 45, 48, 51, 52, 58, 59, 62, 65, 66, 69, 73, 74, 76, 80, 83, 84, 87, 90, 91, 94, 97, 101, 104, 105, 107, 115, 118, 122, 125, 126, 129, 132, 136, 139, 143]
- DHD_154
40 [2, 5, 6, 9, 12, 16, 19, 20, 23, 26, 27, 30, 33, 34, 40, 43, 47, 50, 54, 57, 61, 64, 68, 71, 72, 74, 78, 81, 85, 88, 92, 95, 99, 102, 106, 109, 115, 116, 119, 122, 123, 126, 129, 133, 136, 137, 140, 143, 144, 147]
- DHD_155
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- 45 DHD_156
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- DHD_157
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- DHD13_2341
[4, 5, 8, 11, 12, 15, 18, 19, 22, 25, 29, 32, 33, 36, 39, 41, 71, 91, 95, 98, 102, 105, 106, 109, 112, 113, 116, 117, 119, 121, 125, 126, 129, 132, 133, 136, 137, 139, 140, 143, 144, 146, 147, 150, 153, 154]
- 5 DHD13_AAAA
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- 10 DHD13_BAAA
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- DHD13_XAAA
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- DHD13_XAAX
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- DHD13_XAXA
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- DHD15
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- 30 DHD17
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- 35 DHD20
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- DHD21
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- DHD25
45 [2, 5, 6, 9, 12, 13, 16, 19, 23, 26, 27, 30, 44, 47, 48, 51, 54, 55, 58, 61, 62, 65, 68, 69, 72, 75, 76, 83, 86, 87, 90, 93, 94, 97, 100, 101, 104, 107, 114, 115, 117, 118, 121, 124, 128, 131, 132, 135, 138, 142, 145]
- DHD27
50 [5, 6, 9, 12, 15, 16, 19, 22, 23, 26, 29, 30, 33, 37, 41, 42, 45, 49, 52, 56, 59, 63, 66, 70, 73, 74, 77, 81, 84, 88, 91, 95, 98, 102, 105, 108, 115, 116, 119, 122, 123, 126, 129, 130, 133, 136, 137, 140, 141, 143, 144, 147, 148]
- DHD30

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DHD33

5 [1, 5, 6, 9, 12, 13, 16, 19, 20, 23, 26, 27, 30, 33, 34, 39, 40, 43, 46, 47, 50, 57, 60, 61, 64, 68, 71, 75, 78, 82, 85, 86, 89, 92, 93, 96, 99, 100, 103, 112, 113, 116, 119, 120, 123, 126, 127, 130, 133, 134, 137, 140, 141, 144]

DHD34_XAAAA

10 [1, 5, 8, 9, 12, 16, 19, 23, 26, 29, 30, 33, 37, 45, 48, 49, 52, 55, 56, 59, 62, 63, 66, 69, 70, 73, 76, 77, 86, 90, 93, 97, 100, 104, 105, 107, 111, 114, 124, 125, 128, 131, 132, 135, 138, 142, 145, 149, 152, 153, 156, 157]

DHD34_XAAXA

15 [1, 5, 8, 9, 12, 16, 19, 23, 26, 29, 30, 33, 37, 45, 48, 49, 52, 55, 56, 59, 62, 63, 66, 69, 70, 73, 76, 77, 86, 90, 93, 97, 100, 104, 105, 107, 108, 111, 114, 124, 125, 128, 131, 132, 135, 138, 142, 145, 149, 152, 153, 156, 157]

DHD34_XAXXA

20 [1, 5, 8, 9, 12, 16, 19, 22, 23, 26, 29, 30, 33, 37, 45, 48, 49, 52, 55, 56, 59, 62, 63, 66, 69, 70, 73, 76, 77, 86, 90, 93, 97, 100, 104, 105, 107, 108, 111, 114, 117, 124, 125, 128, 131, 132, 135, 138, 142, 145, 149, 152, 153, 156, 157]

DHD36

[2, 6, 9, 13, 16, 20, 23, 27, 30, 33, 34, 39, 40, 42, 43, 46, 47, 50, 51, 53, 54, 57, 58, 61, 64, 75, 79, 82, 86, 89, 93, 96, 97, 100, 103, 110, 114, 117, 118, 121, 124, 125, 128, 131, 132, 135, 138, 139, 142, 143]

DHD37_ABXB

25 [2, 5, 9, 11, 12, 15, 16, 19, 23, 26, 29, 30, 33, 37, 43, 46, 49, 50, 53, 56, 57, 59, 60, 63, 64, 67, 77, 81, 84, 88, 91, 92, 95, 98, 99, 102, 105, 113, 116, 119, 120, 123, 126, 130, 133, 134, 137, 140, 141, 144]

DHD37_AXBB

30 [2, 5, 9, 12, 16, 18, 19, 22, 23, 26, 29, 30, 33, 37, 43, 46, 49, 50, 52, 53, 56, 57, 60, 67, 70, 71, 77, 78, 81, 84, 88, 91, 95, 96, 99, 102, 105, 113, 116, 119, 120, 123, 126, 127, 130, 133, 134, 137, 140, 144]

DHD37_AXXB

35 [2, 5, 9, 11, 12, 15, 16, 19, 22, 23, 26, 29, 30, 33, 37, 43, 46, 49, 50, 53, 56, 57, 59, 60, 63, 64, 67, 77, 81, 84, 88, 89, 92, 95, 98, 99, 102, 105, 113, 116, 119, 120, 123, 126, 130, 133, 134, 137, 140, 141, 144]

DHD37_BBBB

40 [5, 9, 12, 16, 19, 22, 23, 26, 30, 33, 42, 43, 46, 50, 53, 56, 57, 60, 63, 64, 66, 67, 74, 77, 78, 81, 84, 85, 88, 92, 95, 98, 99, 102, 105, 108, 111, 113, 116, 119, 123, 126, 130, 133, 137, 140, 141, 144]

DHD37_XBBA

[2, 5, 8, 9, 12, 16, 19, 23, 25, 26, 29, 30, 33, 37, 43, 45, 46, 49, 50, 53, 60, 63, 64, 67, 70, 71, 77, 78, 81, 84, 85, 88, 91, 95, 98, 102, 103, 105, 113, 116, 119, 120, 123, 126, 127, 130, 133, 137, 140, 141, 144]

DHD37_XBXB

45 [2, 5, 9, 11, 12, 15, 16, 19, 23, 26, 30, 33, 37, 43, 46, 50, 53, 56, 57, 59, 60, 63, 64, 67, 77, 81, 84, 88, 92, 95, 98, 99, 102, 105, 113, 116, 119, 120, 123, 126, 130, 133, 134, 137, 140, 141, 144]

XAAX

50 [3, 6, 9, 13, 16, 20, 23, 27, 30, 34, 43, 47, 50, 54, 57, 61, 64, 68, 71, 72, 3, 6, 9, 13, 16, 20, 23, 27, 30, 34, 43, 47, 50, 54, 57, 61, 64, 68, 71, 72]

XAXA

[3, 6, 9, 13, 16, 20, 23, 27, 30, 34, 43, 47, 50, 54, 57, 61, 64, 68, 71, 72, 3, 6, 9, 13, 16, 20, 23, 27, 30, 34, 43, 47, 50, 54, 57, 61, 64, 68, 71, 72]

5

In one embodiment, the monomer A polypeptide and the monomer B polypeptide have their interaction specificity determined by at least one designed hydrogen bond network at the interface between the monomer A and the monomer B. In some aspects, (i) monomer A comprises 1 helix, and monomer B comprises 1 helix; (ii) monomer A comprises 1 helix and monomer B comprises 2 helices; (iii) monomer A comprises 1 helix and monomer B comprises 3 helices, (iv) monomer A comprises 1 helix and monomer B comprises 4 helices; or (v) monomer A comprises 1 helix and monomer B comprises 5 helices, wherein the monomer A and the monomer B comprise a hydrogen bond network, e.g., hydrogen bonds that are capable of being formed by the interface residues according to Table 2. In some aspects, (i) monomer A comprises 2 helices, and monomer B comprises 1 helix; (ii) monomer A comprises 2 helices and monomer B comprises 2 helices; (iii) monomer A comprises 2 helices and monomer B comprises 3 helices, (iv) monomer A comprises 2 helices and monomer B comprises 4 helices; or (v) monomer A comprises 2 helices and monomer B comprises 5 helices, wherein the monomer A and the monomer B comprise a hydrogen bond network, e.g., hydrogen bonds that are capable of being formed by the interface residues according to Table 2. In some aspects, (i) monomer A comprises 3 helices, and monomer B comprises 1 helix; (ii) monomer A comprises 3 helices and monomer B comprises 2 helices; (iii) monomer A comprises 3 helices and monomer B comprises 3 helices, (iv) monomer A comprises 3 helices and monomer B comprises 4 helices; or (v) monomer A comprises 3 helices and monomer B comprises 5 helices, wherein the monomer A and the monomer B comprise a hydrogen bond network, e.g., hydrogen bonds that are capable of being formed by the interface residues according to Table 2. In some aspects, (i) monomer A comprises 4 helices, and monomer B comprises 1 helix; (ii) monomer A comprises 4 helices and monomer B comprises 2 helices; (iii) monomer A comprises 4 helices and monomer B comprises 3 helices, (iv) monomer A comprises 4 helices and monomer B comprises 4 helices; or (v) monomer A comprises 4 helices and monomer B comprises 5 helices, wherein the monomer A and the monomer B comprise a hydrogen bond network, e.g., hydrogen bonds that are capable of being formed by the interface residues according to Table 2. In some aspects, (i) monomer A comprises 5

helices, and monomer B comprises 1 helix; (ii) monomer A comprises 5 helices and monomer B comprises 2 helices; (iii) monomer A comprises 5 helices and monomer B comprises 3 helices, (iv) monomer A comprises 5 helices and monomer B comprises 4 helices; or (v) monomer A comprises 5 helices and monomer B comprises 5 helices, wherein the monomer A and the monomer B comprise a hydrogen bond network, e.g., hydrogen bonds that are capable of being formed by the interface residues according to Table 2.

In a second aspect, the disclosure provides non-naturally occurring polypeptides comprising a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS: 1-290, or the group consisting of SEQ ID NOS: 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494. The amino acid sequences of SEQ ID NOS: 1-290 are provided in Table 1A, and the amino acid sequences of SEQ ID NOS: 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494 are provided in Table 1B, and can be used, for example, to generate the heterodimers of the disclosure.

In some aspects, the disclosure provides non-naturally occurring polypeptides comprising a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS: 1-290, wherein GlySer at amino acid positions 1 and 2 of SEQ ID NO: 1, 55, 81, 83, 101, 105, 115, 117, 119, 121, 123, 125, 127, 131, 133, 135, 137, 139, 141, 143, 145, 147, 149, 151, 153, 155, 157, 159, 161, 163, 165, 167, 169, 171, 173, 175, 177, 179, 181, 183, 185, 187, 189, 191, or 193 are optional, e.g., not present. The amino acid sequences of SEQ ID NOS: 1-290 are provided in Table 1A, and can be used, for example, to generate the heterodimers of the disclosure.

In some aspects, the disclosure provides non-naturally occurring polypeptides comprising a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS: 1-290, wherein GlySer at amino acid positions 1 and 2 of SEQ ID NO: 6, 8, 14, 16, 26, 30, 32,

34, 36, 38, 40, 42, 46, 48, 54, 56, 58, 60, 62, 66, 68, 70, 72, 74, 76, 78, 80, 82, 84, 86, 88, 90, 92, 94, 96, 98, 100, 102, 104, 106, 108, 110, 112, 114, 116, 118, 120, 122, 124, 128, 130, 132, 134, 136, 138, 140, 142, 144, 146, 148, 150, 152, 154, 156, 158, 160, 162, 164, 166, 168, 170, 172, 176, 178, 180, 182, 184, 186, 188, 190, 192, or 194 are

5 optional, e.g., not present.

In some aspects, the disclosure provides non-naturally occurring polypeptides comprising a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS: 331, 332,
10 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494. The amino acid sequences of SEQ ID NOS: 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484,
15 486, 488, 490, 493, and 494 are provided in Table 1B, and can be used, for example, to generate the heterodimers of the disclosure.

In some aspects, the disclosure provides non-naturally occurring polypeptides comprising a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the
20 amino acid sequence selected from the group consisting of SEQ ID NOS: 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, wherein the SEQ ID NOS: 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-
25 460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494 is not linked to GlySer at the immediate N terminus or the polypeptide does not comprise GlySer at the N terminus.

In some aspects, the disclosure provides non-naturally occurring polypeptides comprising a polypeptide consisting of an amino acid sequence at least 70%, 75%,
30 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS: 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494. In some aspects, the disclosure

provides non-naturally occurring polypeptides comprising a polypeptide consisting of the sequence of SEQ ID NOs: 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494.

5 In one embodiment, the amino acid changes from the reference amino acid sequence are conservative amino acid substitutions. In another embodiment, amino acid residues at 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of defined interface positions are invariant compared to the reference amino acid
10 sequence. The defined interface residues are as provided in Table 2.

 In a second aspect, the disclosure provides proteins comprising 2, 3, 4, or more non-naturally occurring polypeptides having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS:
15 1-290, wherein the 2, 3, 4, or more naturally occurring polypeptides are covalently linked. In some aspects, the disclosure provides proteins comprising 2, 3, 4, or more non-naturally occurring polypeptides having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS:
20 1-290 and 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, wherein the 2, 3, 4, or more naturally occurring polypeptides are covalently linked. In some aspects, the sequences of monomer A and monomer B listed herein can be modified (substituted) such that the
25 resulting amino acid sequence maintains a hydrogen bond network of the original amino acid sequence as described in Tables 1A and 1B.

 In this aspect, the proteins can be used to generate scaffolds that can be used for any suitable purpose including but not limited to those disclosed herein. In one embodiment, each of the 2, 3, 4, or more non-naturally occurring polypeptides are
30 different. In another embodiment, the 2, 3, 4, or more non-naturally occurring polypeptides may include 2, 3, 4, or more identical polypeptides. In all embodiments, the 2, 3, 4, or more non-naturally occurring polypeptides may, for example, be covalently linked as part of a fusion protein. The 2, 3, 4, or more non-naturally

occurring polypeptides may each be separated by an amino acid linker. Any suitable amino acid linker may be used.

In some aspects, the linker is a flexible linker. In some aspects, the linker is a GS linker. In other aspects, the GS linker comprises (GGS)_n, (GSEGS)_n (SEQ ID NO:423) or (GGGS)_n (SEQ ID NO:425), wherein n is an integer between 1 and 100. In some aspects, the linker comprises an amino acid sequence having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence as set forth in GSEGS_nGSEGS_n (SEQ ID NO:427) or GSEGS_nGSEGS_nGG (SEQ ID NO:461). In some aspects, the linker comprises an amino acid sequence having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence as set forth in GSEGS_nGSEGS_n (SEQ ID NO:429). In some aspects, the linker comprises an amino acid sequence having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence as set forth in (GSEGS)_n, wherein n is 1 to 10, e.g., 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 (SEQ ID NO:423).

In one embodiment, each of the 2, 3, 4, or more non-naturally occurring polypeptides have at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an odd-numbered SEQ ID NO: selected from the group consisting of SEQ ID NOS:1-290. In one embodiment, each of the 2, 3, 4, or more non-naturally occurring polypeptides have at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an odd-numbered SEQ ID NO: selected from the group consisting of SEQ ID NOS:1-290, wherein GlySer at amino acid positions 1 and 2 of SEQ ID NO: 1, 55, 81, 83, 101, 105, 115, 117, 119, 121, 123, 125, 127, 131, 133, 135, 137, 139, 141, 143, 145, 147, 149, 151, 153, 155, 157, 159, 161, 163, 165, 167, 169, 171, 173, 175, 177, 179, 181, 183, 185, 187, 189, 191, or 193 are optional, e.g., not present. In another embodiment, each of the 2, 3, 4, or more non-naturally occurring polypeptides have at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an even-numbered SEQ ID NO: selected from the group consisting of SEQ ID NOS:1-290. In another embodiment, each of the 2, 3, 4, or more non-naturally occurring polypeptides have at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%,

94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an even-numbered SEQ ID NO: selected from the group consisting of SEQ ID NOS:1-290, wherein GlySer at amino acid positions 1 and 2 of SEQ ID NO: 6, 8, 14, 16, 26, 30, 32, 34, 36, 38, 40, 42, 46, 48, 54, 56, 58, 60, 62, 66,
 5 68, 70, 72, 74, 76, 78, 80, 82, 84, 86, 88, 90, 92, 94, 96, 98, 100, 102, 104, 106, 108, 110, 112, 114, 116, 118, 120, 122, 124, 128, 130, 132, 134, 136, 138, 140, 142, 144, 146, 148, 150, 152, 154, 156, 158, 160, 162, 164, 166, 168, 170, 172, 176, 178, 180, 182, 184, 186, 188, 190, 192, or 194 are optional, e.g., not present. In a further embodiment, the 2, 3, 4, or more non-naturally occurring polypeptides include:

10 (a) polypeptides having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an odd-numbered SEQ ID NO: selected from the group consisting of SEQ ID NOS:1-290, or selected from the group consisting of SEQ ID NOS: 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442,
 15 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494; and

(b) polypeptides having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an even-numbered SEQ ID NO: selected from the group
 20 consisting of SEQ ID NOS:1-290, or selected from the group consisting of SEQ ID NOS: 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494.

In some aspects, the 2, 3, 4, or more non-naturally occurring polypeptides
 25 include:

(a) a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of chain a in Table 1A and/or 1B; and

(b) a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%,
 30 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of chain b in Table 1A and/or 1B.

In some aspects, the protein of the present disclosure comprises a heterotrimer. In some aspects, the heterotrimer comprises a monomer having an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%,

99%, or 100% sequence identity along the length of the amino acid sequence described in Tables 1A and 1B, wherein the amino acid sequence forms a hydrogen bond network, e.g., hydrogen bond network formed by the interface residues according to Table 2. In some aspects, the heterotrimer of the present disclosure comprises at least two monomers, wherein each of the monomers comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence described in Tables 1A and 1B, wherein the amino acid sequence forms a hydrogen bond network, e.g., hydrogen bond network formed by the interface residues according to Table 2.

10 In some aspects, the heterotrimer of the present disclosure comprises at least three monomers, wherein each of the monomers comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence described in Tables 1A and 1B, wherein the amino acid sequence forms a hydrogen bond network, e.g., hydrogen bond network formed by the interface residues according to Table 2.

15 In some aspects, the heterotrimer of the present disclosure comprises at least one heterodimer, wherein the heterodimer comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence described in Tables 3 and 4, wherein the amino acid sequence forms a hydrogen bond network, e.g., hydrogen bond network formed by the interface residues according to Table 2.

20 In some aspects, the heterotrimer of the present disclosure comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence described in Tables 3 and 4, wherein the amino acid sequence forms a hydrogen bond network.

25 In some aspects, the protein of the present disclosure comprises a heterotetramer. In some aspects, the heterotetramer comprises a monomer having an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence described in Tables 1A and 1B, wherein the amino acid sequence forms a hydrogen bond network, e.g., hydrogen bond network formed by the interface residues according to Table 2. In some aspects, the heterotetramer of the present disclosure comprises at least two monomers, wherein each of the monomers comprises an amino

acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence described in Tables 1A and 1B, wherein the amino acid sequence forms a hydrogen bond network, e.g., hydrogen bond network formed by the interface residues according to Table 2.

In some aspects, the heterotetramer of the present disclosure comprises at least three monomers, wherein each of the monomers comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence described in Tables 1A and 1B, wherein the amino acid sequence forms a hydrogen bond network, e.g., hydrogen bond network formed by the interface residues according to Table 2.

In some aspects, the heterotetramer of the present disclosure comprises at least four monomers, wherein each of the monomers comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence described in Tables 1A and 1B, wherein the amino acid sequence forms a hydrogen bond network, e.g., hydrogen bond network formed by the interface residues according to Table 2.

In some aspects, the heterotetramer of the present disclosure comprises at least one heterodimer, wherein the heterodimer comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence described in Tables 1A and 1B, wherein the amino acid sequence forms a hydrogen bond network, e.g., hydrogen bond network formed by the interface residues according to Table 2.

In some aspects, the heterotetramer of the present disclosure comprises at least two heterodimers, wherein each of the two heterodimers comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence described in Tables 1A and 1B, wherein the amino acid sequence forms a hydrogen bond network, e.g., hydrogen bond network formed by the interface residues according to Table 2.

In some aspects, the heterotetramer of the present disclosure comprises at least one heterotrimer, wherein the heterotrimer comprises an amino acid sequence at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or

100% sequence identity along the length of the amino acid sequence described in Tables 3 and 4, wherein the amino acid sequence forms a hydrogen bond network.

In another embodiment, the protein comprises the amino acid sequence having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 5 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS:291, 294, 296, 299, and 302-305. The amino acid sequence of SEQ ID NOS:291, 294, 296, 299, and 302-305 is provided in Table 3. These are merely exemplary such proteins of this aspect of the disclosure, and those of skill in the art will understand that any suitable combination of the monomers 10 of the disclosure can be used in generating the proteins of this aspect.

Table 3

Design name	Oligomerization State	Chain	Design Sequence
DHDSC_9-13-37	Heterotetramer (linker italicized and underlined)	9a-13a-37a	PKEEARELIRKQKELIKEQKKLIKEAKQKSDSRDAERIWKRSREINRESKK INKRIKELIKS <u>GSGSEGSSTKEDILERQRKIIERAQEIHRQOEILE</u> ELERIIRKPGSSEEAMKRMLKLEESLRLKLELLESESAQLLYEQRGSE <u>GSGSEGS</u> SDSDEHLKCLKTFLENLRRHLDRDKHIKQLRDLSENPEDER VKDVIDLSERSVRIVKTVIKIFEDSVRKE SEQ ID NO: 291
DHD9-13	Heterotrimer	9a-13a	PKEEARELIRKQKELIKEQKKLIKEAKQKSDSRDAERIWKRSREINRESKK INKRIKELIKS GSGSEGSSTKEDILERQRKIIERAQEIHRQOEILE ELERIIRKPGSSEEAMKRMLKLEESLRLKLELLESESAQLLYEQR SEQ ID NO: 294
DHD15-37	Heterotrimer	15b-37a	TERKLLERSRRLQEESKRLLDEMAEIMRRIKKLLDDPDSEDIAREIKELLR RLKEIIERNQRIAKEHEYIARERSGPGSGSEGSDEHLKCLKTFLENLRR HLDRDKHIKQLRDLSENPEDERVKDVIDLSERSVRIVKTVIKIFEDSVR KKE SEQ ID NO: 296
DHD13-37	Heterotrimer	13b-37b	TEKRLLEEAEAHREQKEI I KKAQELHRRLEETVRQSGSSEEAKKEAKKIL EEIRELSKRSLELLEIILYLSQEQKSGSEGSDEKLDKLLDTLEK ILQTATKIIDDANKLLEKLRRESERKDPKVVETVYVELLRHEKAVKELLEIA KTHAKKVE SEQ ID NO: 299
OPHD_15-9	Heterotrimer	15b-9a	TERKLLERSRRLQEESKRLLDEMAEIMRRIKKLLDDPDSEDIAREIKELLR RLKEIIERNQRIAKEHEYIARERSGSEGSSEGSSEGSSEGSSEGSSEGSSEGS I KEQKKLIKEAKQKSDSRDAERIWKRSREINRESKKINKRIKELIKS SEQ ID NO: 302
OPHD_37-9	Heterotrimer	37a-9a	PKKEAEELAESEELHDRSEKLERAEQSSNSEARKILEDIERISERIEE ISDRIERLLRSGSEGSSEGSDEKLDKLLDTLEKI LQTATKIIDDANK LLEKLRRESERKDPKVVETVYVELLRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 303
OPHD_13-9	Heterotrimer	13a-9a	TKEDILERQRKIIERAQEIHRQOEILEELERIIRKPGSSEEAMKRMLKLL EESLRLKLELLESESAQLLYEQRGSEGSSEGSSEGSSEGSSEGSSEGS LIKEQKKLIKEAKQKSDSRDAERIWKRSREINRESKKINKRIKELIKS SEQ ID NO: 304
OPHD_9-37	Heterotrimer	9b-37a	PKKEAEELAESEELHDRSEKLERAEQSSNSEARKILEDIERISERIEE ISDRIERLLRSGSEGSSEGSDEHLKCLKTFLENLRRHLDRDKHIKQL RDLSENPEDERVKDVIDLSERSVRIVKTVIKIFEDSVRKE SEQ ID NO: 305

In a third aspect, the disclosure provides protein scaffolds, comprising

a) a first designed component comprised of any number of monomer A polypeptides and/or monomer B polypeptides, each from different heterodimers, connected into a single component by amino acid linkers.

5 b) a second designed component, comprising corresponding monomers for each monomer A and/or monomer B in the first designed component one;

wherein the first and second designed components interact to form the protein scaffold, and wherein each monomer A only interacts in the scaffold with its monomer B binding partner. In one embodiment, the first designed component may comprise the protein of any embodiment or combination of embodiments disclosed herein, and/or the
 10 second designed component may comprise a plurality of individual polypeptides of embodiment or combination of embodiments disclosed herein. In non-limiting embodiments, the first designed component and the second designed component may comprise a set of three (Heterotrimer) or four (Heterotetramer) binding partners as
 15 shown in Table 4. As will be understood by those of skill in the art based on the teachings herein, heterotrimers of the disclosure (including but not limited to the exemplary heterotrimers shown in Table 4) include a first component fusion protein of two polypeptides of the disclosure, and the second component comprises two separate polypeptides that are binding partners of the two polypeptides in the fusion protein.
 20 For example, the DHD9-13 scaffold comprises a first designed component comprising the DHD9 A monomer covalently linked to the DHD13 A monomer, and the second designed component comprises individual DHD9 B and DHD13 B monomers. Different scaffolds are separated in the Table by a blank row. As will be understood by those of skill I the art, these are merely exemplary; the monomers in the first designed
 25 component may be linked in any order, and any monomers may be included in the designed components. As will be further understood by those of skill based on the teachings herein, heterotetramers (including but not limited to the exemplary heterotetramer shown in Table 4) include a first component fusion protein of three polypeptides of the disclosure, and the second component comprises three separate
 30 polypeptides that are binding partners of the three polypeptides in the fusion protein.

Table 4

DHD9-13	Heterotrimer	9a-13a	PKEEARELIRKQKELIKEQKKLIKEAKQKSDSRDAERIWKRSREINRESKK INKRIKELIKSGSEGSSEGSSTKEDILERQRKIIERAQEIHRQQEILE
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			ELERIIRKPGSSEAMKRMKLLEESLRLKELLESEESAQLLYEQR SEQ ID NO: 294
DHD9-13	Heterotrimer	9b	PKKEAEELAESEELHDRSEKLERAEQSSNSEARKILEDIERISERIEE ISDRIERLLRS SEQ ID NO: 2
DHD9-13	Heterotrimer	13b	GSHHHHHGSGSENLYFQGSTKRLLEEAERAHREQKEIKKAQELHRRLE EIVRQSGSSEAKKEAKKILEEIRELSKRSLELLREILYLSQEQK SEQ ID NO: 295
DHD9-13	Heterotrimer	13b	HHHHHHGSGSENLYFQGSTKRLLEEAERAHREQKEIKKAQELHRRLEEI VRQSGSSEAKKEAKKILEEIRELSKRSLELLREILYLSQEQK SEQ ID NO: 431
DHD15-37	Heterotrimer	15b-37a	TERKLLERSRRLQEESKRLLEMAEIMRRIKLLDDPDSEDIAREIKELLR RLKEI IERNQRIAKEHEHYIARERSGPGSGSEGSDEHLKLLKTFLENLRR HLDRLDKHIKQLRDI LSEN PEDERVKDVIDLSERSVRIVKTVIKI FEDSVR KKE SEQ ID NO: 296
DHD15-37	Heterotrimer	15a	TRELLRENI ELAKEHI EIMREI LELLQKMEELLERQSS EDILEELRKI IE RIRELLDRSRKIHES EEIAYKEE SEQ ID NO: 297
DHD15-37	Heterotrimer	37b	GSHHHHHGSGSENLYFQGSDDKELDKLLDTLEKILQTATKIIDDANKLLE KLRRSERKDPKVVETVVELLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 298
DHD15-37	Heterotrimer	37b	HHHHHHGSGSENLYFQGSDDKELDKLLDTLEKILQTATKIIDDANKLLEKL RRSERKDPKVVETVVELLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 433
DHD13-37	Heterotrimer	13b-37b	TEKRLLEEAERAHREQKEIKKAQELHRRLEEIVRQSGSSEAKKEAKKIL EEIRELSKRSLELLREILYLSQEQKSGSEGSDEHLKLLKTFLENLRRHLDRLDKHIKQLR DILSEN PEDERVKDVIDLSERSVRIVKTVIKI FEDSVRKKE SEQ ID NO: 299
DHD13-37	Heterotrimer	13a	TKEDILERQRKII ERAQEIHRQOEI LEELERIIRKPGSSEAMKRMKLLE EESLRLKELLESEESAQLLYEQR SEQ ID NO: 300
DHD13-37	Heterotrimer	37a	GSSHHHHHS SGENLYFQGSDEHLKLLKTFLENLRRHLDRLDKHIKQLR DILSEN PEDERVKDVIDLSERSVRIVKTVIKI FEDSVRKKE SEQ ID NO: 301
DHD13-37	Heterotrimer	37a	SHHHHHHS SGENLYFQGSDEHLKLLKTFLENLRRHLDRLDKHIKQLRDI LSEN PEDERVKDVIDLSERSVRIVKTVIKI FEDSVRKKE SEQ ID NO: 435
OPHD_15-9	Heterotrimer	15b-9a	TERKLLERSRRLQEESKRLLEMAEIMRRIKLLDDPDSEDIAREIKELLR RLKEI IERNQRIAKEHEHYIARERSGSEGSSEGSDEHLKLLKTFLENLRRHLDRLDKHIKQLR I KEQKLI KEAQKSDSRDAERIWKRSREINRESKKINKRIKELIKS SEQ ID NO: 302
OPHD_15-9	Heterotrimer	15a	TRELLRENI ELAKEHI EIMREI LELLQKMEELLEKARGADEDVAKTIKEL LRRLEKEI IERNQRIAKEHEHYIARERS SEQ ID NO: 19
OPHD_15-9	Heterotrimer	9b	PKKEAEELAESEELHDRSEKLERAEQSSNSEARKILEDIERISERIEE ISDRIERLLRS SEQ ID NO: 2
OPHD_37-9	Heterotrimer	37a-9a	PKKEAEELAESEELHDRSEKLERAEQSSNSEARKILEDIERISERIEE ISDRIERLLRS GSEGSSEGSDEHLKLLKTFLENLRRHLDRLDKHIKQLRDI LSEN PEDERVKDVIDLSERSVRIVKTVIKI FEDSVRKKE SEQ ID NO: 303
OPHD_37-9	Heterotrimer	37b	GSDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETVVE LLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 42
OPHD_37-9	Heterotrimer	37b	DDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETVVELL KRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 352
OPHD_37-9	Heterotrimer	9b	PKKEAEELAESEELHDRSEKLERAEQSSNSEARKILEDIERISERIEE ISDRIERLLRS SEQ ID NO: 2

OPHD_13-9	Heterotrimer	13a-9a	TKEDILERQRKIIERAQEIHRQQEILEELERIIRKPGSSSEAMKRMKLL EESLRLKELLELESESAQLLYEQRGSEGGSEGGSPKKEARELIRKQKE LIKEQKLIKEAKQKSDSRDAERIWKRSREINRESKKINKRIKELIKS SEQ ID NO: 304
OPHD_13-9	Heterotrimer	13b	GTEKRLLEEAEARAHREQKEIIKKAQELHRRLEEVRSQSGSSEEAKKEAKKI LEEIRELSKRSLELLREILYLSQEQKGLVPR SEQ ID NO: 4
OPHD_13-9	Heterotrimer	9b	PKKEAEELAESEELHDRSEKLERAEQSSNSEEARILEDIERISERIEE ISDRIERLLRS SEQ ID NO: 2
OPHD_9-37	Heterotrimer	9b-37a	PKKEAEELAESEELHDRSEKLERAEQSSNSEEARILEDIERISERIEE ISDRIERLLRSSEGGSEGGSDSDEHLKLLKTFLENLRRHLDRDLKHIKQL RDILSENPEDEKVDVIDLSERSVRIVKTVIKIFEDSVRKKE SEQ ID NO: 305
OPHD_9-37	Heterotrimer	9a	GSPKKEARELIRKQKELIKEQKLIKEAKQKSDSRDAERIWKRSREINRES KKINKRIKELIKS SEQ ID NO: 1
OPHD_9-37	Heterotrimer	9a	PKEARELIRKQKELIKEQKLIKEAKQKSDSRDAERIWKRSREINRESKK INKRIKELIKS SEQ ID NO: 331
OPHD_9-37	Heterotrimer	37b	GSDDKELDKLLDTLEKILQATATKIIDANKLLEKLRRSERKDPKVVETYVE LLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 42
OPHD_9-37	Heterotrimer	37b	DDKELDKLLDTLEKILQATATKIIDANKLLEKLRRSERKDPKVVETYVELL KRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 352
DHDSC_9-13-37	Heterotetramer	9a-13a-37a	PKEARELIRKQKELIKEQKLIKEAKQKSDSRDAERIWKRSREINRESKK INKRIKELIKSGSEGGSEGGSTKEDILERQRKIIERAQEIHRQQEILE ELERIIRKPGSSSEAMKRMKLLLEESLRLKELLELESESAQLLYEQRGSE GGSEGGSDSDEHLKLLKTFLENLRRHLDRDLKHIKQLRDILSENPEDEK VKDVIDLSERSVRIVKTVIKIFEDSVRKKE SEQ ID NO: 291
DHDSC_9-13-37	Heterotetramer	9b	PKKEAEELAESEELHDRSEKLERAEQSSNSEEARILEDIERISERIEE ISDRIERLLRS SEQ ID NO: 2
DHDSC_9-13-37	Heterotetramer	13b	TEKRLLEEAEARAHREQKEIIKKAQELHRRLEEVRSQSGSSEEAKKEAKKI LEEIRELSKRSLELLREILYLSQEQK SEQ ID NO: 292
DHDSC_9-13-37	Heterotetramer	37b	GSSHHHHHSSGENLYFQGSDDKELDKLLDTLEKILQATATKIIDANKLLE KLRRSERKDPKVVETYVELLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 293
DHDSC_9-13-37	Heterotetramer	37b	SHHHHHSSGENLYFQGSDDKELDKLLDTLEKILQATATKIIDANKLLEK LRRSERKDPKVVETYVELLKRHEKAVKELLEIAKTHAKKVE SEQ ID NO: 437

In these embodiments, the scaffold may be stable up to 95°C and has a guanidine denaturation midpoint of 4 M, as described in the examples that follow.

In some aspects, the heterotrimer or heterotetramer of the present disclosure does not comprise a His tag.

In another aspect, the disclosure provides protein scaffolds, comprising

(a) a fusion protein comprising of 2, 3, 4, or more polypeptides, wherein each polypeptide present in the fusion protein is a non-naturally occurring polypeptide comprises 1-5 alpha helices, wherein adjacent alpha helices may optionally be connected by an amino acid linker;

wherein each polypeptide in the fusion protein is capable of non-covalently interacting with a binding partner, and wherein the fusion protein does not comprise a binding partner for any polypeptide present in the fusion protein; and

5 (b) a binding partner for at least one of the polypeptides present in the fusion protein;

wherein the fusion protein and the binding partner non-covalently interact to form the protein scaffold, wherein an interaction specificity between the binding partner and the at least polypeptide in the fusion protein are determined by at least one hydrogen bond network at the interface between the binding partner and the at least one polypeptide.

10 Binding partners are polypeptides capable of forming heterodimers with a polypeptide present in the fusion protein, and are exemplified above with respect to SEQ ID NO:1-290. The binding partner for at least one polypeptide in the fusion protein may comprise a binding partner for 2, 3, 4, or all polypeptides in the fusion protein. As will be understood, when more than one binding partner is present, they are present as individual binding partner polypeptides, and not linked together.

The fusion protein may comprise 2, 3, 4, or more polypeptides. In certain embodiments, the fusion protein comprises at least 3 or 4 polypeptides in total. Exemplary embodiments of such fusion proteins are provided herein, for example in describing heterotrimer and heterotetramer embodiments in Table 4. The polypeptides in the fusion protein may all be the same, may all be different, or may include both identical and distinct polypeptides. In one specific embodiment, each polypeptide in the fusion protein is a different polypeptide.

In one embodiment,

25 (i) the fusion protein comprises 2, 3, 4, or more polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of SEQ ID NOS: 1-290, or SEQ ID NOS: 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494; and

30 (ii) the binding partner comprises a binding partner as defined herein for each polypeptide in (i), wherein each binding partner has at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group SEQ ID NOS: 1-

290, or selected from the group consisting of SEQ ID NOS: 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494. As described herein, the odd-numbered SEQ ID NOS: between
5 SEQ ID NO:1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494 are noted as “A” monomers and the even-numbered SEQ ID NOS between SEQ ID NO:1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494 are the “B” monomers, with adjacent A and B monomers in Tables 1A and 1B capable of forming heterodimers as described in detail herein. Thus, for example, if the fusion protein included the polypeptide of SEQ ID NO:1, then binding partner may include SEQ ID NO:2, while if the fusion protein included the polypeptide
10 of SEQ ID NO:2, then binding partner may include SEQ ID NO:1. The numerous combinations of fusion protein polypeptides and binding partners will be clear to those of skill in the art based on the teachings herein.

In one embodiment, amino acid changes in the fusion protein and the binding partner from the reference amino acid sequence are conservative amino acid
20 substitutions. In another embodiment amino acid residues at 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of defined interface positions in the polypeptides in the fusion protein and the binding partner are invariant compared to the reference amino acid sequence. In a further embodiment, the at least one hydrogen bond network
25 is asymmetric. In a further embodiment, the binding interface comprises at least 25% hydrophobic residues. In another embodiment, the scaffold is stable up to 95°C and has a guanidine denaturation midpoint of 4 M.

In another embodiment, the disclosure provides methods of forming the
30 designed heterodimer disclosed herein, comprising:

- a) providing two of the monomers as unlinked monomers;
- b) providing the other two monomers as linked monomers

whereby the unlinked monomers associate with their respective monomer of the same heterodimer, and not with any of the other monomers. Further details of this aspect are provided in the examples that follow.

5 In another embodiment, the disclosure provides a designed heterodimer protein comprising:

a) asymmetric buried hydrogen bond networks incorporated into regularly repeating backbone structures; and

b) helix hairpin helix monomers wherein the supercoil phases of the helices are
10 fixed at 0, 90, 180, or 270 degrees and the supercoil twist (ω_0) and helical twist (ω_1) are held constant for either a two layer left handed super coil ($\omega_0=-2.85$ and $\omega_1=102.85$), or a 5 layer untwisted bundle ($\omega_0=0$ and $\omega_1=100$) 27. Further details of this aspect are provided in the examples that follow.

In another embodiment, the disclosure provides uses of the polypeptide, protein,
15 heterodimer protein, protein scaffold, nucleic acid, expression vector, and/or cell of any embodiment or combination of embodiments for any suitable purpose, including but not limited to those disclosed herein such as designing protein logic gates

In a fourth aspect, the disclosure provides fusion proteins comprising a
20 polypeptide of the formula X-B-Z, wherein:

(a) the X domain is a non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein the X domain is capable of non-covalently binding to a first target;

(b) the Z domain is a non-naturally occurring polypeptide comprising 1, 2, or 3
25 alpha helices, wherein the Z domain is capable of non-covalently binding to either (i) a second target that differs from the first target, or (ii) a different non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices; and

(c) the B domain is an amino acid linker;

wherein a combined number of alpha helices from the X domain and the Z domain is
30 4, 5, or 6; and

wherein the X domain and the Z domain interact at a binding interface, wherein the binding interface comprises a hydrogen bond network in which at least one side chain in each alpha helix hydrogen of the X domain bonds with a side chain in an alpha helix in the Z domain, and wherein the binding interface comprises a plurality of hydrophobic residues.

Each helix in the X domain H-bonds with at least one helix in the Z domain and each helix in the Z domain H-bonds with at least one helix in the X domain.

In a fifth aspect, the disclosure provides kits or compositions, comprising at least two
5 fusion proteins comprising the formula X-B-Z, wherein
the B domain in each fusion protein is independently a polypeptide linker;
the X domain in each fusion protein comprises a first non-naturally occurring
polypeptide comprising 1, 2, or 3 alpha helices;
10 the Z domain in each fusion protein comprises a second non-naturally occurring
polypeptide comprising 1, 2, or 3 alpha helices, wherein a combined number of alpha helices
from the X domain and the Z domain in each individual fusion protein is 4, 5, or 6; wherein
the X domain and the Z domain interact at a binding interface, wherein the binding interface
comprises a hydrogen bond network in which at least one side chain in each X domain alpha
15 helix bonds with a side chain in an alpha helix in the Z domain; wherein
the X domain in a first fusion protein is capable of non-covalently binding to a first
target;
the Z domain in a second fusion protein is capable of non-covalently binding to a
second target; and
the X domains and Z domains in each individual fusion protein that are not capable of
20 non-covalently binding to the first target or the second target are capable of non-covalently
binding to an X or a Z domain of a different fusion protein in the plurality of fusion proteins.

The fusion proteins and kits can be used, for example, in the methods disclosed herein
such as for logic gate construction, and for any other suitable use as will be appreciated by
those of skill in the art based on the teachings herein. Specifically, fusion proteins can be
25 used for designing 2-input AND and OR logic gates built from *de novo* designed proteins
that regulate the association of arbitrary protein units ranging from split enzymes to
transcriptional machinery in vitro, and in living cells. Binding interaction cooperativity
makes the gates largely insensitive to stoichiometric imbalances in the inputs, and the
modularity of the approach enables ready extension to 3-input OR, AND, and disjunctive
30 normal form gates. The modularity and cooperativity of the control elements, coupled with
the ability to *de novo* design an essentially unlimited number of protein components, enables
design of sophisticated post-translational control logic over a wide range of biological
functions.

In one embodiment, the Z domain is a non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein the Z domain is capable of non-covalently binding to a second target that differs from the first target. This embodiment is useful, for example, for generating single component dimerizers for use in AND/NOR gates. In another embodiment, the Z domain is a non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein the Z domain is capable of non-covalently binding to a different non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices. This embodiment is useful, for example, for generating 2 or 3-component dimerizers for use in AND/NOR gates.

The first targets and second targets may be any target suitable for an intended use. In non-limiting embodiments, the first target and/or the second target may comprise polypeptides or nucleic acids.

In one embodiment of the kit or composition,

- (i) the first fusion protein has the formula X1-B1-Z1, wherein the X1 domain is capable of non-covalently binding to the first target; and
- (ii) the second fusion protein has the formula X2-B2-Z2, wherein the Z2 domain is capable of non-covalently binding to the second target; and wherein the Z1 and X2 domains are capable of non-covalently binding to each other.

In another embodiment of the kit or composition,

- (i) the first fusion protein has the formula X1-B1-Z1, wherein the X1 domain is capable of non-covalently binding to the first target; and
- (ii) the second fusion protein has the formula X2-B2-Z2,
- (iii) the at least two fusion proteins comprise a third fusion protein of formula X3-B3-Z3, wherein the Z3 domain is capable of non-covalently binding to the second target; wherein
 - (A) the Z1 and X2 domains are capable of non-covalently binding to each other; and
 - (B) the Z2 and X3 domains are capable of non-covalently binding to each other.

In one embodiment of the fusion protein or the kits or compositions, the binding interface comprises at least 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60% or greater hydrophobic residues. The B domain linker may be any suitable amino acid sequence, including but not limited to those described herein. In one embodiment, the B domain for each fusion protein is independently between 6-12, 6-11, 6-10, 7-12, 7-11, 7-10, 8-12, 8-11, 8-10, 9-12, 9-11, 9-10, 10-12, 10-11, 11-12, 6, 7, 8, 9, 10, 11, or 12 amino acids in length.

In another embodiment, the combined number of alpha helices from the X and Z domains in an individual fusion protein is 4. In a further embodiment, the X domain of each fusion protein has 2 alpha helices and the Z domain of each fusion protein has 2 alpha helices. In one embodiment, either the X domain or the Z domain of each fusion protein has 1 alpha helix and the other has 3 alpha helices.

In one embodiment, each X domain and each Z domain comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to SEQ ID NOS:1-290, or selected from the group consisting of SEQ ID NOS: 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, with the proviso that the X domain and the Z domain do not form a heterodimer (a-b) pair. In one embodiment, each X domain and each Z domain comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to SEQ ID NOS:1-290 and 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, with the proviso that the X domain and the Z domain do not form a heterodimer (a-b) pair. In one non-limiting embodiment, at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions as defined in Table 2 are invariant in the polypeptides relative to the reference polypeptide.

A different nomenclature is used in the examples that follow. Table 5 provides correspondence between the names used in the examples and in Tables 1A and 1B. The first column is the numbering used in the examples, while the second column lists the corresponding name in Tables 1A and 1B. For example, polypeptide 1 in the examples is DHD37_ABXB (a), 1' is DHD37_ABXB (b). Polypeptide 2 is DHD15 (a), 2' is DHD15 (b), and so on.

Table 5

- 1: DHD 37_ABXB
- 2: DHD 15
- 3: DHD 131
- 4: DHD 101

- 5: DHD 9
- 6: DHD 150
- 7: DHD 154
- 8: DHD 17
- 5 9: DHD 13_XAAA
- 10: DHD 39
- 11: DHD 155

In one embodiment, each fusion protein independently comprises a polypeptide that is 10 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide having the amino acid sequence of a sequence selected from the group consisting of SEQ ID NO: 302, 303, 306-326, 439, 441, 443, 445, 447, 449, 451, 453, 455, and 457:

- 15 2'-1'_2-residue_linker
 GSTERKLLERSRRLQEESKRLLEDMAEIMRRIKKLLDDPDSEDIAREIKELLRRLKEI IERNQRIAKEHEHYIARE
 RSAADDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVET YVELLKRHEKAVKELLEIAKTHAKK
 VE (SEQ ID NO: 306)
 2'-1'_2-residue_linker
- 20 TERKLLERSRRLQEESKRLLEDMAEIMRRIKKLLDDPDSEDIAREIKELLRRLKEI IERNQRIAKEHEHYIARERS
 AADDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVET YVELLKRHEKAVKELLEIAKTHAKKVE
 (SEQ ID NO:439)
 2'-1'_6-residue_linker
- 25 RSGGSGSPDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVET YVELLKRHEKAVKELLEIAKT
 HAKKVE (SEQ ID NO: 307)
 2'-1'_6-residue_linker
- 30 TERKLLERSRRLQEESKRLLEDMAEIMRRIKKLLDDPDSEDIAREIKELLRRLKEI IERNQRIAKEHEHYIARERS
 GSGSGPDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVET YVELLKRHEKAVKELLEIAKTHA
 KKVE (SEQ ID NO:441)
 2'-1'_12-residue_linker
- 35 GSTERKLLERSRRLQEESKRLLEDMAEIMRRIKKLLDDPDSEDIAREIKELLRRLKEI IERNQRIAKEHEHYIARE
 RSGGSGSPGGSGSPDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVET YVELLKRHEKAVKEL
 LEIAKTHAKKVE (SEQ ID NO: 308)
 TERKLLERSRRLQEESKRLLEDMAEIMRRIKKLLDDPDSEDIAREIKELLRRLKEI IERNQRIAKEHEHYIARERS
 GSGSGPGGSGSPDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVET YVELLKRHEKAVKELLE
 IAKTHAKKVE (SEQ ID NO:443)

- 2'-1'_24-residue_linker
 GSTERKLLERSRRLQEESKRLLEDMAEIMRRIKLLDDPSEDIAREIKELLRRLKEI IERNQRIAKEHEIYIARE
 RSGGSGSPGGSGSPGGSGSPGGSGSPDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETVYE
 LLKRHEKAVKELLEIAKTHAKKVE (SEQ ID NO: 309)
- 5 2'-1'_24-residue_linker
 TERKLLERSRRLQEESKRLLEDMAEIMRRIKLLDDPSEDIAREIKELLRRLKEI IERNQRIAKEHEIYIARERS
 GSGSGSPGGSGSPGGSGSPGGSGSPDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETVVELL
 KRHEKAVKELLEIAKTHAKKVE (SEQ ID NO:445)
- 11-7'
- 10 PEDDVVRIIKEDLESNREVLREQKEIHRILELVTRGEVSEEAI DRV LKRQEDLLKKQKESD KARKVVEERGSE
 GSGSEGS DLEDLLRRLRRLVDEQRRLVEELERVSRRLKAVRDNEDERELARLSREHSDIQDKHDKLAREILEVL
 KRLLERTE (SEQ ID NO: 310)
- 1'-4'
- 15 GSDAYDLDRIVKEHRRLVEEQRELVEELEKLVRRQEDHRVDKESHEILERLERI IRRSTRILTELEKLTDEFER
 RTRGSEGS GSEGS SDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETVVELLKRHEKAVKE
 LLEIAKTHAKKVE (SEQ ID NO: 311)
- 1'-4'
- 20 DAYDLDRIVKEHRRLVEEQRELVEELEKLVRRQEDHRVDKESHEILERLERI IRRSTRILTELEKLTDEFERRT
 RGSEGS GSEGS SDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETVVELLKRHEKAVKELL
 EIAKTHAKKVE (SEQ ID NO:447)
- 4-3'
- GSDDEDLELERLLREYHRVLREYEKLEELRRLYEEYKRGEVSEEE SDRI LREI KEILDKSERLWDLSEE VWR TLL
 YQAE GSEGS GSEGS DEKDYHRRLIEHLEDLVR RHEELIKRQKQVVEELERRGLDERLRRVDRFRRSERWEEVI
 ERFRQVVDKLRKSVE (SEQ ID NO: 312)
- 25 4-3'
- DEDELELERLLREYHRVLREYEKLEELRRLYEEYKRGEVSEEE SDRI LREI KEILDKSERLWDLSEE VWR TLLYQ
 AEGSEGS GSEGS DEKDYHRRLIEHLEDLVR RHEELIKRQKQVVEELERRGLDERLRRVDRFRRSERWEEVIER
 FRQVVDKLRKSVE (SEQ ID NO:449)
- 3-2'*
- 30 GSTERKLLERSRRLQEESKRLLEDMAEIMRRIKLLKKARGADEKVLDELRKI IERIRELLDRSRKIHERSEEIA
 YKEEGSEGS GSEGS SDES DRIRKIVEESDEIVKESRKLAEARELIKESDKRVSEERNERLLEELLRILDENA
 ELLKRNLELLKEVLYRTR (SEQ ID NO: 313)
- 3-2'*
- 35 TERKLLERSRRLQEESKRLLEDMAEIMRRIKLLKKARGADEKVLDELRKI IERIRELLDRSRKIHERSEEIAYK
 EEGSEGS GSEGS SDES DRIRKIVEESDEIVKESRKLAEARELIKESDKRVSEERNERLLEELLRILDENAEL
 LKRNLELLKEVLYRTR (SEQ ID NO:451)
- 1'-3'
- 40 GSDDEDLELERLLREYHRVLREYEKLEELRRLYEEYKRGEVSEEE SDRI LREI KEILDKSERLWDLSEE VWR TLL
 YQAE GSEGS GSEGS SDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETVVELLKRHEKAVK
 ELLEIAKTHAKKVE (SEQ ID NO: 314)
- 1'-3'

DEDELERLLREYHRVREYKLLLEELRRLYEYKRGVSEEEEDRIEREIKEILDKSERLWDLSEEVWRTLLYQ
 AEGSESGSGSESGSDDKELDKLLDTLEKILQATATKIIDDANKLLEKLRRSERKDPKVVETYVELLKRHEKAVKEL
 LEIAKTHAKKVE (SEQ ID NO:453)
 1'-5

5 PKKEAEELAESEELHDRSEKLHERAEQSSNSEARKILEDIERISERIEEISDRIERLLRSGSESGSESGSD
 DKELDKLLDTLEKILQATATKIIDDANKLLEKLRRSERKDPKVVETYVELLKRHEKAVKELLEIAKTHAKKVE
 (SEQ ID NO: 303)
 5'-2'

10 TERKLLERSRRLQEE SKRLLEDMAEIMRRIKLLDDPSEDIAREIKELLRRLKEI IERNQRIAKEHEYIARERS
 GSESGSESGSGSPKEEARELIRKQKELIKEQKKLIKEAKQKSDSRDAERIWKRSREINRESKKINKRIKELIKS
 (SEQ ID NO: 302)
 1-6

15 DSDEHLKCLKTFLLENLRRHLDRDKHIKQLRDI LSEN PEDERVKDVIDLSERSVRIVKTVIKI FEDSVRKKEGSE
 GSGSESGSESGSESGSESGSPTDEVIEVLKELLRIHRENLRVNEEIVEVNERASRVTDREELERLLRR
 SNELIKRSRELNEESKKLIEKLERLAT (SEQ ID NO: 315)
 1'-7

20 DDKELDKLLDTLEKILQATATKIIDDANKLLEKLRRSERKDPKVVETYVELLKRHEKAVKELLEIAKTHAKKVEGS
 EGSSEGSTAEELLE VHKKSDRVTK EHLRVSEEILKVVEVLTRGEVSSEVLKRVLRKLEELTDKLRVTEEQRRV
 VEKLN (SEQ ID NO: 316)
 6'-7

25 DNEEIIKEARRVVEEYKKA VDRLEELVRRRAENAKHASEKELKDIVREILRISKELNKVSERLIELWERSQERARG
 SEGSGSEGSTAEELLE VHKKSDRVTK EHLRVSEEILKVVEVLTRGEVSSEVLKRVLRKLEELTDKLRVTEEQRR
 VVEKLN (SEQ ID NO: 317)
 1'-6-7

30 DDKELDKLLDTLEKILQATATKIIDDANKLLEKLRRSERKDPKVVETYVELLKRHEKAVKELLEIAKTHAKKVEGS
 EGSSESGSESGSESGSESGSPTDEVIEVLKELLRIHRENLRVNEEIVEVNERASRVTDREELERLLR
 RSNELIKRSRELNEESKKLIEKLERLATGSESGSESGSESGSESGSESGSEGSTAEELLE VHKKSDRVTK
 EHLRVSEEILKVVEVLTRGEVSSEVLKRVLRKLEELTDKLRVTEEQRRVVEKLN (SEQ ID NO: 318)
 11-1

35 DSDEHLKCLKTFLLENLRRHLDRDKHIKQLRDI LSEN PEDERVKDVIDLSERSVRIVKTVIKI FEDSVRKKEGSE
 GSGSESGPEDDVVRI IKEDLESNREVLREQKEIHRI LELVTRGEVSEE AIDRVLKRQEDLLKKQKESTDKARKVV
 EERR (SEQ ID NO: 319)
 11-6'

40 DNEEIIKEARRVVEEYKKA VDRLEELVRRRAENAKHASEKELKDIVREILRISKELNKVSERLIELWERSQERARG
 SEGSGSESGPEDDVVRI IKEDLESNREVLREQKEIHRI LELVTRGEVSEE AIDRVLKRQEDLLKKQKESTDKARK
 VVEERR (SEQ ID NO: 320)
 11-7'

40 DLEDLLRRLRRLVDEQRRLVEELERVSRRLKAVRDNEDERELARLSREHSDIQDKHDKLAREILEVLKRLRLERT
 EGSESGSESGSESGSESGSPTDEVIEVLKELLRIHRENLRVNEEIVEVNERASRVTDREELERLLRR
 RVLKRQEDLLKKQKESTDKARKVVEERR (SEQ ID NO: 321)
 1'-6

- GSDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETYVELLKRHEKAVKELLEIAKTHAKKVE
 GSEGSSEGSPTDEVIEVLKELLRIHRENLRVNEEIVEVNERASRVTDREELERLLRRSNELIKRSRELNEESKK
 LIEKLERLAT (SEQ ID NO: 322)
 1'-6
- 5 DDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETYVELLKRHEKAVKELLEIAKTHAKKVEGS
 EGSSEGSPTDEVIEVLKELLRIHRENLRVNEEIVEVNERASRVTDREELERLLRRSNELIKRSRELNEESKKLI
 EKLERLAT (SEQ ID NO:455)
 7-1
- 10 DSDEHLKCLKTFLLENLRRHLDRDKHIKQLRDLSENPEDEKVDVIDLSERSVRIVKTVIKIFEDSVRKKEGSE
 GSGSEGSTAEELLEHVHKS DRVTKHELRVSEEILKVVEVLTRGEVSSEVLKRVLRKLEELTDKLRVTEEQRRVV
 EKLN (SEQ ID NO: 323)
 4'-2'*
- 15 GTERKLLERSRRLQEEKRLLEMAEIMRRIKLLKKARGADEKVLDELRKIIERIRELLDRSRKIIHERSEEIAY
 KEEGSEGSSEGSSEGS DAYDLDRIVKEHRLVVEEQRELVEELEKLVRRQEDHRVDDKESHEILERLERIIRSTR
 LTELEKLTDEFERRTR (SEQ ID NO: 324)
 2*-1'
- TREELLRENIELAKEHIEIMREILELLQKMEELLEKARGADEDVAKTIKELLRRLKEIERNQRIAKEHEHYIARE
 RSGSEGSSEGSSEGSDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETYVELLKRHEKAVKEL
 LEIAKTHAKKVE (SEQ ID NO: 325)
- 20 1'-9
- GSDDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETYVELLKRHEKAVKELLEIAKTHAKKVE
 GSEGSSEGSSEGSPTKEDILRQRKIIERAQEIHRROQEIILEELERIIRKPGSSEEAMKRMLKLLLEESLRLKELLE
 SEESAQLLYEQR (SEQ ID NO: 326)
 1'-9
- 25 DDKELDKLLDTLEKILQTATKIIDDANKLLEKLRRSERKDPKVVETYVELLKRHEKAVKELLEIAKTHAKKVEGS
 EGSSEGSSEGSPTKEDILRQRKIIERAQEIHRROQEIILEELERIIRKPGSSEEAMKRMLKLLLEESLRLKELLE
 ESAQLLYEQR (SEQ ID NO:457)

In some aspects, each fusion protein independently comprises a polypeptide that is
 30 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%,
 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group
 including, but not limited to, a polypeptide having the amino acid sequence of SEQ ID NO:
 302, 303, 306-326, 439, 441, 443, 445, 447, 449, 451, 453, 455, and 457, wherein GlySer at
 35 amino acid residues 1 and 2 of any of 302, 303, 306-326, 439, 441, 443, 445, 447, 449, 451,
 453, 455, and 457 are optional, e.g., not present.

In another embodiment, the kits or compositions further comprising the first target
 and the second target. In one embodiment, the first target and the second target each
 independently comprise a polypeptide of the formula X10-B10-Z10, wherein

(a) the X10 domain is a non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices;

(b) the Z10 domain is a non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices; and

5 (c) the B10 domain is an amino acid linker;

wherein the X domain and the Z domain interact at a target binding interface, wherein the target binding interface comprises a hydrogen bond network in which at least one side chain in each alpha helix hydrogen of the X domain bonds with a side chain in a different alpha helix in the Z domain, and wherein the target binding interface comprises a plurality of
10 hydrophobic residues. In one embodiment, the target binding interface comprises at least 25% hydrophobic residues. In another embodiment, the B10 domain for the first target and the second target is independently between 6-12, 6-11, 6-10, 7-12, 7-11, 7-10, 8-12, 8-11, 8-10, 9-12, 9-11, 9-10, 10-12, 10-11, 11-12, 6, 7, 8, 9, 10, 11, or 12 amino acids in length. In another embodiment, the combined number of alpha helices from the X and Z domains in the
15 first target and the second target protein is 4. In a further embodiment,

(a) the X10 domain of each of the first target and the second target has 2 alpha helices and the Z10 domain of each of the first target and the second target has 2 alpha helices; or

(b) either the X10 domain or the Z10 domain of each of the first target and the
20 second target has 1 alpha helix and the other has 3 alpha helices. In one embodiment, each X10 domain and each Z10 domain comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide having the amino acid sequence selected from SEQ ID NOS: 1-290, 331,
25 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, with the proviso that the X10 domain forms a heterodimer (a-b) pair with the X domain of the fusion protein, and the Z10 domain forms a heterodimer (a-b) pair with the Z domain of the fusion protein. In one embodiment, at least 20%, 25%, 30%, 35%, 40%, 45%,
30 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of the first target and/or the second target are invariant compared to the reference polypeptide amino acid sequence (interface residues shown in Table 2).

In another embodiment, the first target and/or the second target further comprise one or more effector polypeptide domains linked to one or more of the X10 and/or Z10 domains, for example, wherein the one or more effector polypeptide domains may comprise a polypeptide including, but not limited to, nucleic acid binding proteins, transcription factors, receptor binding proteins, split enzymes, effectors of membrane receptors, etc.

In a sixth aspect, the disclosure provides methods, comprising:

(i) contacting the fusion protein of embodiment or combination of embodiments of the fifth or sixth aspects disclosed herein with a biological sample under conditions to promote non-covalent binding of the fusion protein with first target and second target present in the sample, and

(ii) detecting non-covalent binding of the one or more fusion proteins to the first target and/or the second target in the biological sample.

The detecting may comprise any suitable means for detecting binding, including but not limited to mass spectrometry, yeast-2-hybrid detection, functional assays, or any other suitable assay as will be clear to those of skill in the art based on the current disclosure. In one embodiment, the method comprises detecting cooperative non-covalent binding of the one or more fusion proteins to the first target and the second target in the biological sample. This embodiment comprises use of the fusion proteins in AND gate logic, as described in more detail in the examples that follow. As used herein, “cooperative” binding means binding the fusion protein cannot bind to the first target without also binding to the second target, and the fusion protein cannot bind to the second target without binding to the first target.

In another embodiment, the method comprises detecting non-covalent binding of the one or more fusion proteins to the first target or the second target in the biological sample. This embodiment comprises use of the fusion proteins in OR gate logic, as described in more detail in the examples that follow.

In another embodiment, the disclosure provides methods comprising:

(a) contacting a biological sample with at least two fusion proteins, wherein each of the at least two fusion proteins comprises the formula X-B-Z, wherein each B is independently a polypeptide linker; each X domain comprises a first non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices; each Z domain comprises a second non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein a combined number of alpha helices from the X domain and the

Z domain in each individual fusion protein is 4, 5, or 6; wherein the X domain and the Z domain interact at a binding interface, wherein the binding interface comprises a hydrogen bond network in which at least one side chain in each X domain alpha helix bonds with a side chain in an alpha helix in the Z domain; wherein

5 the X domain in a first fusion protein is capable of non-covalently binding to a first target;

 the Z domain in a second fusion protein is capable of non-covalently binding to a second target; and

 the X domains and Z domains in each individual fusion protein that are not capable of
10 non-covalently binding to the first target or the second target are capable of non-covalently binding to an X or a Z domain of a different fusion protein in the plurality of fusion proteins;

(b) detecting non-covalent binding of the two or more fusion proteins to the first target and/or the second target in the biological sample. This embodiment comprises use of the fusion proteins in 2 component AND or OR gate logic, as described in more detail in the
15 examples that follow.

In one embodiment of the AND or OR gate logic, the detecting comprises detecting cooperative non-covalent binding of the two or more fusion proteins to the first target and the second target in the biological sample. In another embodiment,

(i) the first fusion protein has the formula X1-B1-Z1, wherein the X1 domain is
20 capable of non-covalently binding to the first target; and

(ii) the second fusion protein has the formula X2-B2-Z2, wherein the Z2 domain is capable of non-covalently binding to the first target; and wherein the Z1 and X2 domains are capable of non-covalently binding to each other.

In a further embodiment,

25 (i) the first fusion protein has the formula X1-B1-Z1, wherein the X1 domain is capable of non-covalently binding to the first target; and

(ii) the second fusion protein has the formula X2-B2-Z2,

(iii) the at least two fusion proteins comprise a third fusion protein of formula X3-B3-Z3, wherein the Z3 domain is capable of non-covalently binding to the second target;
30 wherein

(A) the Z1 and X2 domains are capable of non-covalently binding to each other; and

(B) the Z2 and X3 domains are capable of non-covalently binding to each other.

In another embodiment, the X domains, Y domains, B domains, and or fusion proteins are as recited in any embodiment or combination of embodiments disclosed herein, such as in the fourth and fifth aspects. In one embodiment, at least one of the fusion proteins comprises one or more effector polypeptide domains linked to one or more of the X and/or Z domains, and wherein the detecting step comprises detecting an output signal caused by binding the first target and/or the second target. In another embodiment, the detecting step comprises detecting an output signal from the one or more effector polypeptide caused by cooperative non-covalently binding of the first target and the second target. Such detection may be by any suitable means dependent in part on the output signal to be detected, including but not limited to those disclosed herein. The output signal to be detected may be any suitable output signal including but not limited to fluorescence activity, functional activity, etc.

Any suitable effector polypeptide domain may be employed as suitable for an intended use. In one embodiment, the one or more effector polypeptide domains may comprise a polypeptide including, but not limited to, nucleic acid binding proteins, transcription factors, receptor binding proteins, nucleic acid binding proteins, transcription factors, receptor binding proteins, split enzymes, effectors of membrane receptors, etc.

In a seventh aspect, the disclosure provides compositions comprising

(a) a first polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding a first target; and

(b) a second polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding to the second polypeptide, and wherein the second polypeptide is capable of non-covalently binding a second target that differs from the first target; wherein:

(i) a binding affinity of the first polypeptide for the first target is approximately equal to a binding affinity of the second polypeptide for the second target; and

(ii) the binding affinity of the first polypeptide for the first target and the binding affinity of the second polypeptide for the second target are greater than the binding affinity of the first target and the second target for each other.

Compositions of this seventh aspect can be used, for example, as NOR gates as described in detail in the examples that follow.

In one embodiment, the composition further comprises the first target and the second target. The first targets and second targets may be any target suitable for an intended use. In non-limiting embodiments, the first target and/or the second target may comprise

polypeptides or nucleic acids. In another embodiment, the first target and/or the second target further comprise one or more effector polypeptide domains. Any effector polypeptide domains may be used as suitable for an intended use. In one embodiment, the one or more effector polypeptide domains may comprise a polypeptide including, but not limited to,

5 nucleic acid binding proteins, transcription factors, receptor binding proteins, split enzymes, effectors of membrane receptors, etc. In another embodiment, the first polypeptide and/or the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide

10 having the amino acid sequence selected from the group consisting of SEQ ID NOS:1-290 and 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, as listed in Tables 1A and 1B. In one embodiment, at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%,

15 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of the first polypeptide and/or the second polypeptide are invariant compared to the reference polypeptide amino acid sequence (interface residues shown in Table 2).

In one non-limiting and exemplary embodiment,

(a) the first polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%,

20 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one having the amino acid sequence of SEQ ID NO:3

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GTKEDILERQRKII ERAQEI HRRQQEILEELERII RKPGSSEEAMKRMLKLL EESL RLLKELLELSEESAQLLYE

25 QR (SEQ ID NO: 3); and

(b) the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one

30 having the amino acid sequence of SEQ ID NO:58.

10'

GSSADDVLEDILKII RELIEILDQIL SLLNQLL KLLRHGVPNAKKVVEKYKEILELYLQLVSLFLKIVKTHADAV
SGKIDKKAEEI IKKEEKI KEKLRQAKDILKKLQEEIDKTR (SEQ ID NO: 58)

In one non-limiting and exemplary embodiment,

(a) the first polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one having the amino acid sequence of SEQ ID NO:3

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GTKEDILERQRKII ERAQEI HRRQQEILEELERII RKP GSSEEAMKRMLK LLEESL RLLKELLELSEESAQLLYE
QR (SEQ ID NO: 3); and

(b) the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one having the amino acid sequence of SEQ ID NO:362.

10 10'

SADDVLEDILKII RELIEILDQI LSLNQLLKL LRHGVPNAKKVVEKYKEILELYLQLVSLFLKIVKTHADAVSG
KIDKKAEEI KKEEIKI KEKLRQAKDILKKLQEEIDKTR (SEQ ID NO: 362)

15

In another embodiment, the first target and/or the second target each comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide having the amino acid sequence selected from the group consisting of SEQ ID NO:1-290, with the proviso that the first target forms a heterodimer (a-b) pair with the first polypeptide, and the second target forms a heterodimer (a-b) pair with the second polypeptide. In another embodiment, the first target and/or the second target each comprises a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide having the amino acid sequence selected from the group consisting of SEQ ID NO:1-290 and 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, with the proviso that the first target forms a heterodimer (a-b) pair with the first polypeptide, and the second target forms a heterodimer (a-b) pair with the second polypeptide. Heterodimer A-B pairs among the polypeptides of SEQ ID NOS:1-290 and 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494 are described at length above (See also FIG. 16).

In one embodiment, at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of the first target and/or the second target are invariant compared to the reference polypeptide amino acid sequence (interface residues
5 shown in Table 2).

The compositions of this seventh aspect can be used for any suitable purpose, including in designing NOR logic gates. In one embodiment, the disclosure provides methods comprising

- (a) contacting a biological sample with the composition of any embodiment or
10 combination of embodiments of the seventh aspect of the disclosure; and
- (b) detecting binding, of the first polypeptide to the first target and binding of the second polypeptide to the second target in the sample, such as detecting an output signal caused by actions of effector polypeptides upon binding. Additional details of the use of the compositions of the seventh aspect of the disclosure in NOR logic gates re described in detail
15 in the examples that follow.

In an eighth aspect, the disclosure provides compositions comprising:

- (a) a first polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding a first target; and
20
- (b) a second polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding to the second polypeptide, and wherein the second polypeptide is capable of non-covalently binding a second target that differs from the first target; wherein:
 - (i) a binding affinity of the first polypeptide for the second polypeptide is
25 greater than a binding affinity of the second polypeptide for the second target;
 - (ii) a binding affinity of the first polypeptide for the first target is approximately equal to a binding affinity of the second polypeptide for the second target; and
 - (iii) the binding affinity of the first polypeptide for the first target and the binding affinity of the second polypeptide for the second target are greater than the binding
30 affinity of the first target and the second target for each other.

Compositions of this eighth aspect can be used, for example, as XNOR gates as described in detail in the examples that follow. In one embodiment, the composition further comprises the first target and the second target. The first targets and second targets may be any target suitable for an intended use. In non-limiting embodiments, the first target and/or

the second target may comprise polypeptides or nucleic acids. In another embodiment, the first target and/or the second target further comprise one or more effector polypeptide domains. Any effector polypeptide domains may be used as suitable for an intended use. In one embodiment, the one or more effector polypeptide domains may comprise a polypeptide including, but not limited to, nucleic acid binding proteins, transcription factors, receptor binding proteins, split enzymes, effectors of membrane receptors, etc. In another embodiment, the first polypeptide and/or the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide having the amino acid sequence selected from the group consisting of SEQ ID NOS: 1-290 and 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, as listed in Tables 1A and 1B. In one embodiment, at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of the first polypeptide and/or the second polypeptide are compared to the reference polypeptide amino acid sequence (interface residues shown in Table 2).

In one non-limiting and exemplary embodiment,

(a) the first polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one having the amino acid sequence of SEQ ID NO:3

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 25 GTKEDILERQRKII ERAQEI HRRQQEILEELERII RKPGSSEEAMKRMLKLEESLRLLLKELLEELSEESAQLLYE
 QR (SEQ ID NO: 3); and

(b) the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one having the amino acid sequence of SEQ ID NO:42.

1' (b)
 30 GSDDKELDKLLD'TLEKILQTATKI IDDANKLLEKLRRSERKDPKVVETVVELLKRHEKAVKELLEIAKTHAKKVE
 (SEQ ID NO: 42)

In one non-limiting and exemplary embodiment,

(a) the first polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one having the amino acid sequence of SEQ ID NO:3

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GTKEDILERQRKII ERAQEI HRRQQEILEELERII RKPGSSEEAMKRMLKLEESLRLKELLELSEESAQLLYE
QR (SEQ ID NO: 3); and

(b) the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one having the amino acid sequence of SEQ ID NO:352.

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1' (b)

DDKELDKLLDTLEKILQTATKIIDANKLLEKLRRSERKDPKVVETVVELLKRHEKAVKELLEIAKTHAKKVE
(SEQ ID NO: 352)

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In another embodiment, the first target and/or the second target each comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide having the amino acid sequence selected from the group consisting of SEQ ID NO:1-290 and 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, with the proviso that the first target forms a heterodimer (a-b) pair with the first polypeptide, and the second target forms a heterodimer (a-b) pair with the second polypeptide. Heterodimer A-B pairs among the polypeptides of SEQ ID NOS:1-290 and 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494 are described at length above (See also FIG. 16). In one embodiment, at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of the first target and/or the second target are invariant compared to the reference polypeptide amino acid sequence (interface residues shown in Table 2).

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The compositions of this eighth aspect can be used for any suitable purpose, including in designing XNOR logic gates. In one embodiment, the disclosure provides methods comprising

- 5 (a) contacting a biological sample with the composition of any one of claims 50-56; and
- (b) detecting binding interactions between the first polypeptide and the first target, the second polypeptide and the second target, the first polypeptide and the second polypeptide, and the first target and the second target in the sample, such as detecting an output signal caused by actions of effector polypeptides upon binding. Additional details of
10 the use of the compositions of the eighth aspect of the disclosure in XNOR logic gates re described in detail in the examples that follow.

In a ninth aspect, the disclosure provides compositions comprising:

- 15 (a) a first polypeptide comprising 4 alpha helices, wherein the first polypeptide is capable of non-covalently binding a first target; and
- (b) a second polypeptide comprising 4 alpha helices, wherein the second polypeptide is capable of non-covalently binding a second target that differs from the first target; wherein:
 - 20 (i) a binding affinity of the first target for the second target is greater than a binding affinity of the first polypeptide for the first target;
 - (ii) a binding affinity of the first polypeptide for the first target is approximately equal to a binding affinity of the second polypeptide for the second target; and
 - (iii) the sum of the binding affinity of (A) the first polypeptide for the first target and (B) the binding affinity of the second polypeptide for the second target, is greater
25 than the binding affinity of the first target and the second target.

Compositions of this ninth aspect can be used, for example, as NAND gates as described in detail in the examples that follow. In one embodiment, the composition further comprises the first target and the second target. The first targets and second targets may be any target suitable for an intended use. In non-limiting embodiments, the first target and/or
30 the second target may comprise polypeptides or nucleic acids. In another embodiment, the first target and/or the second target further comprise one or more effector polypeptide domains. Any effector polypeptide domains may be used as suitable for an intended use. In one embodiment, the one or more effector polypeptide domains may comprise a polypeptide including, but not limited to, nucleic acid binding proteins, transcription factors, receptor

binding proteins, split enzymes, effectors of membrane receptors, etc. In another embodiment, the first polypeptide and/or the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide having the amino acid sequence selected from the group consisting of SEQ ID NOS: 1-290 and 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, as listed in Tables 1A and 1B. In one embodiment, at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of the first polypeptide and/or the second polypeptide are invariant compared to the reference polypeptide amino acid sequence (interface residues shown in Table 2).

In one non-limiting and exemplary embodiment,

(a) the first polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one having the amino acid sequence of SEQ ID NO:42

1' (b)
 20 GSDDKELDKLLDTLEKILQTATKIIDANKLLEKLRRSERKDPKVVETVVELLKRHEKAVKELLEIAKTHAKKVE
 (SEQ ID NO: 42)

(b) the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one having the amino acid sequence of SEQ ID NO: 57.

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 25 DHSRKLEEI LDRLRKHVKRLLLEHLRELLSLVKENPEDKDLVEVLELSLAILRRSLEAVEAFLKSVTKKDPDDEDL
 RRKADEIRKEVEEIKKSLAEVEKEIYKLG (SEQ ID NO: 57)

In one non-limiting and exemplary embodiment,

(a) the first polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one having the amino acid sequence of SEQ ID NO:352

1' (b)

DDKELDKLLDTLEKILQATATKIIDDANKLLEKLRRSERKDPKVVETVVELLKRHEKAVKELLEIAKTHAKKVE
(SEQ ID NO: 352)

(b) the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to one
5 having the amino acid sequence of SEQ ID NO: 57.

10

DHSRKLEEILDRLRKHVKRLLEHLRELLSLVKENPEDKDLVEVLELSLAILRRSLEAVEAFLKSVTKKDPDDEDL
RRKADEIRKEVEEIKKSLAEVEKEIYKLK (SEQ ID NO: 57)

10 In another embodiment, the first target and/or the second target each comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide having the amino acid sequence selected from the group consisting of SEQ ID NO:1-290 and 331, 332, 334, 336-
15 422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, with the proviso that the first target forms a heterodimer (a-b) pair with the first polypeptide, and the second target forms a heterodimer (a-b) pair with the second polypeptide. Heterodimer A-B pairs among the polypeptides of SEQ ID NOS:1-290 and 331,
20 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494 are described at length above (See also FIG. 16). In one embodiment, at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined
25 interface positions of the first target and/or the second target are compared to the reference polypeptide amino acid sequence (interface residues shown in Table 2).

The compositions of this ninth aspect can be used for any suitable purpose, including in designing NAND logic gates. In one embodiment, the disclosure provides methods comprising

- 30 (a) contacting a biological sample with the composition of any one of claims 60-66; and
(b) detecting binding interactions between the first polypeptide and the first target, the second polypeptide and the second target, and the first target and the second target in the

sample, such as detecting an output signal caused by actions of effector polypeptides upon binding.

As used throughout the present application, the term "polypeptide" is used in its
5 broadest sense to refer to a sequence of subunit amino acids. The polypeptides of the
invention may comprise L-amino acids, D-amino acids (which are resistant to L-amino acid-
specific proteases *in vivo*), or a combination of D- and L-amino acids. The polypeptides
described herein may be chemically synthesized or recombinantly expressed. The
polypeptides may be linked to other compounds to promote an increased half-life *in vivo*,
10 such as by PEGylation, HESylation, PASylation, glycosylation, or may be produced as an
Fc-fusion or in deimmunized variants. Such linkage can be covalent or non-covalent as is
understood by those of skill in the art.

As will be understood by those of skill in the art, the polypeptides of the invention
may include additional residues at the N-terminus, C-terminus, or both that are not present in
15 the polypeptides of the invention; these additional residues are not included in determining
the percent identity of the polypeptides of the invention relative to the reference polypeptide.

As noted above, the polypeptides of the invention may include additional residues at
the N-terminus, C-terminus, or both. Such residues may be any residues suitable for an
intended use, including but not limited to detection tags (i.e.: fluorescent proteins, antibody
20 epitope tags, etc.), linkers, therapeutic agents, ligands suitable for purposes of purification
(His tags, etc.), ligands to drive localization, and peptide domains that add functionality to
the polypeptides.

In a tenth aspect, the disclosure provides nucleic acids encoding the polypeptide,
protein, fusion protein, scaffold, or design component of any embodiment or combination of
25 embodiments disclosed herein. The nucleic acid sequence may comprise single stranded or
double stranded RNA or DNA in genomic or cDNA form, or DNA-RNA hybrids, each of
which may include chemically or biochemically modified, non-natural, or derivatized
nucleotide bases. Such nucleic acid sequences may comprise additional sequences useful for
promoting expression and/or purification of the encoded polypeptide, including but not
30 limited to polyA sequences, modified Kozak sequences, and sequences encoding epitope
tags, export signals, and secretory signals, nuclear localization signals, and plasma membrane
localization signals. It will be apparent to those of skill in the art, based on the teachings
herein, what nucleic acid sequences will encode the polypeptides of the disclosure.

In an eleventh aspect, the disclosure provides expression vector comprising one or more nucleic acids of the disclosure operatively linked to a control sequence. "Expression vector" includes vectors that operatively link a nucleic acid coding region or gene to any control sequences capable of effecting expression of the gene product. "Control sequences" operably linked to the nucleic acid sequences of the disclosure are nucleic acid sequences capable of effecting the expression of the nucleic acid molecules. The control sequences need not be contiguous with the nucleic acid sequences, so long as they function to direct the expression thereof. Thus, for example, intervening untranslated yet transcribed sequences can be present between a promoter sequence and the nucleic acid sequences and the promoter sequence can still be considered "operably linked" to the coding sequence. Other such control sequences include, but are not limited to, polyadenylation signals, termination signals, and ribosome binding sites. Such expression vectors can be of any type, including but not limited plasmid and viral-based expression vectors. The control sequence used to drive expression of the disclosed nucleic acid sequences in a mammalian system may be constitutive (driven by any of a variety of promoters, including but not limited to, CMV, SV40, RSV, actin, EF) or inducible (driven by any of a number of inducible promoters including, but not limited to, tetracycline, ecdysone, steroid-responsive). The expression vector must be replicable in the host organisms either as an episome or by integration into host chromosomal DNA. In various embodiments, the expression vector may comprise a plasmid, viral-based vector, or any other suitable expression vector.

In a twelfth aspect, the disclosure provides cells comprising one or more nucleic acid, expression vector, polypeptide, protein, heterodimer protein, and/or protein scaffold of any embodiment or combination of embodiments disclosed herein. Nucleic acids or expression vectors may be episomal or chromosomally integrated. Any suitable cell type may be used, such prokaryotic or eukaryotic cells. The cells can be transiently or stably engineered to incorporate the expression vector of the disclosure, using techniques including but not limited to bacterial transformations, calcium phosphate co-precipitation, electroporation, or liposome mediated-, DEAE dextran mediated-, polycationic mediated-, or viral mediated transfection

In addition, the disclosure provides methods of producing a polypeptide, fusion protein, protein, heterodimer, etc. (collectively referred to as polypeptide) disclosed herein. In one embodiment, the method comprises the steps of (a) culturing a host according to this aspect of the disclosure under conditions conducive to the expression of the polypeptide, and (b) optionally, recovering the expressed polypeptide. The expressed polypeptide can be

recovered from the cell free extract or recovered from the culture medium. In another embodiment, the method comprises chemically synthesizing the polypeptides.

5 Example 1 Design of orthogonal protein heterodimers

Abstract: Here we demonstrate that heterodimeric interaction specificity can be achieved using extensive and modular buried hydrogen bond networks. We used the Crick generating equations to produce millions of four helix backbones with varying degrees of supercoiling around a central axis, identified those accommodating extensive hydrogen bond
10 networks, and designed connected pairs of helices with short loops and optimize the remainder of the sequence. 65 of 97 such designs expressed in *E. coli* formed constitutive heterodimers, and crystal structures of four designs were in close agreement with the computational models and confirmed the designed hydrogen bond networks. In cells, a set of six heterodimers were found to be fully orthogonal, and in vitro, following mixing of 32
15 chains from sixteen heterodimer designs, denaturation in 5M GdnHCl and reannealing, the vast majority of the interactions were between the designed cognate pairs. The ability to design orthogonal protein heterodimers enables sophisticated protein based control logic for synthetic biology, and illustrates that nature has not fully explored the possibilities for programmable biomolecular interaction modalities. Hydrogen bond networks, including
20 modular hydrogen bond networks are described in published patent application number WO2017173356, incorporated by reference herein.

Orthogonal sets of protein-protein and protein-peptide interactions play important roles in biological systems. Creation of new specificities by sequence redesign has been difficult, often resulting in promiscuous binding. We hypothesized that large sets of
25 designed heterodimers could be generated by incorporating asymmetric buried hydrogen bond networks into regularly repeating backbone structures. We generated helical bundle heterodimers in which each monomer is a helix-turn-helix starting from four-helix backbones. For each of the four helices, we exhaustively sampled the helical phase ($\Delta\phi_1$), supercoil radius (R) and offset along the Z-axis (Z offset) (FIG. 1A), restricting the supercoil
30 phases of the helices to 0, 90, 180 and 270 degrees, and the supercoil twist (ω_0) and helical twist (ω_1) to the ideal values for either a two layer left handed super coil ($\omega_0=-2.85$ and $\omega_1=102.85$), or a 5 layer untwisted bundle ($\omega_0=0$ and $\omega_1=100$) (Fig. 5A-B). This yielded 27 million untwisted and 60 million left-handed supercoiled backbones for both parallel and antiparallel orientations of opposing helices (Fig. 1B).

To identify the modular hydrogen bond network equivalents to DNA base pairs, we used ROSETTA™ HBNET²¹ to design buried hydrogen bond networks in the central repeat units of each backbone, and obtained 2251 hydrogen bond networks involving at least 4 side chain residues with all heavy-atom donors and acceptors participating in hydrogen bonds, and connecting all 4 helices (Fig. 1c; Fig. 6, Table 6). We then identified all of the geometrically compatible placements of these hydrogen bond networks in each backbone (Fig. 1d), selected backbones accommodating at least two networks, and connected pairs of helices with short loops (Fig. 1e). Low energy sequences were identified using ROSETTADesign™²² calculations in which the hydrogen bond networks were held fixed. Designs with fully satisfied hydrogen bond networks and tight hydrophobic packing were selected for experimental characterization, excluding those with networks with C2 symmetry to disfavor homodimerization of monomers. Designed heterodimers (DHDs) are referred to by numbers with monomers labeled a or b; for example, DHD15_a refers to monomer “a” of design DHD15.

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Table 6: The frequency of observing each hydrogen bond networks during the systematic search.

HBNet composition	Frequency	Percentage (%)
(S/T) 2Q1Y1	13954	3.87071362
(S/T) 3Q1	9959	2.76253669
(S/T) 2D1H1	8452	2.34450849
(S/T) 1Q2Y1	7603	2.10900356
(S/T) 1D1Q1Y1	7359	2.04132016
(S/T) 3D1	6332	1.75643963
(S/T) 3D1Q1	5525	1.53258512
(S/T) 1D1Q3	5071	1.40664962
(S/T) 1D1Q2	5062	1.4041531
(S/T) 2N1Y1	5046	1.39971484
(S/T) 1N1Q1Y1	4921	1.36504097
(S/T) 2H2	4683	1.29902192
(S/T) 2H1Q1	4572	1.26823152
(S/T) 3H1	3955	1.09708129
(S/T) 2D1Q2	3946	1.09458477
(S/T) 1D1N1Q2	3862	1.07128393
(S/T) 3N1	3783	1.04937005
(S/T) 2D1Y1	3762	1.04354484
(S/T) 2D1Q1	3669	1.01774747

(S/T) 1D1H1Q1	3653	1.01330922
(S/T) 1D1Q1W1	3409	0.94562582
(S/T) 1D1Q2Y1	3342	0.92704063
(S/T) 2Q3	3111	0.86296331
(S/T) 2D1N1Q1	2999	0.83189552
(S/T) 2Q2Y1	2850	0.79056427
(S/T) 1D1W1Y1	2849	0.79028688
(S/T) 2N1Q2	2741	0.76032865
(S/T) 2D1Q1Y1	2723	0.75533562
(S/T) 1D1N1Q1Y1	2684	0.74451737
(S/T) 2Q1W1	2641	0.73258956
(S/T) 2H1N1Q1	2591	0.71872001
(S/T) 2Q2	2582	0.71622349
(S/T) 2N1Q1	2554	0.70845654
(S/T) 2D1W1	2467	0.68432353
(S/T) 2H1N1	2377	0.65935834
(S/T) 4Q1	2305	0.63938619
(S/T) 1N1Q2Y1	2296	0.63688967
(S/T) 1D2Q2	2285	0.63383837
(S/T) 2D1H1Q1	2276	0.63134185
(S/T) 2D1Q1W1	2267	0.62884533
(S/T) 1H1Q1Y1	2222	0.61636274
(S/T) 1D1N1Q1	2207	0.61220187
(S/T) 2H1Y1	2150	0.59639059
(S/T) 1D1N1Y1	2109	0.58501756
(S/T) 1Q1Y2	1962	0.54424109
(S/T) 1H1Q2	1957	0.54285413
(S/T) 1Q1W1Y1	1954	0.54202196
(S/T) 2N1Q1Y1	1935	0.53675153
(S/T) 3H1Q1	1901	0.52732024
(S/T) 1D1H1W1	1879	0.52121764

94 of the 97 selected designs were well-expressed in *E. coli* with both monomers co-purifying by Ni-affinity chromatography (only one monomer contains a hexahistidine tag). For 85/94, the dominant species observed in size exclusion chromatography (SEC) had the expected size (Fig. 1f). Three designs characterized by CD spectroscopy were found to be all alpha helical and stable at 95°C (Fig. 1g, Fig. 6). Sequences and other information on the designs are provided in Tables 1A-B (above).

We explored the extent to which the heterodimer set could be expanded by permuting the hydrogen bond networks in the different helical repeat units, and by permuting the backbone connectivity. Assigning each unique network a letter, DHD37_XBBA indicates a

variant where the second, third and fourth repeat units have hydrogen bond networks B, B, and A, and the first heptad has exclusively hydrophobic residues in the core, while DHD103_1:423 indicates a heterodimer where one monomer consists of the first helix of DHD103 and the other monomer consists of helices 2 through 4 (Fig. 7). 13 of 14 hydrogen bond network permuted variants and 9 of 10 “3+1” backbone-permuted heterodimers (generated from five starting “2+2” heterodimers) ran as single peaks on SEC.

SAXS spectra collected for 44 designs were consistent with the design models (Fig. 1h, Fig. 2f-h). The X-ray crystal structures of DHD131, DHD37_1:234, DHD127 and DHD15 had backbone C α atom RMSDs to the design models ranging from 0.95 to 1.7 Å. The extensive five-residue buried hydrogen bond network of DHD131 (involving two serines, an asparagine, a tyrosine, and a tryptophan) is nearly identical in the crystal structure, with an additional water molecule bridging the interactions (Fig. 2a). The two designed hydrogen bond networks in DHD37_1:234, which contain buried histidine and tyrosine aromatic side chains sterically disfavoring homodimers, are in close agreement with the crystal structure (Fig. 2b). In DHD127, the histidines in the two hydrogen bond networks adopt a rotamer different from the design model (Fig. 2c), making a hydrogen bond with a water molecule. A crystal structure of DHD15 at pH 7.0 is similar to the design model (Fig. 2d), while a structure at pH 6.5 is of a domain-swapped, hetero-tetramer conformation.

We built three induced dimerization systems by fusing one monomer each from two different heterodimers via a flexible linker, and testing whether the remaining two monomers from each pair could be brought together by the fusion (Fig. 3a). In each case, the three components co-purified by Ni-NTA chromatography (one monomer has a hexahistidine tag); In yeast two-hybrid assays (Y2H) with monomers from two different heterodimers fused to the DNA binding domain (DBD) and transcriptional activation domain (AD), expression of the heterodimerizer fusion as a separate polypeptide chain increased signal significantly over background (Fig. 3b).

We covalently linked the monomer chain “a” subunits of 3 DHDs via flexible linkers (FIG. 3C), and co-expressed this “scaffold” and the 3 separate chain “b” monomers, one with a hexahistidine tag, in *E. coli*. The scaffold plus monomer assembly is stable at 95°C and has a guanidine denaturation midpoint of 4 M (FIG. 9).

By generating interfaces with many polar groups which are energetically costly to bury without geometrically matched hydrogen bonding interactions, our design protocol implicitly disfavors non-cognate interactions (explicit negative design to disfavor non-cognate interactions is computationally intractable given the very large number of possible

off-target binding modes). For 24 designs, strong interactions were observed by Y2H with the two partners fused to DBD and AD, but not when either partner was fused to both domains; the designed heterodimers, but not the homodimers, form in cells (Fig. 4A). The 24 monomers in 12 of these designs were crossed in an all-by-all Y2H experiment; interactions were observed for all cognate pairs, and 27 of the 552 possible non-cognate interactions (Fig. 9). Orthogonality was higher for an 8 DHD subset: of 240 possible non-cognate interactions, only 4 were observed (Fig. 4B; the interacting polar residues are depicted schematically in Fig. 10). Co-expression of unfused monomers eliminated off-target interactions (Fig. 4C); the cognate interactions are evidently stronger than the non-cognate interactions.

Our results demonstrate that the domain of unbounded sets of orthogonal heterodimeric biomolecules constructed from a single repeating backbone is not limited to nucleic acids. Interaction specificity arises from extensive buried hydrogen bond networks such as the fully connected TYR-SER-TRP-ASN-SER (SEQ ID NO:333) crystallographically confirmed network in Fig. 2a, and heterogeneity in the size of the residues at the designed interface (Fig. 9d-i), analogous to the contribution of steric effects to Watson-Crick base pairing specificity. Our large set of orthogonal interactions, together with the retention of specificity in the fused monomer systems (the induced dimerizer and scaffold of Fig. 3), and the interaction strength hierarchy illustrated by the cognate interaction competition experiment (Fig. 4c), can be used, by way of non-limiting example, to prepare protein based cellular control circuits with faster response times and better integration with signaling inputs and outputs than current nucleic acid based circuitry.

Methods for Example 1

Computational Design

1. Systematic sampling of parametric helical backbones

We used a generalization of the Crick coiled-coil parameters⁵ to independently sample all four helices of the heterodimers supercoiled around the same axis. The supercoil twist (ω_0) and helical twist (ω_1) were coupled and ideal values were used²⁰ with ω_0 and ω_1 held constant among the helices. A left-handed supercoil results from $\omega_0=-2.85$ and $\omega_1=102.85$, and a straight bundle with no supercoiling from $\omega_0=0$ and $\omega_1=100$. The supercoil phases ($\Delta\phi_0$) for the helices were fixed at 0° , 90° , 180° and 270° , respectively. The offset along the Z-axis (Z offset) for the first helix was fixed to 0 as a reference point, with the rest of the helices independently sampling from -1.51 \AA to 1.51 \AA , with a step size of 1.51 \AA . All helices sampled helical phases ($\Delta\phi_1$) independently, from 0° to 90° , with a step size of 10° .

Two of the helices with a $\Delta\phi_0$ separation of 180° sampled the radius from Z-axis (R) from 5 Å to 8 Å, while the other two sampled from 7 Å to 10 Å, all with a step size of 1 Å. Each helix is set to have 35 residues to accommodate 5 heptad repeats. After removing redundant sample points from the overlapping regions of radii sampling, the supercoiled helical bundles

5 contained more than 60 million unique backbones, and the straight helical bundles contained more than 27 million unique backbones.

2. HBNet Search

For each parametrically generated backbone, HBNetTM 21 was used to search the middle heptad for hydrogen bond networks that connect all four helices, contain at least four
10 side chains contributing hydrogen bonds, have all heavy atom donors and acceptors satisfied, and span the intermolecular interface. Symmetry was not enforced during the HBNetTM search. For buried interface positions, only non-charged polar amino acids were considered; for residues that were at the boundary between protein core and surface, all polar amino acids were considered. A subsequent RosettaTM design calculation was performed to optimize
15 hydrophobic packing, with atom pair restraints from HBNetTM being put on the newly identified hydrogen bond networks. Finally, a minimization step and side chain repacking step was performed without atom pair restraints on hydrogen bonding residues to evaluate how well the networks remained intact in the absence of the constraints. Designs with at most 5 alanines in the middle heptad and no buried unsatisfied polar heavy atoms were selected for
20 downstream design.

3. Generating combinations of HBNetTM with heptad stacking

The purpose of this step is to identify five-heptad backbones (full backbones) that can accommodate at least 2 HBNetTM. Instead of generating one-heptad backbones and full
25 backbones separately, searching for HBNetTM in the one-heptad backbones and aligning them to all full backbones, we reasoned the heptad stacking method remains the same if one simply searches for HBNetTM in the middle heptad on all full backbones, extracts the middle heptads, and aligns them to all full backbones. We therefore extracted the middle heptads containing HBNetTM, generated all variants of chain ordering, and did pairwise alignment of
30 middle heptads to full backbones using TMalign³⁰. All alignments with root mean square deviation (RMSD) less than 0.3 were identified and full backbones that can accommodate at least 2 middle heptads were selected for final design.

4. Connecting parametric helical backbones

Helical backbones are connected with short 2-5 residue loops such that the RMSD of each loop is less than 0.4 RMSD to a nine residues stretch in a native protein. Distance and directionality between helices limit what loops can connect, as such, our closure extends and shrinks helices by up to 3 residues. We then superimpose all short loops from the PDB onto the first and last two helical residues. The loops with the lowest stub-RMSD are minimized using the RosettaTM score function onto the helical endpoints to ensure a near perfect closure. Loop quality is assessed by measuring the distance in RMSD to the closest nine stretch in the PDB. The loop with the lowest RMSD is returned as the solution. We repeat this procedure to connect all helices and report the solution with the lowest RMSD.

10

5. Design calculations

Backbones were regularized using Cartesian space minimization in RosettaTM to alleviate any torsional strain introduced by heptad stacking. Two consecutive RosettaTM packing rounds were performed with increasing weight on the repulsive energy to optimize hydrophobic packing, while constraining the hydrogen bond network residues. A FastDesign step was subsequently used within a generic Monte Carlo mover to optimize secondary structure shape complementarity, while allowing at most 8% alanine, 3 methionine and 3 phenylalanine in the protein core. The last step of minimization and side chain repacking to identify the movement of HBNets without atom pair constraints is the same as what was described in Step 2.

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6. Selection criteria and metrics used to evaluate designs

Designs were selected based on the following criteria: change in polar surface area upon binding (dSASA_polar) greater than 800 Å²; secondary structure shape complementarity (ss_sc) score greater than 0.65; holes score around HBNets less than -1.4; no buried unsatisfied heavy atoms; at least one buried bulky polar side chains per monomer. Selected designs were then visually inspected for good packing of hydrophobic side chains, especially the interdigitation of isoleucine, leucine and valine. Surface tyrosines were added at non-interfering positions to aid protein concentration measurement by recording OD280. Surface charge residues for a few of the designs were redesigned to shift the theoretical isoelectric point away from buffer pH.

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RMSD calculations

Crystal structures and the corresponding design models were superimposed with TMalign using all heavy atoms. From this alignment, RMSD was calculated across all alpha-carbon atoms, and also across heavy atoms of the hydrogen bond network residues.

5 Logistic Regression

Designs were first scored with various filters in RosettaTM with the filter values reported. Experimental results and RosettaTM filter values were used as input to a logistic regression method³¹ to find correlations between computational metrics and experimental observations.

10

Visualization and figures

All structural images for figures were generated using PyMOL³².

Buffer and media recipe

15 TBM-5052: 1.2% [wt/vol] tryptone, 2.4% [wt/vol] yeast extract, 0.5% [wt/vol] glycerol, 0.05% [wt/vol] D-glucose, 0.2% [wt/vol] D-lactose, 25 mM Na₂HPO₄, 25 mM KH₂PO₄, 50 mM NH₄Cl, 5 mM Na₂SO₄, 2 mM MgSO₄, 10 μM FeCl₃, 4 μM CaCl₂, 2 μM MnCl₂, 2 μM ZnSO₄, 400 nM CoCl₂, 400 nM NiCl₂, 400 nM CuCl₂, 400 nM Na₂MoO₄, 400 nM Na₂SeO₃, 400 nM H₃BO₃

20 Lysis buffer: 20 mM Tris, 300 mM NaCl, 20 mM Imidazole, pH 8.0 at room temperature

Wash buffer: 20 mM Tris, 300mM NaCl, 30 mM Imidazole, pH 8.0 at room temperature

25 Elution buffer: 20 mM Tris, 300 mM NaCl, 250 mM Imidazole, pH 8.0 at room temperature

Buffer W: 100 mM Tris-HCl pH 8.0, 150 mM NaCl and 1 mM EDTA

Buffer E: Buffer W containing 2.5 mM D-desthiobiotin

TBS buffer: 20 mM Tris pH 8.0, 100 mM NaCl

30 Construction of synthetic genes

For the expression of heterodimers, both monomers were encoded in the same plasmid, separated by a ribosome binding sequence (GAAGGAGATATCATC; SEQ ID NO:327). Synthetic genes were ordered from Genscript Inc. (Piscataway, N.J., USA) and delivered in pET21-NESG *E. coli* expression vector, inserted between the NdeI and XhoI

sites. For the pET21-NESG constructs, a hexahistidine tag and a tobacco etch virus (TEV) protease cleavage site (GSSHHHHHHSSGENLYFQGS; SEQ ID NO:328) were added in frame at the N-terminus of the second monomer. A stop codon was introduced at the 3' end of the second monomer to stop expression of the C-terminal hexahistidine tag in the vector.

- 5 For purification with Strep-tactin resin, a streptavidin tag (SAWSHPQFEKGGGSGGGSGGSAWSHPQFEKSGENLYFQGS; SEQ ID NO:329) coding sequence was cloned in frame 5' of the first monomer sequence.

For the co-expression of 3 and 4 proteins from the same plasmid (induced dimerization and synthetic scaffold designs), synthetic genes were cloned in the pRSFDuet-1
10 expression vector. The first (in the case of 3 proteins) or first two (in the case of 4 proteins) genes were cloned between NcoI and HindIII sites, with a ribosome binding site separating the 2 proteins in the latter case. The last two genes were cloned between NdeI and XhoI sites, separated by a ribosome binding site. A hexahistidine tag and a TEV protease cleavage site coding sequence were cloned in frame 5' of the last gene.

- 15 Genes for yeast-two-hybrid (Y2H) studies were cloned into plasmids bearing the GAL4 transcription activation domain (poAD) and the GAL4 DNA-binding domain (poDBD).

Protein expression

- 20 Plasmids were transformed into chemically competent *E. coli* expression strains BL21(DE3)Star (Invitrogen) or Lemo21TM (DE3) (New England Biolabs) for protein expression. Single colonies were picked from agar plates following transformation and growth overnight, and 5 ml starter cultures were grown at 37°C in Luria-Bertani (LB) medium containing 100 µg/mL carbenicillin (for pET21-NESG vectors) or kanamycin (for
25 pRSFDuet-1 vectors) with shaking at 225 rpm for 18 hours at 37°C. Starter cultures were diluted into 500 ml TBM-5052 containing 100 µg/mL carbenicillin or kanamycin, and incubated with shaking at 225 rpm for 24 hours at 37°C.

- For expression of ¹³C¹⁵N- or ¹⁵N-labeled protein, the plasmids were transformed into the Lemo21TM (DE3) *E. coli* expression strain and plated on M9/glucose plates containing 50
30 µg/mL carbenicillin. For the starter culture, a single colony was used for inoculation of 50 mL LB medium with 50 µg/mL carbenicillin in a 250 mL baffled flask, and incubated with shaking at 225 rpm for 18 hours at 37°C. 10 mL of the starter culture was then transferred to a 2 L baffled flask containing 500 mL of Terrific BrothTM (Difco), with 25 mM Na₂HPO₄, 25 mM KH₂PO₄, 50 mM NH₄Cl, 5 mM Na₂SO₄, and 100 µg/mL carbenicillin. The culture

was grown at 37°C to an OD600 of approximately 1.0, then centrifuged at 5000 ref for 15 minutes to pellet the cells. The Terrific Broth™ medium was removed, and the cells were washed briefly with 30 mL of phosphate buffered saline (PBS). The cells were then transferred to a fresh 2 L baffled flask containing 500 mL of labeled media (25 mM Na₂HPO₄, 25 mM KH₂PO₄, 50 mM 15NH₄Cl, 5 mM Na₂SO₄, 0.2% (w/v) ¹³C glucose),
5 and 100 µg/mL carbenicillin. The cells were allowed to grow at 37°C for 2 hours, before IPTG (Carbosynth) was added to 1mM and the temperature was reduced to 18°C. The labeled glucose and NH₄Cl were obtained from Cambridge Isotopes.

10 Affinity purification

Cells were harvested by centrifugation for 15 minutes at 5000 ref at 4°C and resuspended in 20 ml lysis buffer. Lysozyme, DNase, and EDTA-free cocktail protease inhibitor (Roche) were added to the resuspended cell pellet before sonication at 70% power for 5 minutes. For Immobilized metal affinity chromatography (IMAC), lysates were clarified
15 by centrifugation at 4°C and 18,000 rpm for at least 30 minutes and applied to Ni-NTA (Qiagen) columns pre-equilibrated with lysis buffer. The column was washed two times with 5 column volumes (CV) of wash buffer, followed by 5 CV of elution buffer. For Strep tag purification, elution fractions from IMAC were applied to Strep-Tactin® Superflow resin (IBA) pre-equilibrated in Buffer W. The column was washed with 5 CV Buffer W, before
20 applying 3 CV Buffer E to elute proteins off the column. Mass and purity of eluted proteins were confirmed using electrospray ionization mass spectrometry (ESI-MS) on a Thermo Scientific TSQ Quantum Access mass spectrometer.

Size-exclusion chromatography (SEC)

25 N-terminal hexahistidine tags and streptavidin tags were cleaved with TEV protease overnight at room temperature, at a ratio of 1 mg TEV for 100 mg of protein. Prior to addition of TEV, buffer was exchanged into lysis buffer. After TEV cleavage, sample was passed over an additional Ni-NTA column and washed with 1.5 CV of lysis buffer, flow through were collected and further purified by SEC using a Superdex™ 75 10/300 increase
30 column (GE Healthcare) in TBS buffer.

Circular dichroism (CD) measurements

CD wavelength scans (260 to 195 nm) and temperature melts (25 to 95°C) were performed using an AVIV model 420 CD spectrometer. Temperature melts were carried out

at a heating rate of 4°C/min and monitored by the change in ellipticity at 222 nm; protein samples were diluted to 0.25 mg/mL in PBS pH 7.4 in a 0.1 cm cuvette. Guanidinium chloride (GdmCl) titrations were performed on the same spectrometer with automated titration apparatus in PBS pH 7.4 at 25°C, with a protein concentration of 0.025 mg/mL in a 1
5 cm cuvette with stir bar. Each titration consisted of at least 40 evenly distributed GdmCl concentration points with one minute mixing time for each step. Titrant solution consisted of the same concentration of protein in PBS + GdmCl.

Crystallization of protein samples

10 Purified protein samples were concentrated to approximately 20 mg/ml in 25 mM Tris pH 8.0 and 150 mM NaCl. Samples were screened with a 5-position deck MosquitoTM crystal (ttplabtech) with an active humidity chamber, utilizing the following crystallization screens: JCSG+TM (Qiagen), Crystal ScreenTM (Hampton Research), PEG/IonTM (Hampton Research), PEGRx HTTM (Hampton Research), IndexTM (Hampton Research) and MorpheusTM
15 (Molecular Dimensions). The optimal conditions for crystallization of the different designs were found as follows: OPHD_37_N3C1, 0.15 M potassium bromide and 30% w/v polyethylene glycol monomethyl ether 2000; OPHD_127, 0.12 M ethylene glycols, 0.1 M buffer system 3 pH 8.5, and 50% v/v precipitate mix 1 from the Morpheus screen; OPHD_15, 0.2 M Ammonium sulfate, 0.1 M BIS-TRIS pH 6.5, 18% v/v Polyethylene glycol 400;
20 OPHD_15, 0.1 M Imidazole pH 7.0, and 25% v/v Polyethylene glycol monomethyl ether 550; OPHD_131, 0.2 M Ammonium acetate, 0.1 M HEPES pH 7.5, 25% w/v Polyethylene glycol 3,350. Crystals were obtained after 1 to 14 days by the hanging drop vapor diffusion method with the drops consisting of a 1:1, 2:1 and 1:2 mixture of protein solution and reservoir solution.

25

X-ray data collection and structure determination

The crystals of the designed proteins were looped and placed in the corresponding reservoir solution, containing 20% (v/v) glycerol if the reservoir solution did not contain cryoprotectant, and flash-frozen in liquid nitrogen. The X-ray data sets were collected at the
30 Advanced Light Source at Lawrence Berkeley National Laboratory with beamlines 8.2.1 and 8.2.2. Data sets were indexed and scaled using either XDS³³ or HKL2000³⁴. Initial models were generated by the molecular-replacement method with the program PHASER³⁵ within the PhenixTM software suite³⁶, using the design models as the initial search models. Efforts were made to reduce model bias through refinement with simulated annealing using

Phenix.refine^{TM 37}, or, if the resolution was sufficient, by using Phenix.autobuild^{TM 38} with rebuild-in-place set to false, simulated annealing and prime-and-switch phasing. Iterative rounds of manual building in COOT³⁹ and refinement in PhenixTM were used to produce the final models. Due to the high degree of self-similarity inherent in coiled-coil-like proteins, datasets for the reported structures suffered from a high degree of pseudo translational non-crystallographic symmetry, as report by Phenix.XtrriageTM, which complicated structure refinement and may explain the higher than expected R values reported. RMSDs of bond lengths, angles and dihedrals from ideal geometries were calculated with Phenix^{TM 36}. The overall quality of all final models was assessed using the program MOLPROBITY^{TM 40}.

10

Small Angle X-ray Scattering (SAXS)

Samples were purified by SEC in 25 mM Tris pH 8.0, 150 mM NaCl and 2% glycerol; fractions preceding the void volume of the column were used as blanks for buffer subtraction. Scattering measurements were performed at the SIBYLS^{TM 12.3.1} beamline at the Advanced Light Source. The X-ray wavelength (λ) was 1.27 Å, and the sample-to-detector distance was 1.5 m, corresponding to a scattering vector q ($q = 4\pi \sin \theta/\lambda$, where 2θ is the scattering angle) range of 0.01 to 0.3 Å⁻¹. A series of exposures, in equal sub-second time slices, were taken of each well: 0.3 second exposures for 10 seconds resulting in 32 frames per sample. For each sample, data was collected for two different concentrations to test for concentration-dependent effects; “low” concentration samples ranged from 2-3 mg/mL and “high” concentration samples ranged from 5-7 mg/mL. Data was processed using the SAXS FrameSliceTM online serve and analyzed using the ScÅtterTM software package^{41,42}. FoXS^{TM 43,44} was used to compare design models to experimental scattering profiles and calculate quality of fit (χ) values.

25

Yeast two-hybrid assay

For each pair of binders tested, chemically competent cells of yeast strain PJ69-4a (MATa trp1-901 leu2-3,112 ura3-52 his3-200 gal4(deleted) gal80(deleted) LYS2::GAL1-HIS3 GAL2-ADE2 met2::GAL7-lacZ) were transformed with the appropriate pair of plasmids containing DNA binding domain or activation domains, using the LiAc/SS carrier DNA/PEG method⁴⁵. In the case of induced dimerization, the heterodimerizer was cloned downstream of one of the “monomer proteins”, separated by a p2a and nuclear localization sequence (GSGATNFSLLKQAGDVEENPGPGDKAELIPEPPKRRKVELGTA; SEQ ID NO:330). The p2a sequence ensures translational cleavage to make the heterodimerizer a

30

separate protein from the “monomer protein”. The selection of transformed yeast cells was performed in synthetic dropout (SDO) media lacking tryptophan and leucine for 48 hours with shaking at 1000 rpm at 30°C. The resulting culture was diluted 1:100 and grown for 16 hours in fresh SDO media lacking tryptophan and leucine, before transferring to a 96 well plate and diluted 1:100 into SDO media containing 100 mM 3-Amino-1,2,4-triazole (3-AT), lacking tryptophan, leucine and histidine (5 mM 3-AT in the case of induced dimerization). The culture was incubated with shaking at 1000 rpm at 30°C. Since bringing the DNA binding domain and the transcription activation domain into proximity is necessary for the growth of yeast cells in media lacking histidine, binding of two proteins was indicated by the growth of yeast cells^{46,47}. The optical density of yeast cells was recorded after 48 hours. For Y2H assay on agar plates, the 1:100 diluted overnight culture was transferred onto Nunc™ OmniTray™ (Thermo Fisher) using a 96 Solid Pin Multi-Blot Replicator (V&P Scientific), with the agar lacking tryptophan, leucine and histidine, and containing 100 mM 3-AT. The plates were imaged daily until Day 5 to monitor the sizes of colonies. Images were analyzed by the ColonyArea⁴⁸ package on ImageJ.

References for Example 1

1. Jones, S. & Thornton, J. M. Principles of protein-protein interactions. *Proc. Natl. Acad. Sci. U. S. A.* **93**, 13–20 (1996).
2. Harbury, P. B., Zhang, T., Kim, P. S. & Alber, T. A switch between two-, three-, and four-stranded coiled coils in GCN4 leucine zipper mutants. *Science* **262**, 1401–1407 (1993).
3. Diss, M. L. & Kennan, A. J. Orthogonal recognition in dimeric coiled coils via buried polar-group modulation. *J. Am. Chem. Soc.* **130**, 1321–1327 (2008).
4. Thomas, F., Boyle, A. L., Burton, A. J. & Woolfson, D. N. A set of de novo designed parallel heterodimeric coiled coils with quantified dissociation constants in the micromolar to sub-nanomolar regime. *J. Am. Chem. Soc.* **135**, 5161–5166 (2013).
5. Crick, F. H. C. The Fourier transform of a coiled-coil. *Acta Cryst (1953)*. *Q6*, 685–689 [doi:10.1107/S0365110X53001952] **6**, 1–5 (1953).
6. Zarrinpar, A., Park, S.-H. & Lim, W. A. Optimization of specificity in a cellular protein interaction network by negative selection. *Nature* **426**, 676–680 (2003).
7. Aakre, C. D. *et al.* Evolving new protein-protein interaction specificity through promiscuous intermediates. *Cell* **163**, 594–606 (2015).

8. Joachimiak, L. A., Kortemme, T., Stoddard, B. L. & Baker, D. Computational design of a new hydrogen bond network and at least a 300-fold specificity switch at a protein-protein interface. *J. Mol. Biol.* **361**, 195–208 (2006).
9. Skerker, J. M. *et al.* Rewiring the specificity of two-component signal transduction systems. *Cell* **133**, 1043–1054 (2008).
- 5 10. Crooks, R. O., Baxter, D., Panek, A. S., Lubben, A. T. & Mason, J. M. Deriving Heterospecific Self-Assembling Protein-Protein Interactions Using a Computational Interactome Screen. *J. Mol. Biol.* **428**, 385–398 (2016).
11. Gradišar, H. & Jerala, R. De novo design of orthogonal peptide pairs forming parallel
10 coiled-coil heterodimers. *J. Pept. Sci.* **17**, 100–106 (2011).
12. Thompson, K. E., Bashor, C. J., Lim, W. A. & Keating, A. E. SYNZIP protein interaction toolbox: in vitro and in vivo specifications of heterospecific coiled-coil interaction domains. *ACS Synth. Biol.* **1**, 118–129 (2012).
13. Reinke, A. W., Grant, R. A. & Keating, A. E. A synthetic coiled-coil interactome
15 provides heterospecific modules for molecular engineering. *J. Am. Chem. Soc.* **132**, 6025–6031 (2010).
14. Acharya, A., Rishi, V. & Vinson, C. Stability of 100 homo and heterotypic coiled-coil a¹ a² pairs for ten amino acids (A, L, I, V, N, K, S, T, E, and R). *Biochemistry* **45**, 11324–11332 (2006).
- 20 15. Grigoryan, G. & Keating, A. E. Structure-based prediction of bZIP partnering specificity. *J. Mol. Biol.* **355**, 1125–1142 (2006).
16. Gonzalez, L., Jr, Woolfson, D. N. & Alber, T. Buried polar residues and structural specificity in the GCN4 leucine zipper. *Nat. Struct. Biol.* **3**, 1011–1018 (1996).
17. Lumb, K. J. & Kim, P. S. A buried polar interaction imparts structural uniqueness in a
25 designed heterodimeric coiled coil. *Biochemistry* **34**, 8642–8648 (1995).
18. Tatko, C. D., Nanda, V., Lear, J. D. & DeGrado, W. F. Polar Networks Control Oligomeric Assembly in Membranes. *J. Am. Chem. Soc.* **128**, 4170–4171 (2006).
19. Grigoryan, G. & DeGrado, W. F. Probing Designability via a Generalized Model of Helical Bundle Geometry. *J. Mol. Biol.* **405**, 1079–1100 (2011).
- 30 20. Huang, P.-S. *et al.* High thermodynamic stability of parametrically designed helical bundles. *Science* **346**, 481–485 (2014).
21. Boyken, S. E. *et al.* De novo design of protein homo-oligomers with modular hydrogen-bond network-mediated specificity. *Science* **352**, 680–687 (2016).
22. Leaver-Fay, A. *et al.* ROSETTA^{TM3}: an object-oriented software suite for the simulation

- and design of macromolecules. *Methods Enzymol.* **487**, 545–574 (2011).
23. Ruotolo, B. T. & Robinson, C. V. Aspects of native proteins are retained in vacuum. *Curr. Opin. Chem. Biol.* **10**, 402–408 (2006).
24. Sahasrabudde, A. *et al.* Confirmation of intersubunit connectivity and topology of
5 designed protein complexes by native MS. *Proc. Natl. Acad. Sci. U. S. A.* **115**, 1268–
1273 (2018).
25. Zhou, M., Huang, C. & Wysocki, V. H. Surface-induced dissociation of ion mobility-
separated noncovalent complexes in a quadrupole/time-of-flight mass spectrometer.
Anal. Chem. **84**, 6016–6023 (2012).
- 10 26. Zhou, M. & Wysocki, V. H. Surface induced dissociation: dissecting noncovalent protein
complexes in the gas phase. *Acc. Chem. Res.* **47**, 1010–1018 (2014).
27. Anderson, G. P., Shriver-Lake, L. C., Liu, J. L. & Goldman, E. R. Orthogonal Synthetic
Zippers as Protein Scaffolds. *ACS Omega* **3**, 4810–4815 (2018).
28. Rothmund, P. W. K. Folding DNA to create nanoscale shapes and patterns. *Nature* **440**,
15 297–302 (2006).
29. Qian, L. & Winfree, E. Scaling up digital circuit computation with DNA strand
displacement cascades. *Science* **332**, 1196–1201 (2011).
30. Zhang, Y. & Skolnick, J. TM-align: a protein structure alignment algorithm based on the
TM-score. *Nucleic Acids Res.* **33**, 2302–2309 (2005).
- 20 31. Rocklin, G. J. *et al.* Global analysis of protein folding using massively parallel design,
synthesis, and testing. *Science* **357**, 168–175 (2017).
32. Schrödinger, LLC. The PyMOL Molecular Graphics System, Version 1.8. (2015).
33. Kabsch, W. XDS. *Acta Crystallogr. D Biol. Crystallogr.* **66**, 125–132 (2010).
34. Otwinowski, Z. & Minor, W. Processing of X-ray diffraction data collected in oscillation
25 mode. *Methods Enzymol.* **276**, 307–326 (1997).
35. McCoy, A. J. *et al.* Phaser crystallographic software. *J. Appl. Crystallogr.* **40**, 658–674
(2007).
36. Adams, P. D. *et al.* PHENIX: a comprehensive Python-based system for macromolecular
structure solution. *Acta Crystallogr. D Biol. Crystallogr.* **66**, 213–221 (2010).
- 30 37. Afonine, P. V. *et al.* Joint X-ray and neutron refinement with phenix.refine. *Acta*
Crystallogr. D Biol. Crystallogr. **66**, 1153–1163 (2010).
38. Terwilliger, T. C. *et al.* Iterative model building, structure refinement and density
modification with the PHENIX AutoBuild wizard. *Acta Crystallogr. D Biol. Crystallogr.*
64, 61–69 (2008).

39. Emsley, P. & Cowtan, K. Coot: model-building tools for molecular graphics. *Acta Crystallogr. D Biol. Crystallogr.* 60, 2126–2132 (2004).
40. Davis, I. W. et al. MolProbity: all-atom contacts and structure validation for proteins and nucleic acids. *Nucleic Acids Res.* 35, W375–83 (2007).
- 5 41. Dyer, K. N. et al. High-throughput SAXS for the characterization of biomolecules in solution: a practical approach. *Methods Mol. Biol.* 1091, 245–258 (2014).
42. Rambo, R. P. & Tainer, J. A. Characterizing flexible and intrinsically unstructured biological macromolecules by SAS using the Porod-Debye law. *Biopolymers* 95, 559–571 (2011).
- 10 43. Schneidman-Duhovny, D., Hammel, M. & Sali, A. FoXS: a web server for rapid computation and fitting of SAXS profiles. *Nucleic Acids Res.* 38, W540–4 (2010).
44. Schneidman-Duhovny, D., Hammel, M., Tainer, J. A. & Sali, A. Accurate SAXS profile computation and its assessment by contrast variation experiments. *Biophys. J.* 105, 962–974 (2013).
- 15 45. Schiestl, R. H. & Gietz, R. D. High efficiency transformation of intact yeast cells using single stranded nucleic acids as a carrier. *Curr. Genet.* 16, 339–346 (1989).
46. Chien, C. T., Bartel, P. L., Sternglanz, R. & Fields, S. The two-hybrid system: a method to identify and clone genes for proteins that interact with a protein of interest. *Proc. Natl. Acad. Sci. U. S. A.* 88, 9578–9582 (1991).
- 20 47. Bartel, P. L., Roecklein, J. A., SenGupta, D. & Fields, S. A protein linkage map of *Escherichia coli* bacteriophage T7. *Nat. Genet.* 12, 72–77 (1996).
48. Guzmán, C., Bagga, M., Kaur, A., Westermarck, J. & Abankwa, D. ColonyArea: an ImageJ plugin to automatically quantify colony formation in clonogenic assays. *PLoS One* 9, e92444 (2014).
- 25 49. Dyachenko, A. et al. Tandem Native Mass-Spectrometry on Antibody-Drug Conjugates and Submillion Da Antibody-Antigen Protein Assemblies on an Orbitrap EMR Equipped with a High-Mass Quadrupole Mass Selector. *Anal. Chem.* 87, 6095–6102 (2015).
50. Waitt, G. M., Xu, R., Wisely, G. B. & Williams, J. D. Automated in-line gel filtration for native state mass spectrometry. *J. Am. Soc. Mass Spectrom.* 19, 239–245 (2008).
- 30 51. Marty, M. T. et al. Bayesian deconvolution of mass and ion mobility spectra: from binary interactions to polydisperse ensembles. *Anal. Chem.* 87, 4370–4376 (2015).
52. Bern, M. et al. Parsimonious Charge Deconvolution for Native Mass Spectrometry. *J. Proteome Res.* 17, 1216–1226 (2018).
53. Jones, D. T. Protein secondary structure prediction based on position-specific scoring

matrices. *J. Mol. Biol.* **292**, 195–202 (1999).

Example 2. Orthogonal protein heterodimers for designing modular protein logic gates

5 **Abstract:** The *de novo* design of modular protein logic for regulating protein function at the post-transcriptional level is a challenge for computational protein design and could have wide ranging applications in synthetic biology. Here we describe the design of 2-input AND, OR, NAND, NOR, XNOR, and NOT gates built from *de novo* designed proteins that regulate the association of arbitrary protein units ranging from split enzymes to transcriptional machinery
10 in vitro, and in living cells. Binding interaction cooperativity makes the gates largely insensitive to stoichiometric imbalances in the inputs, and the modularity of the approach enables ready extension to 3-input OR, AND, and disjunctive normal form gates. The modularity and cooperativity of the control elements, coupled with the ability to *de novo* design an essentially unlimited number of protein components, should enable design of
15 sophisticated post-translational control logic over a wide range of biological functions.

Introduction

The ability to *de novo* design protein-based logic gates with modular control of arbitrary protein-protein interactions could open the door to the tunable design of novel bio-
20 orthogonal functionalities.

In principle, it should be possible to design a wide range of logic gates *de novo* using a set of orthogonal heterodimeric molecules. For example, given hypothetical heterodimer pairs $A:A'$, $B:B'$, and $C:C'$, an AND gate modulating the association of A with C' can be constructed by genetically fusing A' and B , and B' and C : association occurs only in the
25 presence of both $A'-B$, and $B'-C$ (here and below “:” denotes noncovalent interaction, and “-”, genetic fusion via flexible linkers). Several building block properties are desirable for constructing such associative logic gates. First, there should be many mutually orthogonal heterodimeric pairs, so that gate complexity is not limited by the number of individual elements. Second, the building blocks should be modular and similar in structure so that
30 differences in building block shape and other properties do not have to be considered when constructing the gates. Third, single building blocks should be able to bind to multiple partners with different, tunable affinities, allowing inputs to perform negation operations by disrupting pre-existing lower affinity interactions. Fourth, the interactions should be cooperative so gate activation is not sensitive to stoichiometric imbalances in the inputs. In

the above AND gate, for example, if the interactions are not cooperative, a large excess of $A'-B$ will pull the equilibrium towards partially assembled complexes ($A'-B$ with either A or $B'-C$ but not both), which will disrupt gate activation.

Here, we explored the possibility of designing logic gates satisfying all four of the
5 above criteria using *de novo* designed protein heterodimers with hydrogen bond network-
mediated specificity (34). Sets of 6 (in vivo) and 15 (in vitro) mutually orthogonal designed
heterodimers (DHDs, hereafter referred to by numbers, e.g. I and I' form one cognate pair.
with hydrogen bond network (see Fig. 11A inset for example) mediated specificity are
available for logic gate construction, satisfying condition 1 (orthogonality). The
10 heterodimeric interfaces all share the same four helix bundle topology (Fig. 11A), satisfying
condition 2 (modularity). The shared interaction interface allows a limited amount of cross
talk between pairs, leading to a hierarchy of binding affinities, satisfying condition 3
(multiple binding specificities). Inspired by cooperatively activatable systems in nature (35,
36), we sought to achieve condition 4 (cooperativity) by constructing the monomer fusions
15 ($A'-B$ and $B'-C$ in the above example) in such a way that the interaction surfaces (with A and
 C') are buried within the fusions. The free energy required to expose these buried interfaces
would oppose gate activation, and we reasoned that the system could be tuned so that only
the binding energy provided by both interactions would be sufficient to overcome this barrier,
thus ensuring cooperative gate activation (Fig. 11B). If condition 2 (modularity) holds, then a
20 single scheme for ensuring cooperativity could in principle work for a wide range of gate
configurations.

Design of cooperativity

To explore the design of cooperative building blocks, we focused on the simple
25 system $A + A'-B + B'$ (we refer to this as induced dimerization below, A and B' as the
monomers, and $A'-B$ as the dimerizer). If binding is not cooperative, the amount of the
trimeric complex decreases when $A'-B$ is in stoichiometric excess relative to A and B' : the
formation of intermediate dimeric species of the linker protein binding to either of the
monomers competes with formation of trimeric complexes. On the contrary, if binding is
30 cooperative such that no binding to either monomer occurs in the absence of the other, the
amount of trimeric complex formed becomes insensitive to an excess of the dimerizer. A
simple thermodynamic model of the effect of binding cooperativity on the stoichiometric
response of such induced dimerization systems (Fig. 11B, supplemental materials modeling
section) shows that as the binding cooperativity decreases, there is a corresponding decrease

in the final concentration of full trimeric complexes at high dimerizer concentrations (Fig. 11C).

We hypothesized that a folded four helix bundle like state of the $A'-B$ dimerizer could oppose binding to either A or B' , as the relatively hydrophobic interacting surfaces would likely be sequestered within the folded structure (Fig. 14A). We tested different flexible linker lengths connecting A' with B using heterodimers $1:1'$ and $2:2'$ as a model system. All designs were found to be folded and stable in circular dichroism (CD) guanidine hydrochloride (GdnHCl) denaturation experiments, with unfolding free energies greater than 13 kcal/mol (Fig. 11D, Table 10). Although $1'-2'$ dimerizer constructs with short linkers of 0 and 2 residues, or with a very long 24 residue linker could be purified as monomers (Fig. 14B), they were prone to aggregation. In contrast, designs with 6 and 12 residue linkers remained largely monomeric (data not shown). Small angle x-ray scattering (SAXS) experiments (37) indicate their hydrodynamic radii are close to those of folded four-helix bundle DHDs (Fig. 11E). Linkers in this length range likely allow the two monomers ($1'$ and $2'$) to fold back on each other such that the largely hydrophobic interaction surfaces are buried against each other; such a structure would have to partially unfold for $1'-2'$ to interact with either 1 or 2 with free energy cost ΔG_{open} (Fig. 11B), the magnitude of which determines the extent of cooperativity for the gate. We selected a linker length of 6- or 12- residues for all of the following experiments.

We studied the cooperativity of the induced dimerizer system *in vitro* using native mass spectrometry (Fig. 11F). 1 , 2 and $1'-2'$ were separately expressed in *E.coli* and purified. We first measured the extent to which 1 activates the binding of 2 to $1'-2'$. At 10 μM each of 2 and $1'-2'$, the fraction of 2 in complex with $1'-2'$ increased from 3% to 100% upon addition of 20 μM 1 (data not shown); a fold increase comparable with naturally occurring allosteric systems (35). To assess how this activation of binding influences the sensitivity of binding to stoichiometric imbalance, 10 μM 1 and 2 were titrated with increasing concentrations of $1'-2'$ (Fig. 11F), and the species formed determined by nMS. The heterotrimeric $1:1'-2':2$ complex was observed over a wide range of $1'-2'$ concentrations (data not shown). Even in the presence of a 6 fold excess of $1'-2'$, there was no decrease in the amount of $1:1'-2':2$ formed, and neither $1:1'-2'$ or $1'-2':2$ were observed (data not shown). We define cooperativity as the ratio of the affinities in the presence and absence of the other monomer, which in our model directly relates to the free energy of opening of the dimerizer ($c = e^{\Delta G_{\text{open}}/RT}$, see supplementary materials). Matching the thermodynamic model to native MS

data (data not shown) produces an estimated c value of 991,000, which corresponds to ΔG_{open} of 7 kcal/mol. This is about half the measured unfolding free energy of $I'-2'$, suggesting that binding may not require complete unfolding of the four helix bundle state of the dimerizer.

With linker units displaying cooperative binding, we reasoned that the lack of
5 dependence on stoichiometric excesses of one of the components should extend to more complex gates. Using nMS, we investigated the cooperativity of a 2-input AND gate constructed from the two inputs $I'-3'$ and $3-2'$, and monomers I and 2 brought together by the two inputs (Fig. 11G). As the concentration of the 2 inputs was increased, the amount of heterotetrameric complex plateaued at a stoichiometry of 2:1, and then remained constant up
10 to a molar ratio of 6:1. Very little partial complexes (heterotrimers and heterodimers) were observed, further indicating high cooperativity (data not shown). We constructed a 3-input AND gate from $I'-4'$, $4-3'$, and $3-2'$, which together should control the association of I and 2 (Fig. 11H). Similar to the 2-input AND gate, the amount of full, pentameric complexes only decreased slightly at greater than stoichiometric concentrations of inputs with no detectable
15 competing tetrameric complexes (data not shown).

Modular logic gate construction

We explored the modular combination of DHDs to generate a range of 2-input Cooperatively Inducible Protein Heterodimer (CIPHR) logic gates. Monomers from
20 individual DHDs were linked to effector proteins of interest via genetic fusion, whose colocalization or dissociation is dependent upon the inputs. Taking advantage of previously measured all-by-all specificity matrices (34), two modes of interactions were explored: cognate binding between designed protein pairs, or competitive binding involving multispecific interactions. The choice of effector proteins is independent from the input
25 proteins, allowing diverse functional outputs (Fig. 12A).

We used a variant of the yeast-two-hybrid (Y2H) assay to characterize the behavior of the designed logic gates, using a setup similar to previously described yeast-four-hybrid systems (38, 39). To construct an AND gate, we fused 2 to the Gal4 activation domain (AD), and I to the Gal4 DNA binding domain (DBD). The colocalization of AD and DBD, and
30 resulting induction of transcription of the *His3* gene, is dependent upon the expression of both input proteins ($I'-5$, $5'-2'$). Growth in media lacking histidine required expression of both inputs (Fig. 12B). An OR gate was similarly constructed by linking the $I-6$ fusion to the AD and $7'$ to the DBD. Expression of either of the inputs $I'-7$ or $6'-7$ results in growth by driving association of AD with DBD (Fig. 12C).

We explored the construction of additional boolean logic gates by exploiting binding affinity hierarchies identified in all by all Y2H experiments (34). **8** not only interacts with **8'** but also forms homodimers (Fig. 15A); hence **8'** must outcompete **8** homodimers to form the heterodimer. We constructed a NOT gate by fusing **8** to both AD or DBD; yeast cells stopped growing in the presence of co-expressed **8'** input protein (Fig. 12D). Based on the affinity hierarchy $9:9' \approx 10:10' > 9:10'$ (Fig. 15B), we constructed a NOR gate in which **9** was fused to the AD, **10'** to the DBD, with **9'** and **10** the two inputs. Either or both of the inputs outcompete the **9:10'** interaction and hinder yeast growth (Fig. 12E). Based on the affinity hierarchy $9':1' > 9:9' \approx 1:1' > 9:1$ (Fig. 15B), an XNOR gate was constructed by fusing **9** to AD, **1** to DBD, and using **9'** and **1'** as the two inputs: the presence of either outcompetes the **9:1** binding and blocks growth, but when both are expressed they instead interact with each other and growth is observed (Fig. 12F). Similarly, a NAND gate was designed based on the interaction hierarchy $1':10' > 1:1' \approx 10:10' > 1:10$ (Fig. 15B). Neither **1** nor **10** alone can outcompete the **1':10'** binding and hence growth occurs, but when both are expressed, the free energy of formation of both **1:1'** and **10:10'** outweighs that of **1':10'** and growth is blocked (Fig. 12G).

3 input CIPHR logic gates

We constructed a 3-input AND gate (Fig. 10M) in which monomers **1** and **2** are brought into proximity by the three inputs **1'-4'**, **4-3'**, and **3-2'**. We experimentally tested all eight possible input combinations (Fig. 13A), quantifying all complexes using nMS with both **1** and **2** present. Consistent with proper function of a 3-input AND gate, **1** and **2** only showed significant co-assembly when all three inputs are present (data not shown).

To test the modularity of CIPHR logic gates, we designed two different 3-input CIPHR logic gates using the same 4 pairs of DHDs and tested them via Y2H. To make a 3-input OR gate, **1'-6-7** was fused to AD, and **11'** to DBD. Either one of the 3 inputs (**11-1**, **11-6'**, **11-7'**) is able to bring AD to DBD via their linked proteins (Fig. 13B). Y2H results confirmed the correct behavior of this logic gate in cells: any of the input proteins induces cell growth (Fig. 13C). We constructed a CIPHR disjunctive normal form (DNF, $[A \text{ AND } B] \text{ OR } C$) gate by fusing **1'-6** to AD, **11'** to DBD with inputs **11-7'**, **7-1**, or **11-6'** (Fig. 13D). In Y2H experiments, the DNF gate functioned as designed, with low yeast growth levels when no input or only one of the **11-7'** and **7-1** input proteins are present, and high yeast growth levels otherwise (Fig. 13E).

References for Example 2

1. R. Nussinov, How do dynamic cellular signals travel long distances? *Mol. Biosyst.* **8**, 22–26 (2012).
2. A. W. Reinke, J. Baek, O. Ashenberg, A. E. Keating, Networks of bZIP protein-protein interactions diversified over a billion years of evolution. *Science.* **340**, 730–734 (2013).
- 5 3. Y. E. Antebi, J. M. Linton, H. Klumpe, B. Bintu, M. Gong, C. Su, R. McCardell, M. B. Elowitz, Combinatorial Signal Perception in the BMP Pathway. *Cell.* **170**, 1184–1196.e24 (2017).
4. B. Z. Harris, W. A. Lim, Mechanism and role of PDZ domains in signaling complex assembly. *J. Cell Sci.* **114**, 3219–3231 (2001).
- 10 5. G. Seelig, D. Soloveichik, D. Y. Zhang, E. Winfree, Enzyme-free nucleic acid logic circuits. *Science.* **314**, 1585–1588 (2006).
6. L. Qian, E. Winfree, Scaling up digital circuit computation with DNA strand displacement cascades. *Science.* **332**, 1196–1201 (2011).
7. M. B. Elowitz, S. Leibler, A synthetic oscillatory network of transcriptional regulators. *Nature.* **403**, 335–338 (2000).
- 15 8. T. S. Gardner, C. R. Cantor, J. J. Collins, Construction of a genetic toggle switch in *Escherichia coli*. *Nature.* **403**, 339–342 (2000).
9. A. Tamsir, J. J. Tabor, C. A. Voigt, Robust multicellular computing using genetically encoded NOR gates and chemical “wires.” *Nature.* **469**, 212–215 (2011).
- 20 10. P. Siuti, J. Yazbek, T. K. Lu, Synthetic circuits integrating logic and memory in living cells. *Nat. Biotechnol.* **31**, 448–452 (2013).
11. J. Bonnet, P. Yin, M. E. Ortiz, P. Subsoontorn, D. Endy, Amplifying genetic logic gates. *Science.* **340**, 599–603 (2013).
12. B. H. Weinberg, N. T. H. Pham, L. D. Caraballo, T. Lozanoski, A. Engel, S. Bhatia, W. W. Wong, Large-scale design of robust genetic circuits with multiple inputs and outputs for mammalian cells. *Nat. Biotechnol.* **35**, 453–462 (2017).
- 25 13. S. Ausländer, D. Ausländer, M. Müller, M. Wieland, M. Fussenegger, Programmable single-cell mammalian biocomputers. *Nature.* **487**, 123–127 (2012).
14. A. S. Khalil, T. K. Lu, C. J. Bashor, C. L. Ramirez, N. C. Pyenson, J. K. Joung, J. J. Collins, A synthetic biology framework for programming eukaryotic transcription functions. *Cell.* **150**, 647–658 (2012).
- 30 15. N. Roquet, A. P. Soleimany, A. C. Ferris, S. Aaronson, T. K. Lu, Synthetic recombinase-based state machines in living cells. *Science.* **353**, aad8559 (2016).
16. L. B. Andrews, A. A. K. Nielsen, C. A. Voigt, Cellular checkpoint control using programmable sequential logic. *Science.* **361**, eaap8987 (2018).
- 35 17. B. Angelici, E. Mailand, B. Haefliger, Y. Benenson, Synthetic Biology Platform for Sensing and

- Integrating Endogenous Transcriptional Inputs in Mammalian Cells. *Cell Rep.* **16**, 2525–2537 (2016).
18. J. J. Lohmueller, T. Z. Armel, P. A. Silver, A tunable zinc finger-based framework for Boolean logic computation in mammalian cells. *Nucleic Acids Res.* **40**, 5180–5187 (2012).
- 5 19. A. A. Green, P. A. Silver, J. J. Collins, P. Yin, Toehold switches: de-novo-designed regulators of gene expression. *Cell.* **159**, 925–939 (2014).
20. A. A. Green, J. Kim, D. Ma, P. A. Silver, J. J. Collins, P. Yin, Complex cellular logic computation using ribocomputing devices. *Nature.* **548**, 117–121 (2017).
21. K. Rinaudo, L. Bleris, R. Maddamsetti, S. Subramanian, R. Weiss, Y. Benenson, A universal
10 RNAi-based logic evaluator that operates in mammalian cells. *Nat. Biotechnol.* **25**, 795–801 (2007).
22. L. Wroblewska, T. Kitada, K. Endo, V. Siciliano, B. Stillo, H. Saito, R. Weiss, Mammalian synthetic circuits with RNA binding proteins for RNA-only delivery. *Nat. Biotechnol.* **33**, 839–841 (2015).
- 15 23. S.-H. Park, A. Zarrinpar, W. A. Lim, Rewiring MAP kinase pathways using alternative scaffold assembly mechanisms. *Science.* **299**, 1061–1064 (2003).
24. P. L. Howard, M. C. Chia, S. Del Rizzo, F.-F. Liu, T. Pawson, Redirecting tyrosine kinase signaling to an apoptotic caspase pathway through chimeric adaptor proteins. *Proc. Natl. Acad. Sci. U. S. A.* **100**, 11267–11272 (2003).
- 20 25. B. J. Yeh, R. J. Rutigliano, A. Deb, D. Bar-Sagi, W. A. Lim, Rewiring cellular morphology pathways with synthetic guanine nucleotide exchange factors. *Nature.* **447**, 596–600 (2007).
26. L. Morsut, K. T. Roybal, X. Xiong, R. M. Gordley, S. M. Coyle, M. Thomson, W. A. Lim, Engineering Customized Cell Sensing and Response Behaviors Using Synthetic Notch Receptors. *Cell.* **164**, 780–791 (2016).
- 25 27. K. T. Roybal, L. J. Rupp, L. Morsut, W. J. Walker, K. A. McNally, J. S. Park, W. A. Lim, Precision Tumor Recognition by T Cells With Combinatorial Antigen-Sensing Circuits. *Cell.* **164**, 770–779 (2016).
28. J. E. Dueber, B. J. Yeh, K. Chak, W. A. Lim, Reprogramming Control of an Allosteric Signaling Switch Through Modular Recombination. *Science.* **301**, 1904–1908 (2003).
- 30 29. R. M. Gordley, R. E. Williams, C. J. Bashor, J. E. Toettcher, S. Yan, W. A. Lim, Engineering dynamical control of cell fate switching using synthetic phospho-regulons. *Proc. Natl. Acad. Sci. U. S. A.* **113**, 13528–13533 (2016).
30. J. E. Dueber, E. A. Mirsky, W. A. Lim, Engineering synthetic signaling proteins with ultrasensitive input/output control. *Nat. Biotechnol.* **25**, 660–662 (2007).
- 35 31. X. J. Gao, L. S. Chong, M. S. Kim, M. B. Elowitz, Programmable protein circuits in living cells. *Science.* **361**, 1252–1258 (2018).
32. T. Fink, J. Lonžarić, A. Praznik, T. Plaper, E. Merljak, K. Leben, N. Jerala, T. Lebar, Ž. Strmšek,

- F. Lapenta, M. Benčina, R. Jerala, Design of fast proteolysis-based signaling and logic circuits in mammalian cells. *Nat. Chem. Biol.* **15**, 115–122 (2019).
33. A. J. Smith, F. Thomas, D. Shoemark, D. N. Woolfson, N. J. Savery, Guiding Biomolecular Interactions in Cells Using de Novo Protein-Protein Interfaces. *ACS Synth. Biol.* **8**, 1284–1293 (2019).
- 5
34. Z. Chen, S. E. Boyken, M. Jia, F. Busch, D. Flores-Solis, M. J. Bick, P. Lu, Z. L. VanAernum, A. Sahasrabudde, R. A. Langan, S. Bermeo, T. J. Brunette, V. K. Mulligan, L. P. Carter, F. DiMaio, N. G. Sgourakis, V. H. Wysocki, D. Baker, Programmable design of orthogonal protein heterodimers. *Nature.* **565**, 106–111 (2019).
- 10
35. K. E. Prehoda, J. A. Scott, R. D. Mullins, W. A. Lim, Integration of multiple signals through cooperative regulation of the N-WASP-Arp2/3 complex. *Science.* **290**, 801–806 (2000).
36. B. Yu, I. R. S. Martins, P. Li, G. K. Amarasinghe, J. Umetani, M. E. Fernandez-Zapico, D. D. Billadeau, M. Machius, D. R. Tomchick, M. K. Rosen, Structural and energetic mechanisms of cooperative autoinhibition and activation of Vav1. *Cell.* **140**, 246–256 (2010).
- 15
37. K. N. Dyer, M. Hammel, R. P. Rambo, S. E. Tsutakawa, I. Rodic, S. Classen, J. A. Tainer, G. L. Hura, High-throughput SAXS for the characterization of biomolecules in solution: a practical approach. *Methods Mol. Biol.* **1091**, 245–258 (2014).
38. A. Pause, B. Peterson, G. Schaffar, R. Stearman, R. D. Klausner, Studying interactions of four proteins in the yeast two-hybrid system: structural resemblance of the pVHL/elongin BC/hCUL-2 complex with the ubiquitin ligase complex SKP1/cullin/F-box protein. *Proc. Natl. Acad. Sci. U. S. A.* **96**, 9533–9538 (1999).
- 20
39. B. Sandrock, J. M. Egly, A yeast four-hybrid system identifies Cdk-activating kinase as a regulator of the XPD helicase, a subunit of transcription factor IIIH. *J. Biol. Chem.* **276**, 35328–35333 (2001).
- 25
40. A. S. Dixon, M. K. Schwinn, M. P. Hall, K. Zimmerman, P. Otto, T. H. Lubben, B. L. Butler, B. F. Binkowski, T. Machleidt, T. A. Kirkland, M. G. Wood, C. T. Eggers, L. P. Encell, K. V. Wood, NanoLuc Complementation Reporter Optimized for Accurate Measurement of Protein Interactions in Cells. *ACS Chem. Biol.* **11**, 400–408 (2016).
41. Y.-C. Kwon, M. C. Jewett, High-throughput preparation methods of crude extract for robust cell-free protein synthesis. *Sci. Rep.* **5**, 8663 (2015).
- 30
42. J. R. Porter, C. I. Stains, B. W. Jester, I. Ghosh, A general and rapid cell-free approach for the interrogation of protein-protein, protein-DNA, and protein-RNA interactions and their antagonists utilizing split-protein reporters. *J. Am. Chem. Soc.* **130**, 6488–6497 (2008).
43. S. L. Maude, T. W. Laetsch, J. Buechner, S. Rives, M. Boyer, H. Bittencourt, P. Bader, M. R. Verneris, H. E. Stefanski, G. D. Myers, M. Qayed, B. De Moerloose, H. Hiramatsu, K. Schlis, K. L. Davis, P. L. Martin, E. R. Nemecek, G. A. Yanik, C. Peters, A. Baruchel, N. Boissel, F. Mechinaud, A. Balduzzi, J. Krueger, C. H. June, B. L. Levine, P. Wood, T. Taran, M. Leung, K.
- 35

- T. Mueller, Y. Zhang, K. Sen, D. Lebwohl, M. A. Pulsipher, S. A. Grupp, Tisagenlecleucel in Children and Young Adults with B-Cell Lymphoblastic Leukemia. *N. Engl. J. Med.* **378**, 439–448 (2018).
44. S. S. Neelapu, F. L. Locke, N. L. Bartlett, L. J. Lekakis, D. B. Miklos, C. A. Jacobson, I. Braunschweig, O. O. Oluwole, T. Siddiqi, Y. Lin, J. M. Timmerman, P. J. Stiff, J. W. Friedberg, I. W. Flinn, A. Goy, B. T. Hill, M. R. Smith, A. Deol, U. Farooq, P. McSweeney, J. Munoz, I. Avivi, J. E. Castro, J. R. Westin, J. C. Chavez, A. Ghobadi, K. V. Komanduri, R. Levy, E. D. Jacobsen, T. E. Witzig, P. Reagan, A. Bot, J. Rossi, L. Navale, Y. Jiang, J. Aycock, M. Elias, D. Chang, J. Wieszorek, W. Y. Go, Axicabtagene Ciloleucel CAR T-Cell Therapy in Refractory Large B-Cell Lymphoma. *N. Engl. J. Med.* **377**, 2531–2544 (2017).
45. J. A. Fraietta, S. F. Lacey, E. J. Orlando, I. Pruteanu-Malinici, M. Gohil, S. Lundh, A. C. Boesteanu, Y. Wang, R. S. O'Connor, W.-T. Hwang, E. Pequignot, D. E. Ambrose, C. Zhang, N. Wilcox, F. Bedoya, C. Dorfmeier, F. Chen, L. Tian, H. Parakandi, M. Gupta, R. M. Young, F. B. Johnson, I. Kulikovskaya, L. Liu, J. Xu, S. H. Kassim, M. M. Davis, B. L. Levine, N. V. Frey, D. L. Siegel, A. C. Huang, E. J. Wherry, H. Bitter, J. L. Brogdon, D. L. Porter, C. H. June, J. J. Melenhorst, Determinants of response and resistance to CD19 chimeric antigen receptor (CAR) T cell therapy of chronic lymphocytic leukemia. *Nat. Med.* **24**, 563–571 (2018).
46. C. H. June, R. S. O'Connor, O. U. Kawalekar, S. Ghassemi, M. C. Milone, CAR T cell immunotherapy for human cancer. *Science*. **359**, 1361–1365 (2018).
47. A. H. Long, W. M. Haso, J. F. Shern, K. M. Wanhainen, M. Murgai, M. Ingaramo, J. P. Smith, A. J. Walker, M. E. Kohler, V. R. Venkateshwara, R. N. Kaplan, G. H. Patterson, T. J. Fry, R. J. Orentas, C. L. Mackall, 4-1BB costimulation ameliorates T cell exhaustion induced by tonic signaling of chimeric antigen receptors. *Nat. Med.* **21**, 581–590 (2015).
48. E. J. Wherry, M. Kurachi, Molecular and cellular insights into T cell exhaustion. *Nat. Rev. Immunol.* **15**, 486–499 (2015).
49. E. John Wherry, S. J. Ha, S. M. Kaech, W. Nicholas Haining, S. Sarkar, V. Kalia, S. Subramaniam, J. Blattman, D. L. Barber, R. Ahmed, Molecular Signature of CD8+ T Cell Exhaustion during Chronic Viral Infection. *Immunity*. **27** (2007), doi:10.1016/j.immuni.2007.11.006.
50. K. E. Pauken, M. A. Sammons, P. M. Odorizzi, S. Manne, J. Godec, O. Khan, A. M. Drake, Z. Chen, D. R. Sen, M. Kurachi, R. A. Barnitz, C. Bartman, B. Bengsch, A. C. Huang, J. M. Schenkel, G. Vahedi, W. N. Haining, S. L. Berger, E. J. Wherry, Epigenetic stability of exhausted T cells limits durability of reinvigoration by PD-1 blockade. *Science*. **354**, 1160–1165 (2016).
51. K. E. Prehoda, W. A. Lim, How signaling proteins integrate multiple inputs: a comparison of N-WASP and Cdk2. *Curr. Opin. Cell Biol.* **14**, 149–154 (2002).

Table 7. Thermodynamic Modeling of Cooperativity

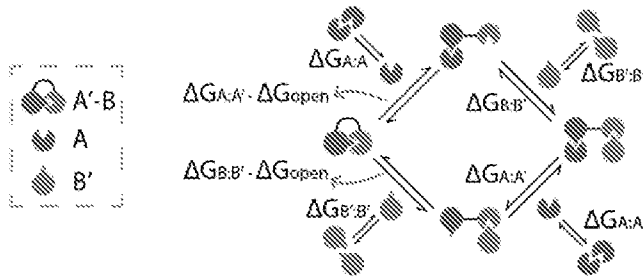
Input	
Output	
Cooperative	Noncooperative
$N-1$	$N-2$

Referring to Table 7, for an induced dimerization system involving proteins A, A'-B, and B', a stoichiometric excess (N) of the A'-B protein results in partially assembled dimeric complexes if the binding is non-cooperative, but fully assembled trimeric complexes if the binding is cooperative.

We model the cooperatively induced dimerization system at thermodynamic equilibrium. Shown below (Table 8), assuming a 'closed' state for A'-B, where the binding interfaces are buried within the four-helix bundle, the binding of A'-B to either A or B' helix hairpins needs to overcome an energy barrier of transitioning from the 'closed' to 'open' state (ΔG_{open}). Therefore the free energy of binding between A'-B to A or B' can be expressed as $\Delta G_{A:A'} - \Delta G_{open}$ and $\Delta G_{B:B'} - \Delta G_{open}$, respectively, where $\Delta G_{A:A'}$ and $\Delta G_{B:B'}$ represent the free energy of binding between the cognate pairs in the absence of the fusion. Once the A:A'-B or A-B':B complexes form, subsequent binding can be simply represented by the binding between cognate heterodimers: $\Delta G_{A:A'}$ or $\Delta G_{B:B'}$. We also observed the presence of (A)₂ and (B')₂ homodimers, therefore added free energy terms describing such processes into the model ($\Delta G_{A:A}$ or $\Delta G_{B':B'}$).

20

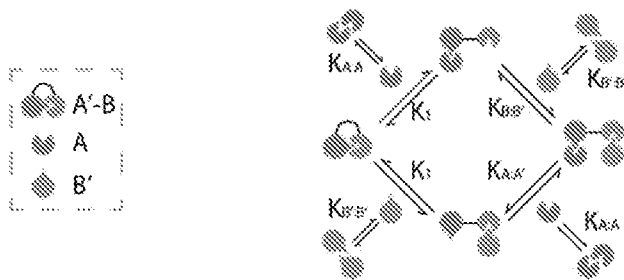
Table 8



ΔG relates to equilibrium constants by $\Delta G = -RT \ln K$, and we further consider the system in terms of K . We make the simplifying assumption that the affinity of A'-B to either A or B' is identical ($K_1 = [A:A'-B]/([A][A'-B]) = [A'-B:B']/([B][A'-B])$). Finally, we define the cooperativity of the system, c , as the ratio between the equilibrium constants in the presence or absence of the other partner ($c = K_{B:B'}/K_1 = K_{A:A'}/K_1$). For an entirely non-cooperative process ($c=1$), $K_{B:B'}=K_1$ and $K_{A:A'}=K_1$ i.e., the first binding event does not affect the affinity of the subsequent binding event.

Since $K_1 = \exp(-(\Delta G_{A:A'} - \Delta G_{open})/RT)$, rewriting the equation for c in terms of free energies leads to $c = \exp(\Delta G_{open}/RT)$. Therefore, the extent of cooperativity is solely determined by the magnitude of the free energy required to partially unfold/expose the buried binding interfaces of the dimerizer A'-B.

We note that explicitly incorporating the equilibrium constants for homodimerization ($K_{A:A}$ and $K_{B':B'}$) only affect the absolute position of each equilibrium, but does not affect the magnitude of the cooperativity (see Table 9). Indeed, taking A as an example, the binding to the closed state becomes $K_1 * K_{A:A}$, and the binding to the open state becomes $K_{A:A'} * K_{A:A}$. Because $K_{A:A}$ is present in both the numerator and the denominator, they cancel out, and c remains purely defined by the relative magnitudes of K_1 and $K_{A:A'}$.



20

Table 9

We solved the following system of equations in *Mathematica* to simulate the amount
 5 of A:A'-B:B' at equilibrium as a function of the initial concentration of A'-B:

$$\begin{aligned}
 K_{A:A} &= \frac{[A_2]}{[A][A]} \\
 K_{B':B'} &= \frac{[B'_2]}{[B'][B']} \\
 K_1 &= \frac{[A:A'-B]}{[A][A'-B]} \\
 K_1 &= \frac{[B':A'-B]}{[B'][A'-B]} \\
 K_{A:A'} &= \frac{[A:A'-B:B']}{[A][A'-B:B']} \\
 K_{B:B'} &= \frac{[A:A'-B:B']}{[B'][A:A'-B]} \\
 [A]_{tot} &= 2 * [A_2] + [A] + [A:A'-B] + [A:A'-B:B'] \\
 [B']_{tot} &= 2 * [B'_2] + [B'] + [A'-B:B'] + [A:A'-B:B'] \\
 [A'-B]_{tot} &= [A'-B:B'] + [A:A'-B] + [A:A'-B:B']
 \end{aligned}$$

We knew from previous native MS titration experiments that the equilibrium
 dissociation constants of cognate designed heterodimers (DHDs) is in the ~10 nM range (1),
 10 therefore $K_{A:A'} = K_{B:B'} = 0.1 \text{ nM}^{-1}$. Varying values of K_1 (and hence the cooperativity factor, c
 $= K_{A:A'}/K_1$) showed different responses of the amount of A:A'-B:B' at equilibrium as a
 function of the initial concentration of A'-B, as shown in Fig. 12C.

We experimentally estimated K_1 using native MS experiments. Mixing 10 μM of *I*
 and *I'-2'* resulted in no detectable amount of the *I:I'-2'* complex, suggesting very weak
 15 binding. The sensitivity of native MS places a lower-bound on the concentration of species
 that can be detected (0.0375 μM). Using this value, a lower-bound for the affinity of *I:I'-2'*
 can be estimated ($1/K_1 \geq 2.65 \text{ mM}$). This is close to the value of 9.91 mM obtained by
 calculating the affinity based on the c value of 991,000 reported in FIG. 13H.

This thermodynamic modeling demonstrates that binding cooperativity can be
 20 achieved for an induced dimerization system through occlusion of the binding interfaces. We
 achieved this by fusing hairpins via a flexible linker, rationalizing that the spontaneous
 folding of these constructs would bury the interaction interfaces on the inside of a four helical
 bundle like topology. Formation of these structures is corroborated by: *i*) SAXS profiles that

are consistent with DHDs structures, *ii*) *m*-values from chemical denaturation experiments consistent with $\Delta SASA$ for the unfolding of DHD topologies, and *iii*) $\Delta G_{\text{open}} < \Delta G_{\text{folding}}$, suggesting that exposing the binding interfaces requires partial unfolding of these fused constructs, but does not exceed the folding free energy of these proteins (a physically
 5 unrealistic scenario).

Materials and Methods

Buffer and media recipe

TBM-5052: 1.2% [wt/vol] tryptone, 2.4% [wt/vol] yeast extract, 0.5% [wt/vol] glycerol, 0.05% [wt/vol] D-glucose, 0.2% [wt/vol] D-lactose, 25 mM Na₂HPO₄, 25 mM
 10 KH₂PO₄, 50 mM NH₄Cl, 5 mM Na₂SO₄, 2 mM MgSO₄, 10 μM FeCl₃, 4 μM CaCl₂, 2 μM MnCl₂, 2 μM ZnSO₄, 400 nM CoCl₂, 400 nM NiCl₂, 400 nM CuCl₂, 400 nM Na₂MoO₄, 400 nM Na₂SeO₃, 400 nM H₃BO₃.

Lysis buffer: 20 mM Tris, 300 mM NaCl, 20 mM Imidazole, pH 8.0 at room temperature.

15 Wash buffer: 20 mM Tris, 300mM NaCl, 30 mM Imidazole, pH 8.0 at room temperature.

Elution buffer: 20 mM Tris, 300 mM NaCl, 250 mM Imidazole, pH 8.0 at room temperature.

TBS buffer: 20 mM Tris pH 8.0, 100 mM NaCl.

20 YPAD buffer: Peptone 20 g/L, yeast extract 10 g/L, Adenine hemisulfate 10 μg/L, dextrose (20 g/L).

C-Trp-Ura-Leu-His+Adenine: hemisulfate+Glucose.

Yeast nitrogen base w/o amino acids (6.7 g/L), synthetic DO media (-Leu/-His/-Trp/-Ura) (1.4 g/L), dextrose (20 g/L), adenine hemisulfate (10 μg/L).

25 Construction of synthetic genes

For the expression of proteins in *E. coli*, synthetic genes were ordered from Genscript Inc. (Piscataway, N.J., USA) and delivered in pET21-NESG *E. coli* expression vector, inserted between the NdeI and XhoI sites. For each expression construct, a hexahistidine tag followed by a tobacco etch virus (TEV) protease cleavage site
 30 (GSSHHHHHSSGENLYFQGS) (SEQ ID NO:328) were added in frame at the N-terminus of the protein. A stop codon was introduced at the 3' end of the protein coding sequence to prevent expression of the C-terminal hexahistidine tag in the vector.

Genes for yeast-two-hybrid (Y2H) studies were cloned into plasmids bearing the GAL4 DNA-binding domain (poDBD) and the GAL4 transcription activation domain (poAD) (2). Input proteins were cloned into plasmids V510 (uracil auxotrophic selection marker) and MX1 (bleomycin selection marker). Genes were expressed under the control of
5 ADH1 promoters.

Protein expression

Plasmids were transformed into chemically competent *E. coli* expression strain Lemo21TM(DE3) (New England Biolabs) for protein expression. Following transformation
10 and overnight growth, single colonies were picked from agar plates into 5 ml Luria-Bertani (LB) medium containing 100 µg/mL carbenicillin (for pET21-NESG vectors) with shaking at 225 rpm for 18 hours at 37°C. Proteins were expressed using the autoinduction method (7): starter cultures were further diluted into 500 ml TBM-5052 containing 100 µg/mL
15 carbenicillin, and incubated with shaking at 225 rpm for 24 hours at 37°C.

Affinity purification

E. coli cells were harvested by centrifugation at 5000 rcf for 15 minutes at 4°C and the pellet resuspended in 18 ml lysis buffer. EDTA-free cocktail protease inhibitor (Roche), lysozyme, and DNase were added to the resuspended cell pellet, followed by cell lysis via
20 sonication at 70% power for 5 minutes. Lysates were clarified by centrifugation at 4°C and 18,000 rpm for 45 minutes and applied to columns containing Ni-NTA (Qiagen) resin pre-equilibrated with lysis buffer. The column was washed two times with 5 column volumes (CV) of wash buffer, followed by 5 CV of elution buffer for protein elution.

25 Size-exclusion chromatography (SEC)

Eluted proteins were buffer exchanged into lysis buffer. N-terminal hexahistidine tags were removed with TEV protease cleavage overnight at room temperature, at a ratio of 1 mg TEV for 100 mg of protein. After TEV cleavage, sample was passed over a fresh Ni-NTA
30 column and washed with 1.5 CV of lysis buffer, collecting flow through. The resulting proteins were purified by SEC using a SuperdexTM 75 10/300 increase column (GE Healthcare) in TBS buffer.

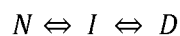
Circular dichroism (CD) measurements

Circular dichroism (CD) wavelength scans (260 - 195 nm) and temperature melts (25 - 95 °C) were performed using an AVIV™ model 420 CD spectrometer, with protein
 5 samples diluted to 0.25 mg/ml in PBS pH 7.4 in a 0.1-cm cuvette. Temperature melts were carried out at a heating rate of 4 °C/min and monitored by the change in ellipticity at 222 nm.

GdmCl titrations were performed on a JASCO™ model J-1500 with automated titration apparatus in PBS pH 7.4 at 25 °C, with protein concentrations between 0.08 mg/ml to 0.025 mg/ml in a 1-cm cuvette with stir bar. Each titration consisted of at least 34 evenly
 10 distributed GdmCl concentration points up to 7.4 M with 30 seconds mixing time for each step. Titrant solution consisted of the same concentration of protein in PBS and GdmCl.

CD data analysis and model fitting

Folding free energies were obtained by fitting equilibrium denaturation data. Fused
 15 hairpin constructs had biphasic unfolding transitions, indicating the existence of an intermediate on their respective energy landscapes. Since native MS showed that Linker 0, Linker 2, Linker 6, and Linker 12 were almost exclusively monomeric in buffer (data not shown), it was concluded that these intermediates were partially folded monomeric species. Thus, the chemical denaturation data of these proteins was fitted to a unimolecular 3-state
 20 model:



where N represents the fully folded state, I a partially folded intermediate, and D the denatured state. The fraction of each species can be written as a function of $K_1 = [I]/[N]$ and
 25 $K_2 = [D]/[I]$, the equilibrium constants for the first and second transitions respectively:

$$\begin{aligned} f_N &= (1 + K_1 + K_1 \cdot K_2)^{-1} \\ f_I &= (1 + K_2 + \frac{1}{K_1})^{-1} \\ f_D &= (1 + \frac{1}{K_2} + \frac{1}{K_1 \cdot K_2})^{-1} \end{aligned}$$

In the context of equilibrium chemical denaturation experiments, the free energy of unfolding is a linear function of denaturant concentration:

30

$$\Delta G_{[den]} = \Delta G_{buffer} - m \cdot [den]$$

where $\Delta G_{[den]}$ represents the free energy of the system at a given concentration of denaturant, ΔG_{buffer} is the corresponding free energy change in the absence of denaturant, and m is a constant of proportionality that relates to the change in solvent-accessible surface area upon unfolding ($\Delta SASA$). Thus, the effect of denaturant on the equilibrium constant relating to each transition can be written as a function of its free energy difference in buffer, and a specific m -value:

$$K_1 = \exp\left(\frac{m_1 \cdot [den] - \Delta G_1}{R \cdot T}\right)$$

$$K_2 = \exp\left(\frac{m_2 \cdot [den] - \Delta G_2}{R \cdot T}\right)$$

By combining these expressions with the definitions for f_N, f_I, f_D , the fractional distribution of each species can be expressed as a function of denaturant concentration, and the free energy change corresponding to each transition (in buffer). Finally, for an ensemble spectroscopic technique such as CD, the observed signal (the dependent variable) as a function of denaturant concentration (the independent variable) can be expressed as a linear combination of the spectroscopic signals corresponding to each species, weighed by their fractional contribution to the ensemble:

$$MRE_{222nm} = f_N \cdot MRE_N + f_I \cdot MRE_I + f_D \cdot MRE_D$$

Where MRE_N, MRE_I, MRE_D represent the spectroscopic signatures (baselines) for the native, intermediate, and denatured states respectively. This equation was used to fit chemical denaturation data for the different linker proteins, and the fitted parameters are reported in Table 10. For Linker 24 in buffer, native MS revealed a significant proportion of dimer (data not shown). Therefore, this model is not entirely appropriate for describing the unfolding, and the fitted values for this construct should be interpreted with care. Nevertheless, denaturation performed at different concentrations of protein revealed that the position of the second transition was concentration-independent, and thus unimolecular. For this event, the model holds.

The total m -values for these linked hairpins were found to be around $3 \text{ kcal mol}^{-1} \text{ M}^{-1}$. It has been shown that m -values correlate with $\Delta SASA$ of unfolding (8). For the folded state, $SASA$ was estimated from the structures of DHDs (1) using PyMOL™ to be 8800 \AA^2 . For the unfolded state, $SASA$ was estimated using ProtSA™ (9, 10), and is about $20,000 \text{ \AA}^2$. Thus,

$\Delta SASA$ for the unimolecular unfolding of a fused hairpin should be around 11,000 \AA^2 , which would have a predicted m -value of 3.3. This number is in close agreement with the fitted parameters reported here, in line with the notion that the folded state for these linker proteins has a four helix bundle topology.

5 **Small Angle X-ray Scattering (SAXS)**

Protein samples were purified by SEC in 25 mM Tris pH 8.0, 150 mM NaCl and 2% glycerol; elution fractions preceding the void volume of the column were used as blanks for buffer subtraction. Scattering measurements were performed at the SIBYLSTM 12.3.1 beamline at the Advanced Light Source. The sample-to-detector distance was 1.5 m, and the X-ray wavelength (λ) was 1.27 \AA , corresponding to a scattering vector q ($q = 4\pi \sin \theta/\lambda$, where 2θ is the scattering angle) range of 0.01 to 0.3 \AA^{-1} . A series of exposures were taken of each well, in equal sub-second time slices: 0.3-s exposures for 10 s resulting in 32 frames per sample. For each sample, data were collected for two different concentrations to test for concentration-dependent effects; ‘low’ concentration samples ranged at 2.5 mg/ml and ‘high’ concentration samples at 5 mg/ml. Data were processed using the SAXS FrameSliceTM online server and analyzed using the ScÅtterTM software package (11, 12). The FoXSTM online server (13, 14) was used to compare experimental scattering profiles to design models and calculate quality of fit (χ) values.

20 **Yeast two-hybrid assay for logic gates**

Chemically competent cells of yeast strain PJ69-4a (MATa trp1-901 leu2-3,112 ura3-52 his3-200 gal4(deleted) gal80(deleted) LYS2::GAL1-HIS3 GAL2-ADE2 met2::GAL7-lacZ) were transformed with the appropriate pair of plasmids containing DNA binding domains (DBD) or activation domains (AD), using the LiAc/SS carrier DNA/PEG method (15). For two input CIPHR logic gates, genes encoding the input proteins (together with selection markers) were genetically integrated into either or both of the Ura3 locus (uracil auxotrophic selection marker) or the YCR043 locus (bleomycin selection marker). In the case of three input CIPHR logic gates, genes encoding two input proteins were genetically integrated as described, with the additional input cloned downstream of either the AD or DBD plasmid, separated by a p2a and nuclear localization sequence (GSGATNFSLLKQAGDVEENPGPGDKAELIPEPPKKRKRKVELGTA; SEQ ID NO:330). The p2a sequence ensures translational cleavage to make the additional input protein a separate protein. The selection of transformed yeast cells was performed in synthetic dropout (SDO) medium lacking tryptophan and leucine for 48 h with shaking at 1,000 r.p.m. at 30 °C.

The resulting culture was diluted 1:100 and grown for 16 h in fresh SDO medium lacking tryptophan and leucine, before being diluted 1:100 in fresh SDO medium lacking tryptophan, leucine and histidine. The culture was incubated with shaking at 1,000 r.p.m. at 30 °C. As it is necessary to bring the DBD and the transcription activation domain into proximity for the growth of yeast cells in medium lacking histidine, successful activation of logic gates was indicated by the growth of yeast cells (16, 17). The optical density of yeast cells was recorded at 24 h, 48 h, and 72 h.

Table 10. Fitted parameters for equilibrium chemical denaturation. Errors represent fitting errors.

	Linker 0	Linker 2	Linker 6	Linker 12	Linker 24
$\Delta G_1^{(N \rightleftharpoons I)}$ (kcal mol ⁻¹)	3.6 (±0.4)	3.5 (±0.2)	3.5 (±0.2)	2.7 (±0.1)	3.7 (±0.3)
$\Delta G_2^{(I \rightleftharpoons D)}$ (kcal mol ⁻¹)	9.8 (±0.6)	10.7 (±0.4)	12.2 (±0.4)	10.6 (±0.5)	10.4 (±0.8)
$\Delta G_{tot}^{(N \rightleftharpoons D)}$ (kcal mol ⁻¹)	13.5 (±0.7)	14.1 (±0.4)	15.7 (±0.5)	13.3 (±0.5)	14.1 (±0.8)
m_1 (kcal mol ⁻¹ M ⁻¹)	1.1 (±0.2)	1.0 (±0.1)	0.9 (±0.1)	0.75 (±0.05)	1.1 (±0.1)
m_2 (kcal mol ⁻¹ M ⁻¹)	1.8 (±0.1)	1.97 (±0.07)	2.22 (±0.08)	1.96 (±0.08)	2.0 (±0.1)
m_{tot} (kcal mol ⁻¹ M ⁻¹)	2.9 (±0.2)	3.0 (±0.1)	3.1 (±0.1)	2.71 (±0.09)	3.1 (±0.2)
MRE_N (deg cm ² dmol ⁻¹)	-23,574 (±114)	-27,561 (±84)	-24,712 (±63)	-33,849 (±131)	-26,438 (±123)
MRE_I (deg cm ² dmol ⁻¹)	-16,330 (±749)	-18,139 (±540)	-14,779 (±710)	-17,362 (±1,158)	-15,567 (±914)
MRE_D (deg cm ² dmol ⁻¹)	-525 (±107)	-785 (±82)	-937 (±68)	-1,104 (±99)	-1,125 (±133)

10

References

1. Z. Chen, S. E. Boyken, M. Jia, F. Busch, D. Flores-Solis, M. J. Bick, P. Lu, Z. L. VanAernum, A. Sahasrabudhe, R. A. Langan, S. Bermeo, T. J. Brunette, V. K.

- Mulligan, L. P. Carter, F. DiMaio, N. G. Sgourakis, V. H. Wysocki, D. Baker, Programmable design of orthogonal protein heterodimers. *Nature*. **565**, 106–111 (2019).
2. S. E. Boyken, Z. Chen, B. Groves, R. A. Langan, G. Oberdorfer, A. Ford, J. M. Gilmore, C. Xu, F. DiMaio, J. H. Pereira, B. Sankaran, G. Seelig, P. H. Zwart, D. Baker, De novo design of protein homo-oligomers with modular hydrogen-bond network-mediated specificity. *Science*. **352**, 680–687 (2016).
 3. A. S. Dixon, M. K. Schwinn, M. P. Hall, K. Zimmerman, P. Otto, T. H. Lubben, B. L. Butler, B. F. Binkowski, T. Machleidt, T. A. Kirkland, M. G. Wood, C. T. Eggers, L. P. Encell, K. V. Wood, NanoLuc Complementation Reporter Optimized for Accurate Measurement of Protein Interactions in Cells. *ACS Chem. Biol.* **11**, 400–408 (2016).
 4. M. E. Lee, W. C. DeLoache, B. Cervantes, J. E. Dueber, A Highly Characterized Yeast Toolkit for Modular, Multipart Assembly. *ACS Synth. Biol.* **4**, 975–986 (2015).
 5. A. S. Khalil, T. K. Lu, C. J. Bashor, C. L. Ramirez, N. C. Pyenson, J. K. Joung, J. J. Collins, A synthetic biology framework for programming eukaryotic transcription functions. *Cell*. **150**, 647–658 (2012).
 6. A. Aranda-Díaz, K. Mace, I. Zuleta, P. Harrigan, H. El-Samad, Robust Synthetic Circuits for Two-Dimensional Control of Gene Expression in Yeast. *ACS Synth. Biol.* **6**, 545–554 (2017).
 7. F. W. Studier, Protein production by auto-induction in high density shaking cultures. *Protein Expr. Purif.* **41**, 207–234 (2005).
 8. J. K. Myers, C. N. Pace, J. M. Scholtz, Denaturant m values and heat capacity changes: relation to changes in accessible surface areas of protein unfolding. *Protein Sci.* **4**, 2138–2148 (1995).
 9. P. Bernadó, M. Blackledge, J. Sancho, Sequence-specific solvent accessibilities of protein residues in unfolded protein ensembles. *Biophys. J.* **91**, 4536–4543 (2006).
 10. J. Estrada, P. Bernadó, M. Blackledge, J. Sancho, ProtSA: a web application for calculating sequence specific protein solvent accessibilities in the unfolded ensemble. *BMC Bioinformatics.* **10**, 104 (2009).
 11. K. N. Dyer, M. Hammel, R. P. Rambo, S. E. Tsutakawa, I. Rodic, S. Classen, J. A. Tainer, G. L. Hura, High-throughput SAXS for the characterization of biomolecules in solution: a practical approach. *Methods Mol. Biol.* **1091**, 245–258 (2014).
 12. R. P. Rambo, J. A. Tainer, Characterizing flexible and intrinsically unstructured biological macromolecules by SAS using the Porod-Debye law. *Biopolymers.* **95**, 559–571 (2011).
 13. D. Schneidman-Duhovny, M. Hammel, A. Sali, FoXS: a web server for rapid

- computation and fitting of SAXS profiles. *Nucleic Acids Res.* **38**, W540–4 (2010).
14. D. Schneidman-Duhovny, M. Hammel, J. A. Tainer, A. Sali, Accurate SAXS profile computation and its assessment by contrast variation experiments. *Biophys. J.* **105**, 962–974 (2013).
- 5 15. R. H. Schiestl, R. D. Gietz, High efficiency transformation of intact yeast cells using single stranded nucleic acids as a carrier. *Curr. Genet.* **16**, 339–346 (1989).
16. C. T. Chien, P. L. Bartel, R. Sternglanz, S. Fields, The two-hybrid system: a method to identify and clone genes for proteins that interact with a protein of interest. *Proc. Natl. Acad. Sci. U. S. A.* **88**, 9578–9582 (1991).
- 10 17. P. L. Bartel, J. A. Roecklein, D. SenGupta, S. Fields, A protein linkage map of Escherichia coli bacteriophage T7. *Nat. Genet.* **12**, 72–77 (1996).
18. Z. L. VanAernum, J. D. Gilbert, M. E. Belov, A. A. Makarov, S. R. Horning, V. H. Wysocki, Surface-Induced Dissociation of Noncovalent Protein Complexes in an Extended Mass Range Orbitrap Mass Spectrometer. *Anal. Chem.* **91**, 3611–3618 (2019).
- 15 19. M. T. Marty, A. J. Baldwin, E. G. Marklund, G. K. A. Hochberg, J. L. P. Benesch, C. V. Robinson, Bayesian deconvolution of mass and ion mobility spectra: from binary interactions to polydisperse ensembles. *Anal. Chem.* **87**, 4370–4376 (2015).
20. Y.-C. Kwon, M. C. Jewett, High-throughput preparation methods of crude extract for robust cell-free protein synthesis. *Sci. Rep.* **5**, 8663 (2015).
- 20 21. M. C. Jewett, J. R. Swartz, Mimicking the Escherichia coli cytoplasmic environment activates long-lived and efficient cell-free protein synthesis. *Biotechnol. Bioeng.* **86**, 19–26 (2004).
22. A. D. Silverman, N. Kelley-Loughnane, J. B. Lucks, M. C. Jewett, Deconstructing Cell-Free Extract Preparation for in Vitro Activation of Transcriptional Genetic Circuitry. *ACS Synth. Biol.* **8**, 403–414 (2019).
- 25 23. J. R. Swartz, M. C. Jewett, K. A. Woodrow, in *Recombinant Gene Expression: Reviews and Protocols*, P. Balbás, A. Lorence, Eds. (Humana Press, Totowa, NJ, 2004), pp. 169–182.
24. V. Muñoz, L. Serrano, Elucidating the folding problem of helical peptides using empirical parameters. III. Temperature and pH dependence. *J. Mol. Biol.* **245**, 297–308 (1995).
- 30 23. V. Muñoz, L. Serrano, Elucidating the folding problem of helical peptides using empirical parameters. III. Temperature and pH dependence. *J. Mol. Biol.* **245**, 297–308 (1995).

We claim

1. A designed heterodimer protein, comprising:
 - (a) a monomer A polypeptide, wherein the monomer A polypeptide is a non-naturally occurring polypeptide comprising 1-5 alpha helices connected by amino acid linkers; and
 - (b) a monomer B polypeptide, wherein the monomer B polypeptide is a non-naturally occurring polypeptide comprising 1-5 alpha helices connected by amino acid linkers,
- 10 wherein monomer A and monomer B non-covalently interact to form the designed heterodimer protein.

2. The designed heterodimer protein of claim 1 wherein monomer A and monomer B have their interaction specificity determined by at least one designed hydrogen bond
- 15 network at the interface between monomer A and monomer B.

3. The designed heterodimer protein of any one of claims 1-2, wherein at least one of monomer A and monomer B comprises a hexahistidine tag, and/or wherein
 - (i) monomer A comprises 2 alpha helices, and monomer B comprises 3 alpha helices;
 - 20 (ii) monomer A comprises 3 alpha helices and monomer B comprises 3 alpha helices;
 - (iii) monomer A comprises 3 alpha helices and monomer B comprises 4 alpha helices,
 - (iv) monomer A comprises 4 alpha helices and monomer B comprises 3 alpha helices;
 - (v) monomer A comprises 4 alpha helices and monomer B comprises 4 alpha helices;
 - (vi) monomer A comprises 5 alpha helices and monomer B comprises 4 alpha helices;
 - 25 (vii) monomer A comprises 4 alpha helices and monomer B comprises 5 alpha helices;
 - (viii) monomer A comprises 5 alpha helices and monomer B comprises 5 alpha helices;
 - (ix) monomer A comprises 2 alpha helices and monomer B comprises 2 alpha helices;
 - or (x) monomer A comprises 3 alpha helices and monomer B comprises 2 alpha helices.

- 30 4. The designed heterodimer protein of any one of claims 1-3, wherein:
 - (i) monomer A comprises a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an odd-numbered SEQ ID NO

selected from the group consisting of selected from the group SEQ ID NOS: 1-290;
and

(ii) monomer B comprises a polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence
5 identity along the length of the amino acid sequence of an even-numbered SEQ ID NO
selected from the group consisting of selected from the group SEQ ID NOS: 1-290,
wherein the even-numbered SEQ ID NO is the binding partner of the odd-numbered
SEQ ID NO. in step (i).

10 5. The designed heterodimer protein of claim 4, wherein amino acid changes from
the reference amino acid sequence are conservative amino acid substitutions.

6. The designed heterodimer protein of claim 4 or 5, wherein at least 20%, 25%,
30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%,
15 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of defined interface positions are
invariant compared to the reference amino acid sequence.

7. A non-naturally occurring polypeptide comprising a polypeptide having at least
70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or
20 100% sequence identity along the length of the amino acid sequence selected from the
group consisting of SEQ ID NOS: 1-290.

8. The non-naturally occurring polypeptide of claim 8, wherein amino acid
changes from the reference amino acid sequence are conservative amino acid
25 substitutions.

9. The non-naturally occurring polypeptide of claim 7 or 8, wherein at least 20%,
25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%,
92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at
30 defined interface positions are invariant compared to the reference amino acid
sequence.

10. A protein comprising 2, 3, 4, or more non-naturally occurring polypeptides
having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%,

98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS: 1-290, wherein the 2, 3, 4, or more naturally occurring polypeptides are covalently linked.

- 5 11. The protein of claim 10, wherein each of the 2, 3, 4, or more non-naturally occurring polypeptides are different.
12. The protein of claim 10 or 11, wherein each of the 2, 3, 4, or more non-naturally occurring polypeptides are present in a fusion protein.
- 10 13. The protein of any one of claims 10-12, wherein each of the 2, 3, 4, or more non-naturally occurring polypeptides have at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an odd-numbered SEQ ID NO: selected from the group consisting of SEQ ID NOS:1-290.
- 15 14. The protein of any one of claims 10-12, wherein each of the 2, 3, 4, or more non-naturally occurring polypeptides have at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an even-numbered SEQ ID NO: selected from the group consisting of SEQ ID NOS:1-290.
- 20 15. The protein of any one of claims 10-12, wherein the 2, 3, 4, or more non-naturally occurring polypeptides include:
- 25 (a) polypeptides having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an odd-numbered SEQ ID NO: selected from the group consisting of SEQ ID NOS:1-290; and
- (b) polypeptides having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an even-numbered SEQ ID NO: selected from the group consisting of SEQ ID NOS:1-290.
- 30

16. The protein of any one of claims 10-15, comprising the amino acid sequence having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group consisting of SEQ ID NOS:291, 294, 296, 299, and 302-305.
- 5
17. The protein of claim any one of claims 10-16, wherein at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions are invariant compared to the reference amino acid sequence.
- 10
18. A protein scaffold, comprising
- a) a first designed component comprised of any number of monomer A polypeptides and/or monomer B polypeptides, each from different heterodimers, connected into a single component by amino acid linkers.
- 15
- b) a second designed component, comprising corresponding monomers for each monomer A and/or monomer B in the first designed component one;
- wherein the first and second designed components interact to form the protein scaffold, and wherein each monomer A only interacts in the scaffold with its monomer B mate.
- 20
19. The protein scaffold of claim 18, wherein the first designed component comprises the protein of any one of claims 10-17, and the second designed component comprises a plurality of individual polypeptides of any one of claims 7-9.
- 25
20. The protein scaffold of claim 18 or 19, wherein any of the monomers in the second designed component comprises a linker.
21. The protein scaffold of any one of claims 18-20, wherein the scaffold is stable up to 95°C and has a guanidine denaturation midpoint of 4 M.
- 30
22. A method of forming the designed heterodimer protein of claims 1-6, comprising:
- a) providing two of the monomers as unlinked monomers;
- b) providing the other two monomers as linked monomers

whereby the unlinked monomers associate with their respective monomer of the same heterodimer, and not with any of the other monomers.

21. A designed heterodimer protein comprising:
- 5 a) asymmetric buried hydrogen bond networks incorporated into regularly repeating backbone structures; and
- b) helix hairpin helix monomers wherein the supercoil phases of the helices are fixed at 0, 90, 180, or 270 degrees and the supercoil twist (ω_0) and helical twist (ω_1) are held constant for either a two layer left handed super coil ($\omega_0=-2.85$ and
- 10 $\omega_1=102.85$), or a 5 layer untwisted bundle ($\omega_0=0$ and $\omega_1=100$)
22. A nucleic acid encoding the polypeptide of claims 7-9 or the protein of any one of claims 10-17.
- 15 23. An expression vector comprising the nucleic acid of claim 22 operatively linked to a promoter.
24. A cell comprising the nucleic acid of claim 22, the expression vector of claim 23, and/or the polypeptide, protein, heterodimer protein, and/or protein scaffold of any
- 20 claim herein.
25. Use of the polypeptide, protein, heterodimer protein, protein scaffold, nucleic acid, expression vector, and/or cell of any preceding claim for any suitable purpose, including but not limited to those disclosed herein such as designing protein logic gates.
- 25
26. A fusion protein comprising a polypeptide of the formula X-B-Z, wherein:
- (a) the X domain is a non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein the X domain is capable of non-covalently binding to a first target;
- 30 (b) the Z domain is a non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein the Z domain is capable of non-covalently binding to either (i) a second target that differs from the first target, or (ii) a different non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices; and
- (c) the B domain is an amino acid linker;

wherein a combined number of alpha helices from the X domain and the Z domain is 4, 5, or 6; and

wherein the X domain and the Z domain interact at a binding interface, wherein the binding interface comprises a hydrogen bond network in which at least one side chain in each alpha helix hydrogen of the X domain bonds with a side chain in an alpha helix in the Z domain, and wherein the binding interface comprises a plurality of hydrophobic residues.

27. A kit or composition, comprising at least two fusion proteins comprising the formula X-B-Z, wherein

10 the B domain in each fusion protein is independently a polypeptide linker;

the X domain in each fusion protein comprises a first non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices;

the Z domain in each fusion protein comprises a second non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein a combined number of alpha helices from the X domain and the Z domain in each individual fusion protein is 4, 5, or 6; wherein the X domain and the Z domain interact at a binding interface, wherein the binding interface comprises a hydrogen bond network in which at least one side chain in each X domain alpha helix bonds with a side chain in an alpha helix in the Z domain; wherein

20 the X domain in a first fusion protein is capable of non-covalently binding to a first target;

the Z domain in a second fusion protein is capable of non-covalently binding to a second target; and

the X domains and Z domains in each individual fusion protein that are not capable of non-covalently binding to the first target or the second target are capable of non-covalently binding to an X or a Z domain of a different fusion protein in the plurality of fusion proteins.

28. The kit or composition of claim 27, wherein

(i) the first fusion protein has the formula X1-B1-Z1, wherein the X1 domain is capable of non-covalently binding to the first target; and

30 (ii) the second fusion protein has the formula X2-B2-Z2, wherein the Z2 domain is capable of non-covalently binding to the second target; and wherein the Z1 and X2 domains are capable of non-covalently binding to each other.

29. The kit or composition of claim 27, wherein:

- (i) the first fusion protein has the formula X1-B1-Z1, wherein the X1 domain is capable of non-covalently binding to the first target; and
- (ii) the second fusion protein has the formula X2-B2-Z2,
- (iii) the at least two fusion proteins comprise a third fusion protein of formula X3-
5 B3-Z3, wherein the Z3 domain is capable of non-covalently binding to the second target;
wherein
- (A) the Z1 and X2 domains are capable of non-covalently binding to each other; and
- (B) the Z2 and X3 domains are capable of non-covalently binding to each
10 other.

30. The fusion protein of claim 26 or the kit or compositions of claims 27-29, wherein the binding interface comprises at least 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60% or greater hydrophobic residues.

15

31. The fusion protein, kit, or composition of any one of claims 26-30, wherein the B domain for each fusion protein is independently between 6-12, 6-11, 6-10, 7-12, 7-11, 7-10, 8-12, 8-11, 8-10, 9-12, 9-11, 9-10, 10-12, 10-11, 11-12, 6, 7, 8, 9, 10, 11, or 12 amino acids in length.

20

32. The fusion protein, kit, or composition of any one of claims 26-31, wherein the combined number of alpha helices from the X and Z domains in an individual fusion protein is 4.

25 33. The fusion protein, kit, or composition of any one of claims 26-32, wherein the X domain of each fusion protein has 2 alpha helices and the Z domain of each fusion protein has 2 alpha helices.

30 34. The fusion protein, kit, or composition of any one of claims 26-33, wherein either the X domain or the Z domain of each fusion protein has 1 alpha helix and the other has 3 alpha helices.

35. The fusion protein, kit, or composition of any one of claims 26-34, wherein each X domain and each Z domain comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%,

80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide comprising the amino acid sequence selected from SEQ ID NO:1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 5 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, with the proviso that the X domain and the Z domain do not do not form a heterodimer (a-b) pair.

36. The fusion protein, kit, or composition of claim 35, wherein at least 20%, 25%, 30%, 10 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of each X domain and each Z domain are invariant compared to the reference amino acid sequence.

15 37. The fusion protein, kit, or composition of any one of claims 26-36, wherein each fusion protein independently comprises a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide having the amino acid sequence selected from the group consisting of SEQ ID 20 NO: 302, 303, 306-326, 439, 441, 443, 445, 447, 449, 451, 453, 455, and 457.

38. The kit or composition of any one of claims 27-37, further comprising the first target and the second target.

25 39. The kit or composition of claim 38, wherein the first target and the second target each independently comprise a polypeptide of the formula X10-B10-Z10, wherein

(a) the X10 domain is a non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices;

(b) the Z10 domain is a non-naturally occurring polypeptide comprising 1, 2, or 3 30 alpha helices; and

(c) the B10 domain is an amino acid linker;

wherein the X domain and the Z domain interact at a target binding interface, wherein the target binding interface comprises a hydrogen bond network in which at least one side chain in each alpha helix hydrogen of the X domain bonds with a side chain in a different

alpha helix in the Z domain, and wherein the target binding interface comprises a plurality of hydrophobic residues.

40. The kit or composition of claim 39, wherein the target binding interface comprises at
5 least 25% hydrophobic residues.

41. The kit, or composition of any one of claims 39-40, wherein the B10 domain for the
first target and the second target is independently between 6-12, 6-11, 6-10, 7-12, 7-11, 7-10,
8-12, 8-11, 8-10, 9-12, 9-11, 9-10, 10-12, 10-11, 11-12, 6, 7, 8, 9, 10, 11, or 12 amino acids
10 in length.

42. The kit, or composition of any one of claims 38-41, wherein the combined number of
alpha helices from the X and Z domains in the first target and the second target protein is 4.

43. The kit, or composition of any one of claims 39-42, wherein
15 (a) the X10 domain of each of the first target and the second target has 2 alpha
helices and the Z10 domain of each of the first target and the second target has 2 alpha
helices; or
(b) either the X10 domain or the Z10 domain of each of the first target and the
20 second target has 1 alpha helix and the other has 3 alpha helices

44. The kit, or composition of any one of claims 39-43, wherein each X10 domain and
each Z10 domain comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%,
85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full
25 length of a polypeptide selected from the group including, but not limited to a polypeptide
comprising the amino acid sequence selected from the group consisting of SEQ ID NO:1-
290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448,
450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486,
488, 490, 493, and 494, with the proviso that the X10 domain forms a heterodimer (a-b) pair
30 with the X domain of the fusion protein, and the Z10 domain forms a heterodimer (a-b) pair
with the Z domain of the fusion protein.

45. The fusion protein, kit, or composition of claim 44, wherein at least 20%, 25%, 30%,
35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%,

95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of each X domain and each Z domain are invariant compared to the reference polypeptide amino acid sequence.

5 46. The kit, or composition of any one of claims 39-45, wherein the first target and/or the second target further comprise one or more effector polypeptide domains linked to one or more of the X10 and/or Z10 domains, for example, wherein the one or more effector polypeptide domains may optionally comprise a polypeptide including, but not limited to, nucleic acid binding proteins, transcription factors, receptor binding proteins, split enzymes,
10 effectors of membrane receptors, etc.

47. Use of the fusion proteins, kits, or compositions of any one of claims 26-46 for any purpose as described herein.

15 48. A method, comprising:

- (i) contacting a fusion protein according to any one of claims 26-37 with a biological sample under conditions to promote non-covalent binding of the fusion protein with first target and second target present in the sample, and
- (ii) detecting non-covalent binding of the one or more fusion proteins to the first
20 target and/or the second target in the biological sample.

49. The method of claim 48, wherein the method comprises detecting cooperative non-covalently binding of the one or more fusion proteins to the first target and the second target in the biological sample.

25

50. The method of claim 48, wherein the method comprises detecting non-covalent binding of the one or more fusion proteins to the first target or the second target in the biological sample.

30 51. A method for target detection, comprising

- (a) contacting a biological sample with at least two fusion proteins, wherein each of the at least two fusion proteins comprises the formula X-B-Z, wherein each B is independently a polypeptide linker;

each X domain comprises a first non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices;

each Z domain comprises a second non-naturally occurring polypeptide comprising 1, 2, or 3 alpha helices, wherein a combined number of alpha helices from the X domain and the Z domain in each individual fusion protein is 4, 5, or 6; wherein the X domain and the Z domain interact at a binding interface, wherein the binding interface comprises a hydrogen bond network in which at least one side chain in each X domain alpha helix bonds with a side chain in an alpha helix in the Z domain; wherein

the X domain in a first fusion protein is capable of non-covalently binding to a first target;

the Z domain in a second fusion protein is capable of non-covalently binding to a second target; and

the X domains and Z domains in each individual fusion protein that are not capable of non-covalently binding to the first target or the second target are capable of non-covalently binding to an X or a Z domain of a different fusion protein in the plurality of fusion proteins;

(b) detecting non-covalent binding of the two or more fusion proteins to the first target and/or the second target in the biological sample.

52. The method of claim 51, wherein the detecting comprises detecting cooperative non-covalent binding of the two or more fusion proteins to the first target and the second target in the biological sample.

53. The method of claim 51 or 52, wherein:

(i) the first fusion protein has the formula X1-B1-Z1, wherein the X1 domain is capable of non-covalently binding to the first target; and

(ii) the second fusion protein has the formula X2-B2-Z2, wherein the Z2 domain is capable of non-covalently binding to the first target; and wherein the Z1 and X2 domains are capable of non-covalently binding to each other.

54. The method of claim 51 or 52, wherein:

(i) the first fusion protein has the formula X1-B1-Z1, wherein the X1 domain is capable of non-covalently binding to the first target; and

(ii) the second fusion protein has the formula X2-B2-Z2,

(iii) the at least two fusion proteins comprise a third fusion protein of formula X3-B3-Z3, wherein the Z3 domain is capable of non-covalently binding to the second target; wherein

- 5 (A) the Z1 and X2 domains are capable of non-covalently binding to each other; and
- (B) the Z2 and X3 domains are capable of non-covalently binding to each other.

55. The method of any one of claims 51-54, wherein the X domains, Y domains, B
10 domains, and or fusion proteins are as recited in any one of claims 26-37.

56. The method of any one of claims 48-55, wherein at least one of the fusion proteins comprises one or more effector polypeptide domains linked to one or more of the X and/or Z domains, and wherein the detecting step comprises detecting an output signal caused by
15 binding the first target and/or the second target.

57. The method of claim 56, wherein the detecting step comprises detecting an output signal from the one or more effector polypeptide caused by cooperative non-covalently binding of the first target and the second target.
20

58. The method of claim 56 or 57, wherein the one or more effector polypeptide domains may comprise a polypeptide including, but not limited to, nucleic acid binding proteins, transcription factors, receptor binding proteins, nucleic acid binding proteins, transcription factors, receptor binding proteins, split enzymes, effectors of membrane receptors, etc..
25

59. The method of any one of claims 56-58, wherein the output signal may include, but is not limited to fluorescence activity, functional activity, etc.

60. The fusion protein, kit or composition of any one of claims 26-46, or the method of
30 any one of claims 48-59, wherein the first target and/or the second target comprise polypeptides or nucleic acids.

61. A composition comprising

(a) a first polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding a first target; and

(b) a second polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding to the second polypeptide, and wherein the second

5 polypeptide is capable of non-covalently binding a second target that differs from the first target; wherein:

(i) a binding affinity of the first polypeptide for the first target is approximately equal to a binding affinity of the second polypeptide for the second target; and

(ii) the binding affinity of the first polypeptide for the first target and the
10 binding affinity of the second polypeptide for the second target are greater than the binding affinity of the first target and the second target for each other.

62. The composition of claim 61, further comprising the first target and the second target.

15 63. The composition of claim 62, wherein the first target and/or the second target further comprise one or more effector polypeptide domains.

64. The composition of claim 63, wherein the one or more effector polypeptide domains may comprise a polypeptide including, but not limited to, nucleic acid binding proteins,
20 transcription factors, receptor binding proteins, split enzymes, effectors of membrane receptors, etc.

65. The composition of any one of claims 61-64, wherein the first polypeptide and/or the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%,
25 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide comprising the amino acid sequence selected from the group consisting of SEQ ID NO:1-
290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448,
450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486,
30 488, 490, 493, and 494.

66. The composition of claim 65, wherein at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of the first

polypeptide and/or the second polypeptide are invariant compared to the reference polypeptide amino acid sequence

67. The composition of any one of claims 61-66, wherein

5 (a) the first polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to the amino acid sequence of SEQ ID NO:3; and

10 (b) the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to the amino acid sequence of SEQ ID NO:58.

68. The composition of any one of claims 61-67, wherein the first target and/or the
15 second target each comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide comprising the amino acid sequence selected from the group consisting of SEQ ID NO:1-
20 290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, with the proviso that the first target forms a heterodimer (a-b) pair with the first polypeptide, and the second target forms a heterodimer (a-b) pair with the second polypeptide.

25 69. The composition of claim 46, wherein at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of the first target and/or the second target are invariant compared to the reference polypeptide amino acid sequence.

30

70. A method, comprising:

(a) contacting a biological sample with the composition of any one of claims 61-69; and

(b) detecting binding, of the first polypeptide to the first target and binding of the second polypeptide to the second target in the sample, such as detecting an output signal caused by actions of effector polypeptides upon binding.

5

71. A composition comprising

(a) a first polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding a first target; and

10 (b) a second polypeptide comprising 2 alpha helices, wherein the first polypeptide is capable of non-covalently binding to the second polypeptide, and wherein the second polypeptide is capable of non-covalently binding a second target that differs from the first target; wherein:

(i) a binding affinity of the first polypeptide for the second polypeptide is greater than a binding affinity of the second polypeptide for the second target;

15 (ii) a binding affinity of the first polypeptide for the first target is approximately equal to a binding affinity of the second polypeptide for the second target; and

(iii) the binding affinity of the first polypeptide for the first target and the binding affinity of the second polypeptide for the second target are greater than the binding affinity of the first target and the second target for each other.

20

72. The composition of claim 71, further comprising the first target and the second target.

73. The composition of claim 72, wherein the first target and/or the second target further comprise one or more effector polypeptide domains.

25

74. The composition of claim 73, wherein the one or more effector polypeptide domains may comprise a polypeptide including, but not limited to, nucleic acid binding proteins, transcription factors, receptor binding proteins, split enzymes, effectors of membrane receptors, etc.

30

75. The composition of any one of claims 71-74, wherein the first polypeptide and/or the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide

comprising the amino acid sequence selected from the group consisting of SEQ ID NO:1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494.

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76. The composition of claim 75, wherein at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of the first polypeptide and/or the second polypeptide are invariant compared to the reference polypeptide amino acid sequence.

10

77. The composition of any one of claims 71-76, wherein

(a) the first polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100%

15

identical to the full length of a polypeptide selected from the group including, but not limited the amino acid sequence of SEQ ID NO:3; and

(b) the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to the amino acid sequence of SEQ ID NO:42.

20

78. The composition of any one of claims 71-77, wherein the first target and/or the second target each comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full length of a polypeptide selected from the group including, but not limited to a polypeptide comprising the amino acid sequence selected from the group consisting of SEQ ID NO:1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, with the proviso that the first target forms a heterodimer (a-b) pair with the first polypeptide, and the second target forms a heterodimer (a-b) pair with the second polypeptide.

25

30

79. The composition of claim 78, wherein at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%,

98%, 99%, or 100% of amino acid residues at defined interface positions of the first target and/or the second target are invariant compared to the reference polypeptide amino acid sequence.

5 80. A method, comprising:
 (a) contacting a biological sample with the composition of any one of claims 71-79; and
 (b) detecting binding interactions between the first polypeptide and the first target.
the second polypeptide and the second target, the first polypeptide and the second
10 polypeptide, and the first target and the second target in the sample, such as detecting an output signal caused by actions of effector polypeptides upon binding.

81. A composition comprising
 (a) a first polypeptide comprising 4 alpha helices, wherein the first polypeptide is
15 capable of non-covalently binding a first target; and
 (b) a second polypeptide comprising 4 alpha helices, wherein the second polypeptide is capable of non-covalently binding a second target that differs from the first target; wherein:
 (i) a binding affinity of the first target for the second target is greater than
20 a binding affinity of the first polypeptide for the first target;
 (ii) a binding affinity of the first polypeptide for the first target is approximately equal to a binding affinity of the second polypeptide for the second target; and
 (iii) the sum of the binding affinity of (A) the first polypeptide for the first target and (B) the binding affinity of the second polypeptide for the second target, is greater
25 than the binding affinity of the first target and the second target.

82. The composition of claim 81, further comprising the first target and the second target.

83. The composition of claim 82, wherein the first target and/or the second target further
30 comprise one or more effector polypeptide domains.

84. The composition of claim 83, wherein the one or more effector polypeptide domains may comprise a polypeptide including, but not limited to, nucleic acid binding proteins,

transcription factors, receptor binding proteins, split enzymes, effectors of membrane receptors, etc.

85. The composition of any one of claims 81-84, wherein the first polypeptide and/or the
5 second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%,
85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full
length of a polypeptide selected from the group including, but not limited to a polypeptide
comprising the amino acid sequence selected from the group consisting of SEQ ID NO:1-
290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448,
10 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486,
488, 490, 493, and 494.

86. The composition of claim 85, wherein at least 20%, 25%, 30%, 35%, 40%, 45%,
50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%,
15 98%, 99%, or 100% of amino acid residues at defined interface positions of the first
polypeptide and/or the second polypeptide are invariant compared to the reference
polypeptide amino acid sequence.

87. The composition of any one of claims 81-86, wherein
20 (a) the first polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%,
70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100%
identical to the full length of a polypeptide selected from the group including, but not limited
the amino acid sequence of SEQ ID NO: 42; and
(b) the second polypeptide comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%,
25 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical
to the full length of a polypeptide selected from the group including, but not limited to the
amino acid sequence of SEQ ID NO:57.

88. The composition of any one of claims 81-87, wherein the first target and/or the
30 second target each comprise a polypeptide that is 50%, 55%, 60%, 65%, 70%, 75%, 80%,
85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% identical to the full
length of a polypeptide selected from the group including, but not limited to a polypeptide
comprising the amino acid sequence selected from the group consisting of SEQ ID NO:1-
290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448,

450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, with the proviso that the first target forms a heterodimer (a-b) pair with the first polypeptide, and the second target forms a heterodimer (a-b) pair with the second polypeptide.

5

89. The composition of claim 88, wherein at least 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of amino acid residues at defined interface positions of the first target and/or the second target are invariant compared to the reference polypeptide amino acid sequence.

10

90. A method, comprising:

(a) contacting a biological sample with the composition of any one of claims 81-88; and

15

(b) detecting binding interactions between the first polypeptide and the first target, the second polypeptide and the second target, and the first target and the second target in the sample, such as detecting an output signal caused by actions of effector polypeptides upon binding.

20

91. A nucleic acid encoding the fusion protein of any one of claims 26 and 30-37.

92. An expression vector comprising the nucleic acid of claim 91 operatively linked to a promoter.

25

93. A host cell comprising the nucleic acid of claim 91 or the expression vector of claim 92.

94. The host cell of claim 93, wherein the nucleic acid or the expression vector is integrated into a host cell chromosome.

30

95. The host cell of claim 93, wherein the nucleic acid or the expression vector is episomal.

96. A heterodimer, comprising:

(a) a monomer A polypeptide, wherein the monomer A polypeptide is a non-naturally occurring polypeptide comprising 1-5 alpha helices, wherein adjacent alpha helices may optionally be connected by an amino acid linker; and

5 (b) a monomer B polypeptide, wherein the monomer B polypeptide is a non-naturally occurring polypeptide comprising 1-5 alpha helices, wherein adjacent alpha helices may optionally be connected by an amino acid linker;

wherein the monomer A polypeptide and the monomer B polypeptide non-covalently interact to form the designed heterodimer protein.

10 97. The heterodimer of claim 96 wherein the monomer A polypeptide and the monomer B polypeptide have their interaction specificity determined by at least one hydrogen bond network at the interface between the monomer A polypeptide and the monomer B polypeptide.

15 98. The heterodimer of claim 96 or 97, wherein

(i) the monomer A polypeptide comprises 2 alpha helices, and the monomer B polypeptide comprises 3 alpha helices;

(ii) the monomer A polypeptide comprises 3 alpha helices and the monomer B polypeptide comprises 3 alpha helices;

20 (iii) the monomer A polypeptide comprises 3 alpha helices and the monomer B polypeptide comprises 4 alpha helices,

(iv) the monomer A polypeptide comprises 4 alpha helices and the monomer B polypeptide 3 alpha helices;

25 (v) the monomer A polypeptide comprises 4 alpha helices and the monomer B polypeptide comprises 4 alpha helices;

(vi) the monomer A polypeptide comprises 5 alpha helices and the monomer B polypeptide comprises 4 alpha helices;

(vii) the monomer A polypeptide comprises 4 alpha helices and the monomer B polypeptide comprises 5 alpha helices;

30 (viii) the monomer A polypeptide comprises 5 alpha helices and the monomer B polypeptide comprises 5 alpha helices;

(ix) the monomer A polypeptide comprises 2 alpha helices and the monomer B polypeptide comprises 2 alpha helices; or

(x) the monomer A polypeptide comprises 3 alpha helices and the monomer B polypeptide comprises 2 alpha helices.

99. The heterodimer of any one of claims 96-98, wherein:

5 (i) the monomer A polypeptide comprises the amino acid sequence having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an odd-numbered SEQ ID NO selected from the group consisting of selected from the group SEQ ID NOS: 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436,
10 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494; and

(ii) the monomer B polypeptide the amino acid sequence having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of an even-
15 numbered SEQ ID NO selected from the group consisting of selected from the group SEQ ID NOS: 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494, wherein the even-numbered SEQ ID NO is the binding partner of the odd-numbered SEQ ID NO. in step (i).

20

100. The heterodimer of claim 99, wherein amino acid changes from the reference amino acid sequence are conservative amino acid substitutions.

101. The heterodimer of claim 99 or 100, wherein amino acid residues at 20%, 25%,
25 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of defined interface positions are invariant compared to the reference amino acid sequence.

102. A protein scaffold, comprising

30 (a) a fusion protein comprising of 2, 3, 4, or more polypeptides, wherein each polypeptide present in the fusion protein is a non-naturally occurring polypeptide comprises 1-5 alpha helices, wherein adjacent alpha helices may optionally be connected by an amino acid linker;

wherein each polypeptide in the fusion protein is capable of non-covalently interacting with a binding partner, and wherein the fusion protein does not comprise a binding partner for any polypeptide present in the fusion protein; and

5 (b) a binding partner for at least one of the polypeptides present in the fusion protein;

wherein the fusion protein and the binding partner non-covalently interact to form the protein scaffold, wherein an interaction specificity between the binding partner and the at least polypeptide in the fusion protein are determined by at least one hydrogen bond network at the interface between the binding partner and the at least one
10 polypeptide.

103. The protein scaffold of claim 102, wherein the binding partner for at least one of the polypeptides present in the fusion protein comprises a binding partner for each polypeptide present in the fusion protein.

15

104. The protein scaffold of claim 102 or 103, wherein the fusion protein comprises at least 3 or 4 polypeptides in total.

105. The protein scaffold of any one of claims 102-104, wherein each polypeptide in
20 the fusion protein are different polypeptides.

106. The protein scaffold of any one of claims 102-105, wherein the fusion protein comprises the protein of any one of claims 20-26, and the binding partner comprises a plurality of individual polypeptides of any one of claims 10-12.

25

107. The protein scaffold of any one of claims 102-106, wherein

(i) the fusion protein comprises 2, 3, 4, or more polypeptide having at least 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence of SEQ ID NOS:
30 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494; and

(ii) the binding partner comprises a binding partner as defined herein for each polypeptide in (i), wherein each binding partner has at least 70%, 75%, 80%, 85%,

90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity along the length of the amino acid sequence selected from the group SEQ ID NOS: 1-290, 331, 332, 334, 336-422, 424, 426, 428, 430, 432, 434, 436, 438, 440, 442, 444, 446, 448, 450, 452, 454, 456, 458-460, 462, 464, 466, 468, 470, 472, 474, 476, 478, 480, 482, 484, 486, 488, 490, 493, and 494.

108. The protein scaffold of claim 107, wherein amino acid changes in the fusion protein and the binding partner from the reference amino acid sequence are conservative amino acid substitutions.

10

109. The protein scaffold of claim 107 or 108, wherein amino acid residues at 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% of defined interface positions in the polypeptides in the fusion protein and the binding partner are invariant compared to the reference amino acid sequence.

15

110. The protein scaffold any one of claims 102-109, wherein the at least one least one hydrogen bond network is assymmetric.

111. The protein scaffold any one of claims 102-111, wherein the binding interface comprises at least 25% hydrophobic residues..

20

112. The protein scaffold any one of claims 102-111, wherein the scaffold is stable up to 95°C and has a guanidine denaturation midpoint of 4 M.

25

113. The heterodimer protein of any one of claims 96 to 101 or protein scaffold of any one of claims 102 to 112, wherein the amino acid linker comprises (GSEGS)_n (SEQ ID NO:423), wherein n is between 1 and 10.

114. The heterodimer protein or protein scaffold of claim 113, wherein the amino acid linker comprises GSEGS₂GSEGS₂ (SEQ ID NO:429) or GSEGS₂GSEGS₂GGS (SEQ ID NO:461).

30

115. The protein scaffold of any one of claims 102 to 114, wherein the hydrogen bond network comprises one or more hydrogen bonds.

116. The protein scaffold of any one of claims 102 to 115, wherein the hydrogen
5 bond network is formed between the interface residues according to Table 2.

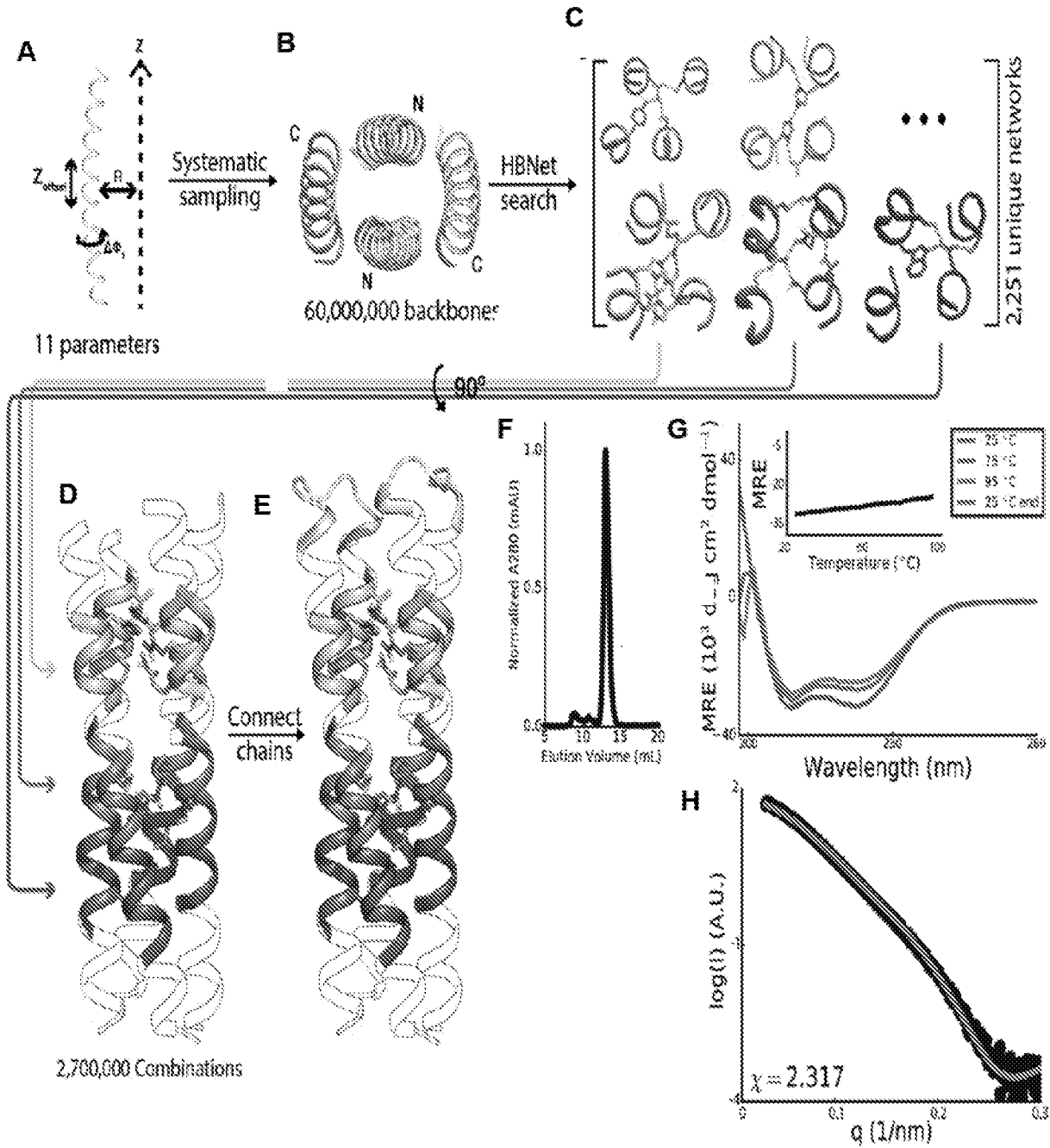


FIGURE 1A – 1H

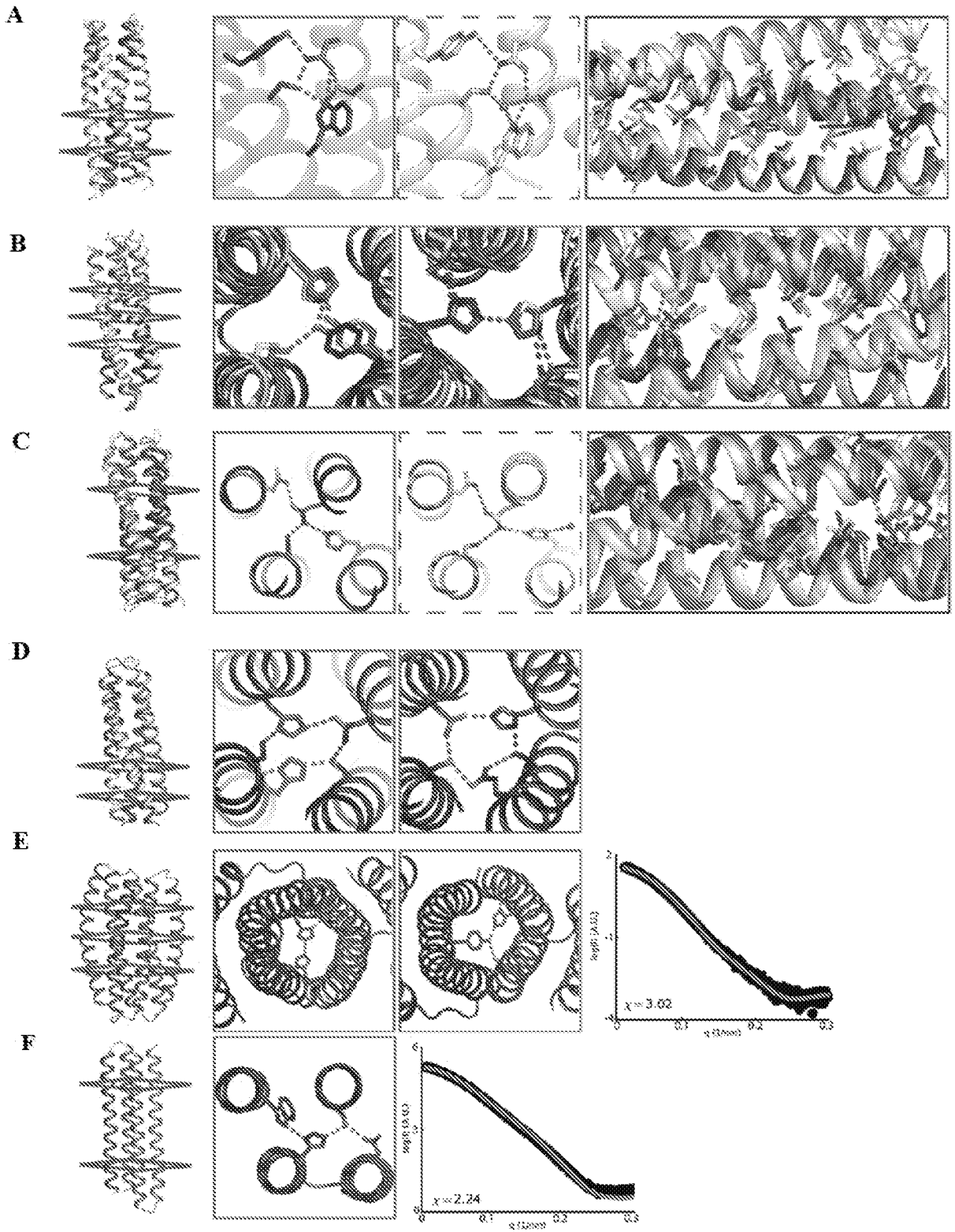


FIGURE 2A – 2F

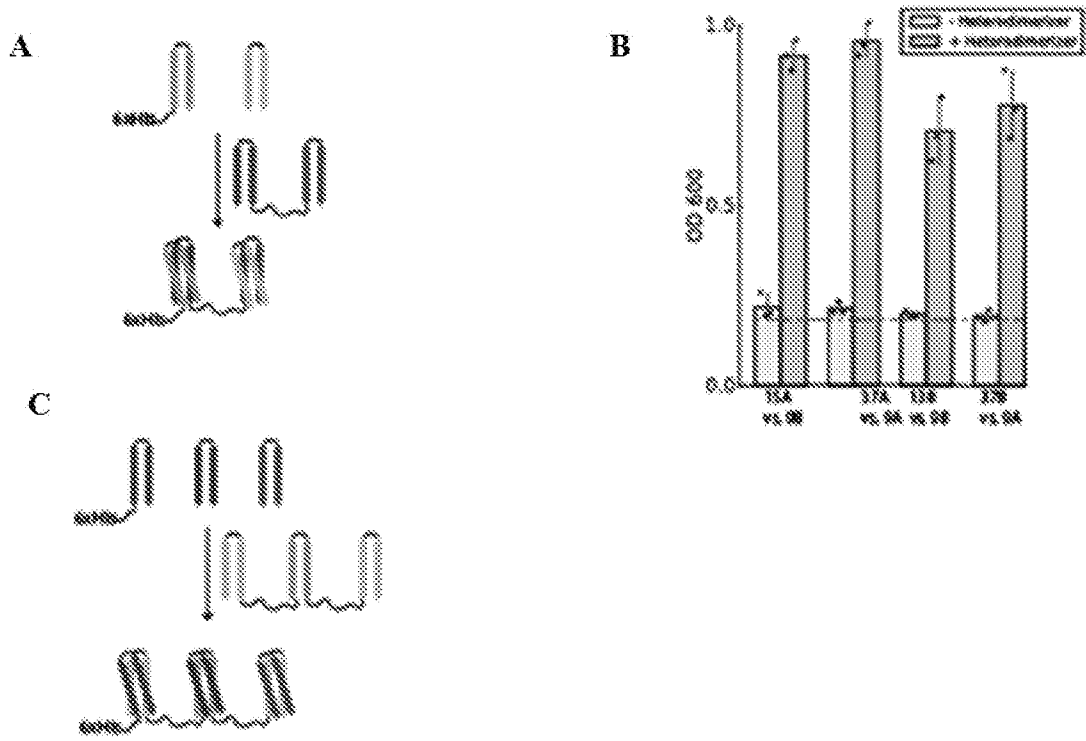


FIGURE 3A – 3C

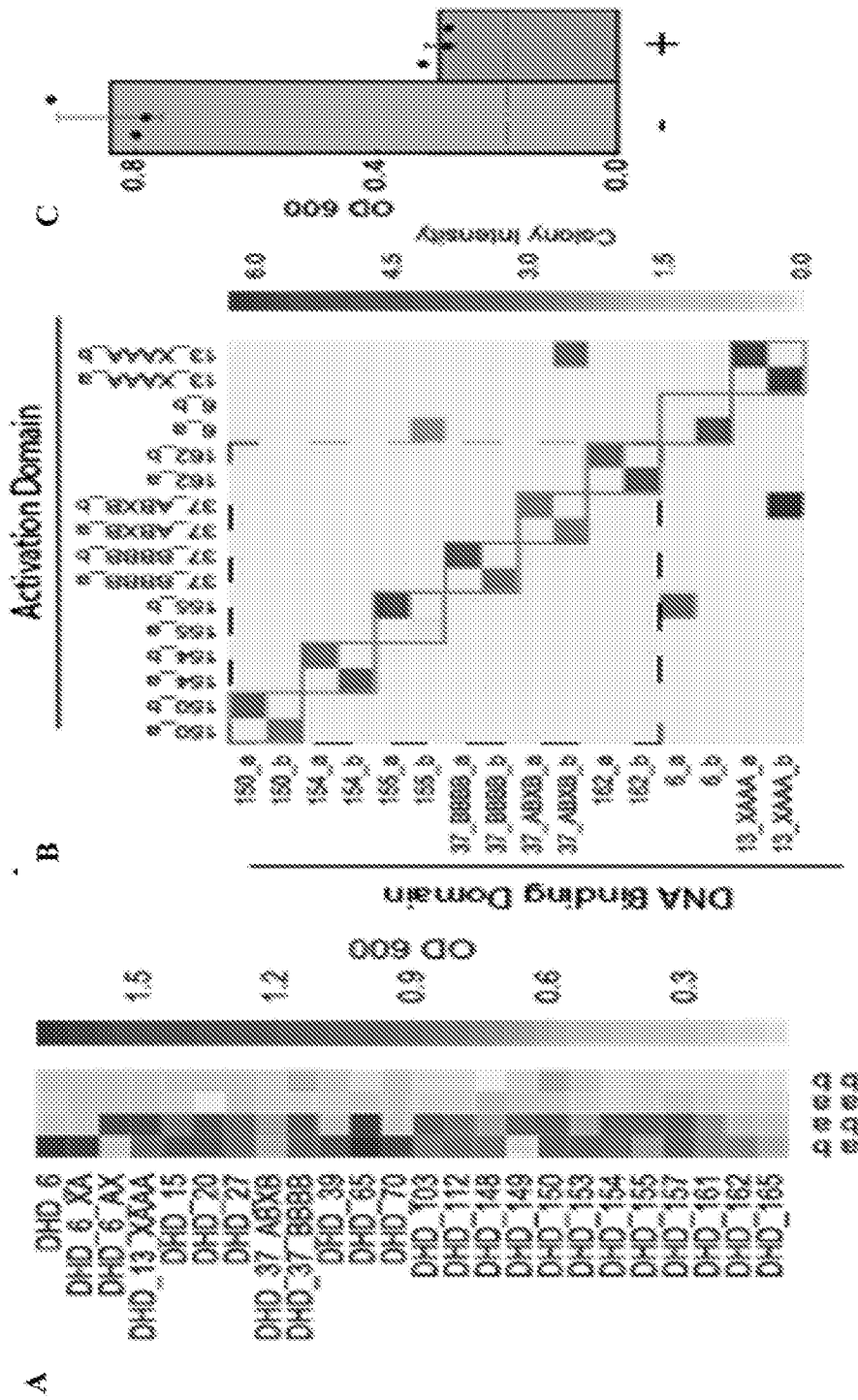
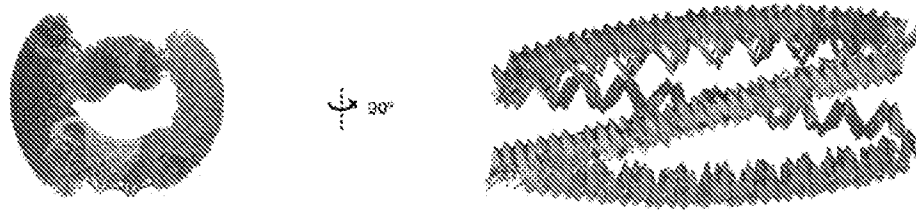


FIGURE 4A - 4C

A



B

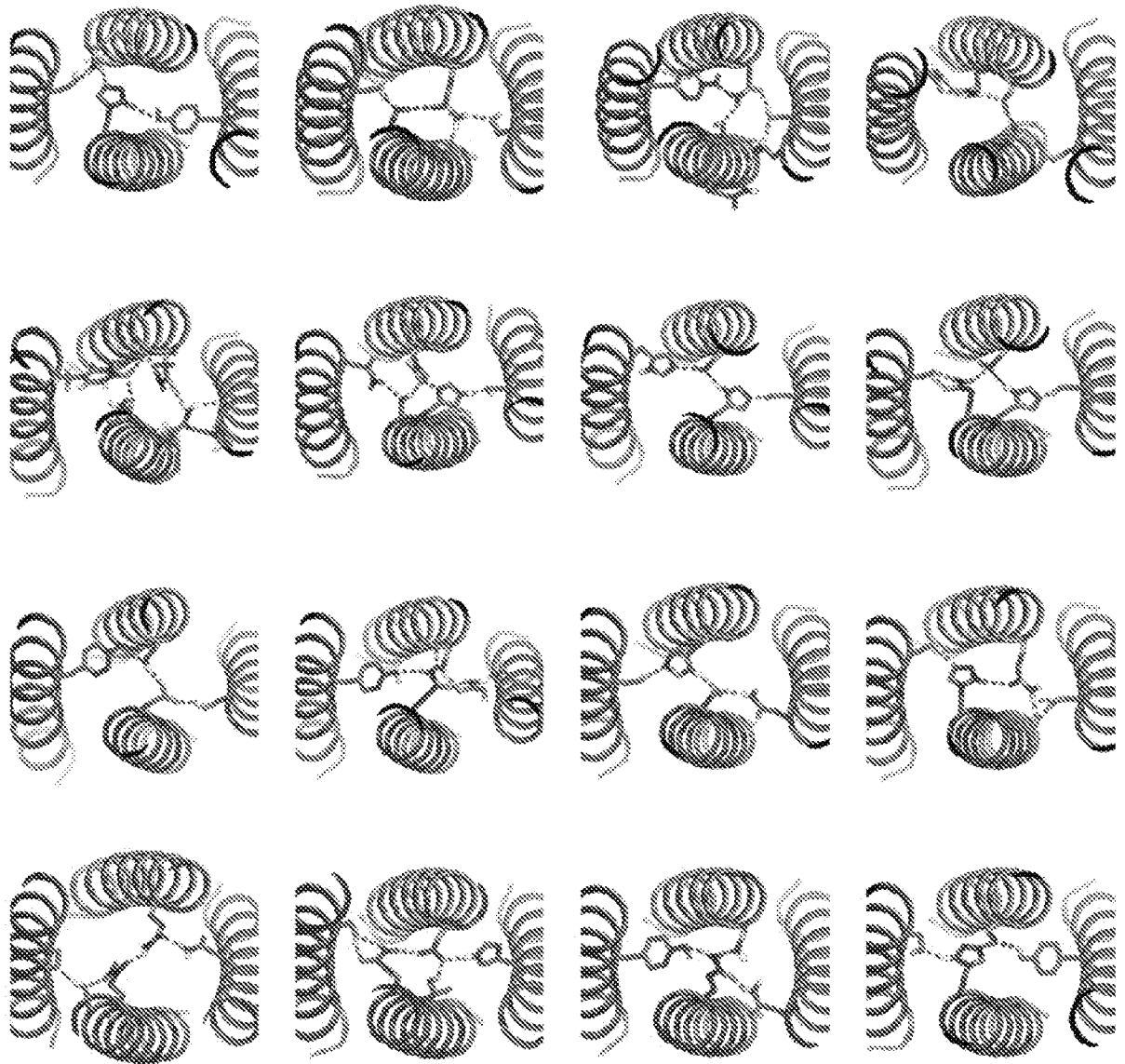
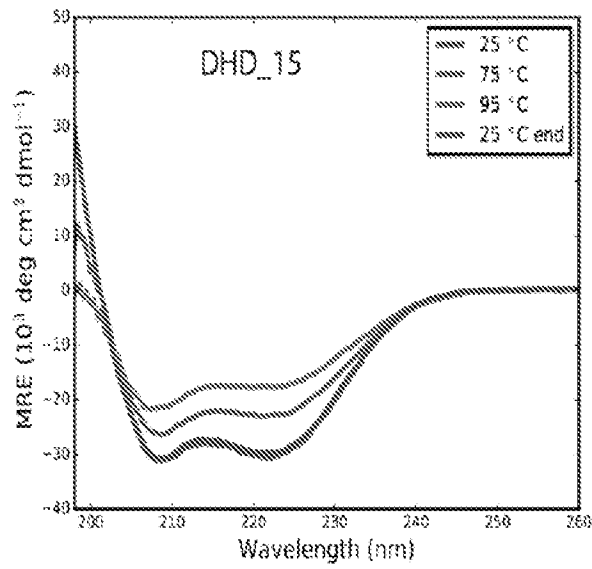
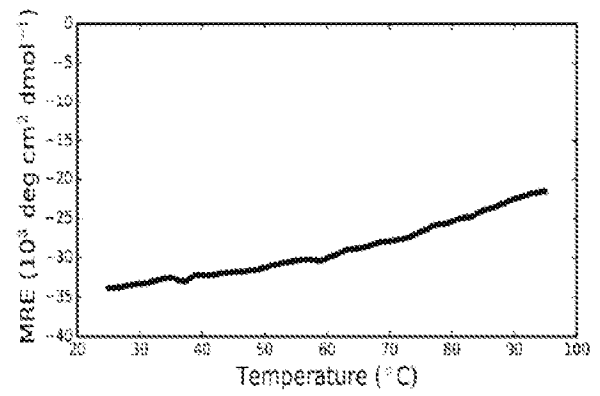
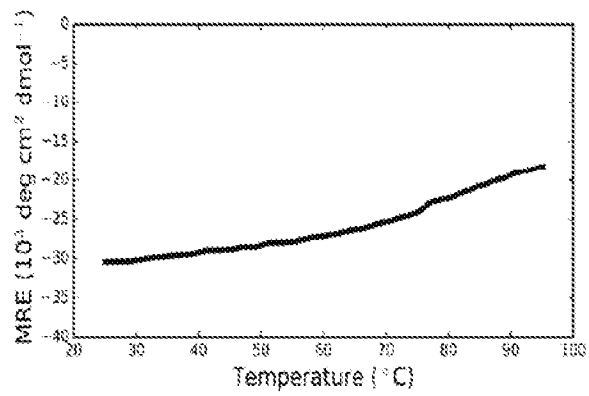
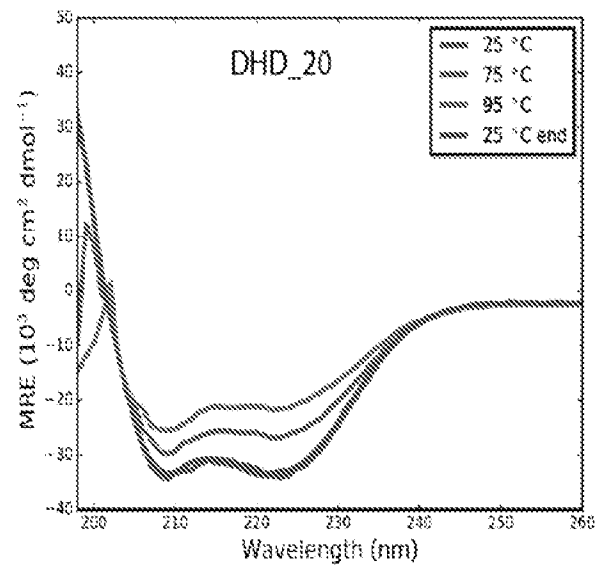


FIGURE 5A - 5B

A



B



C

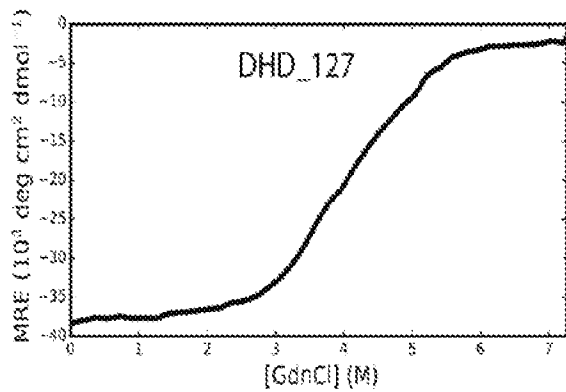


FIGURE 6A – 6C

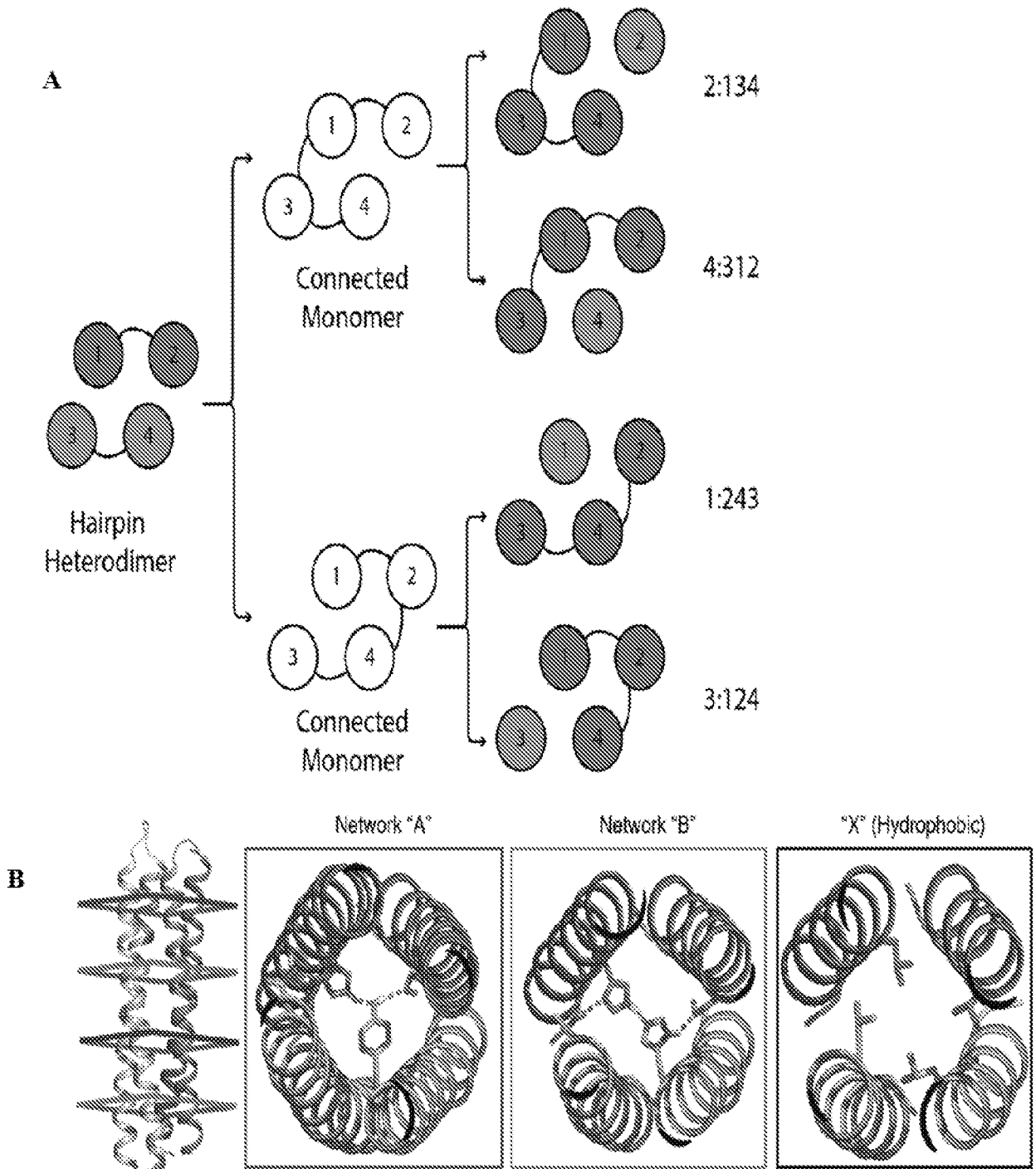


FIGURE 7A – 7B

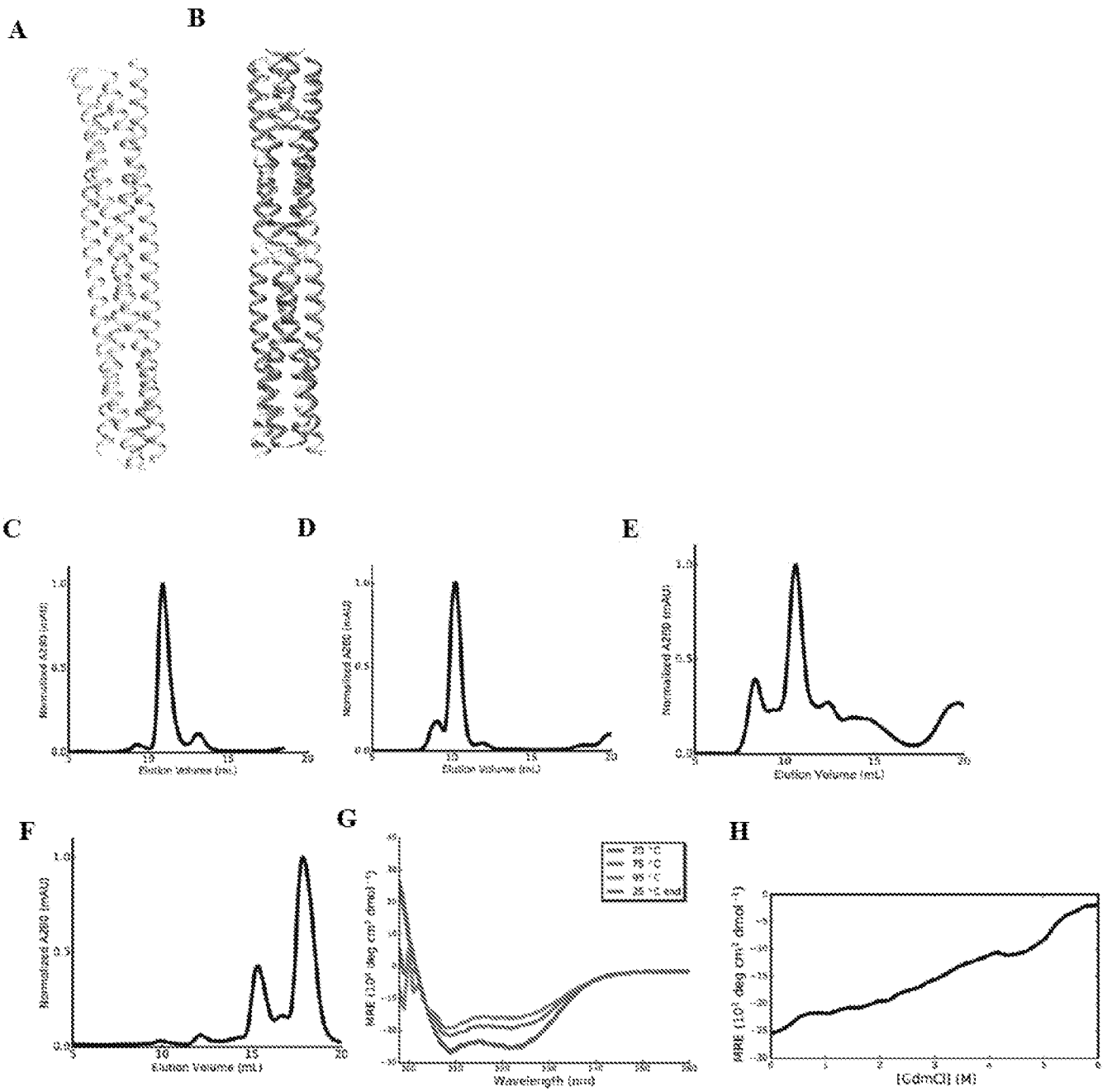


FIGURE 8A – 8H

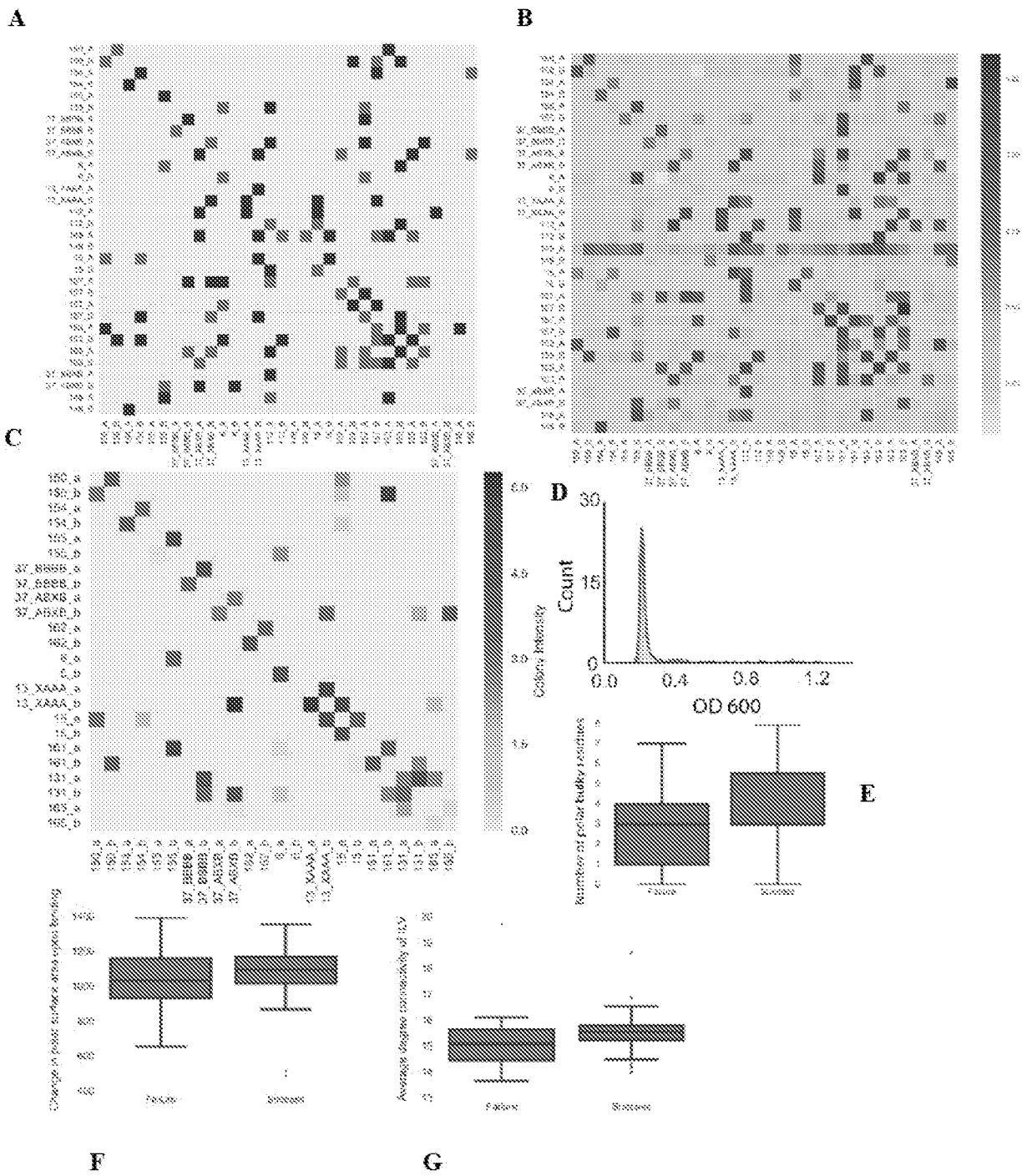


FIGURE 9A – 9G

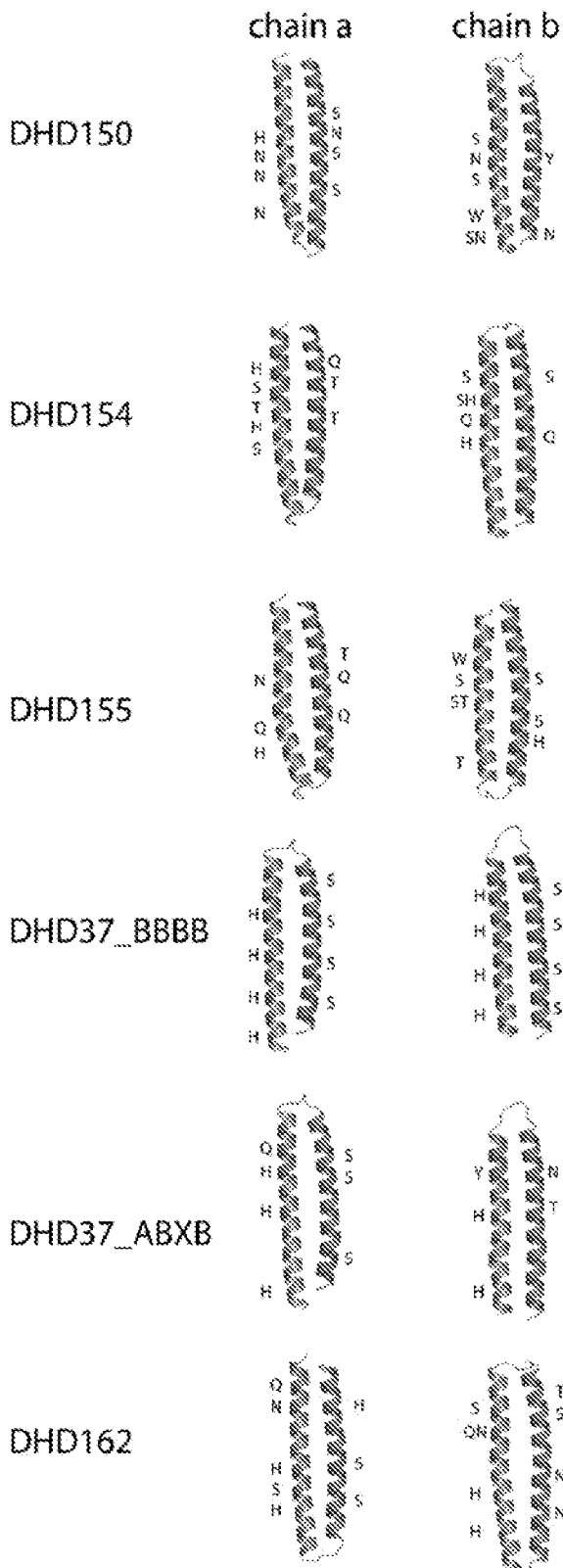


FIGURE 10

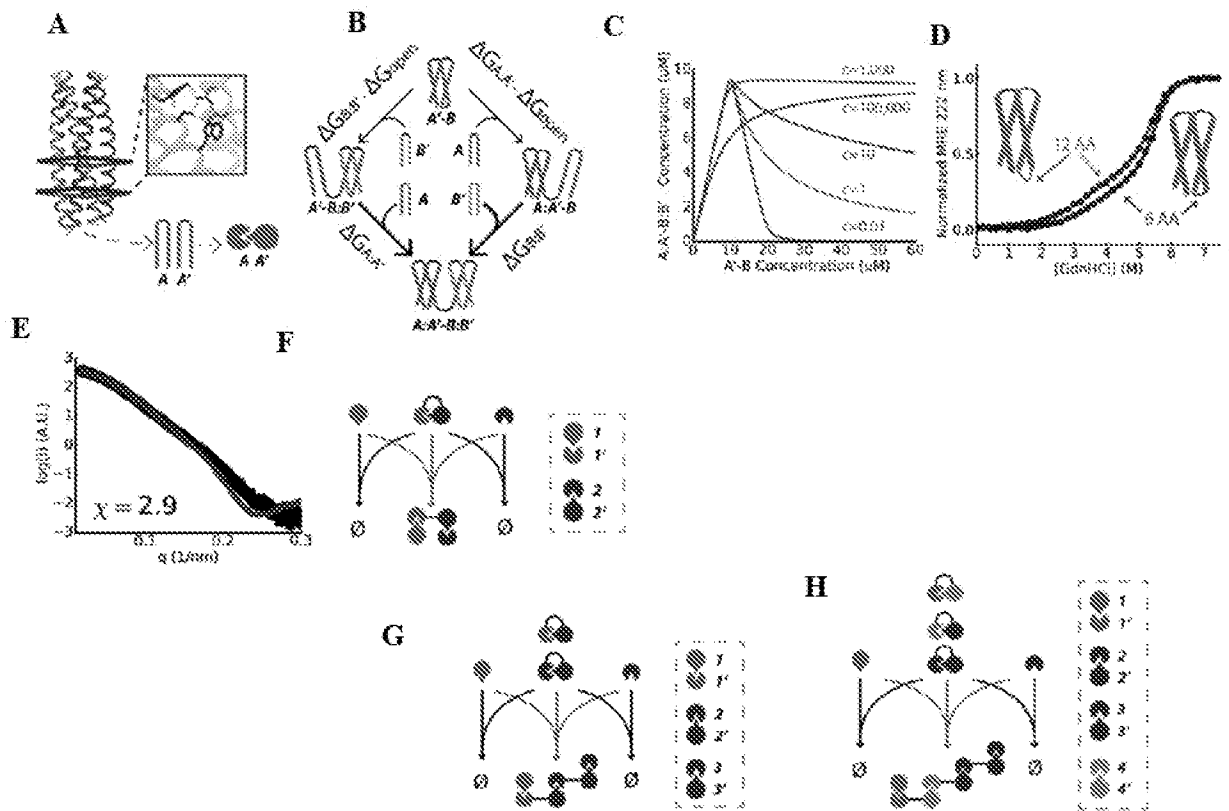


FIGURE 11A – 11H

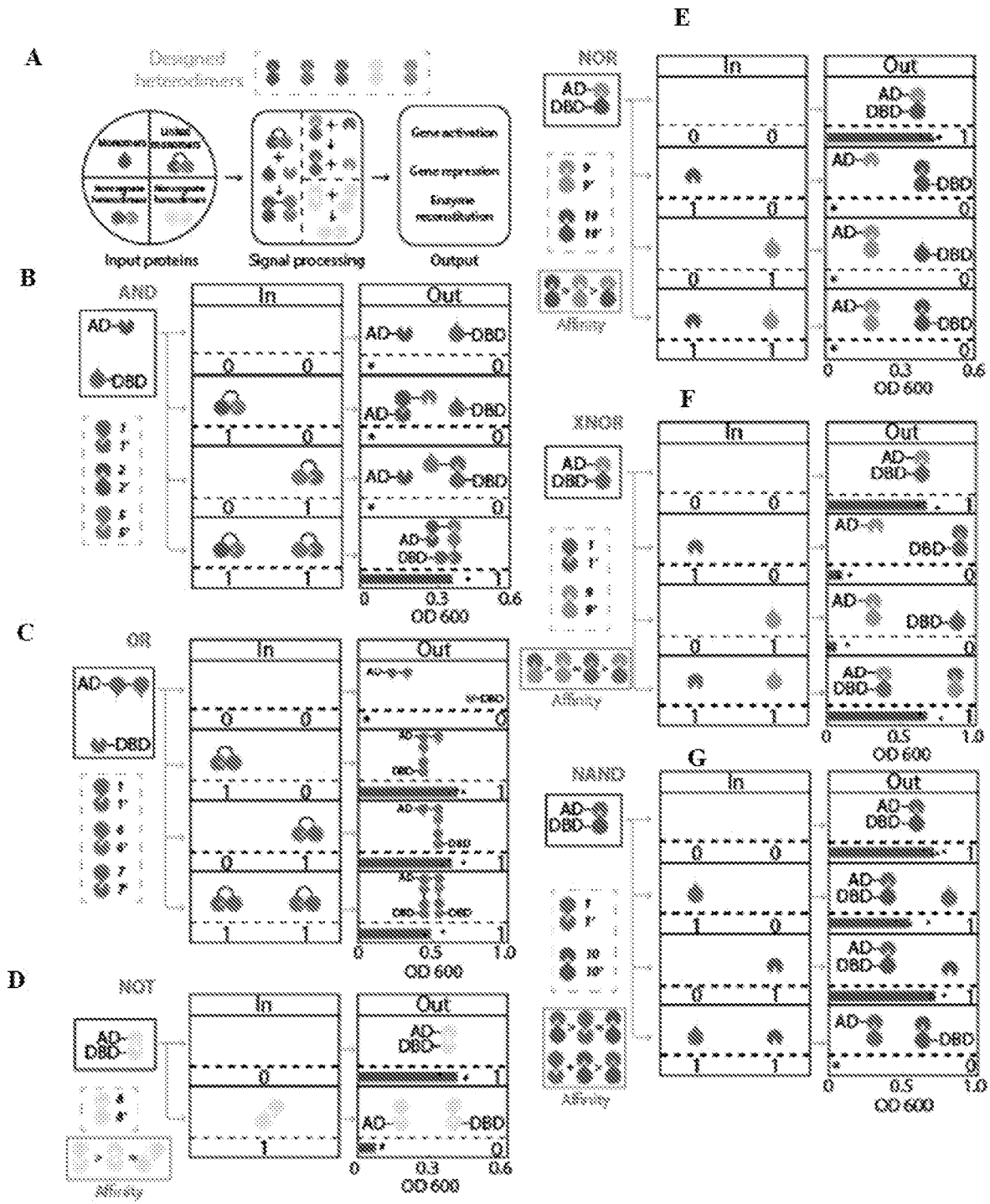


FIGURE 12A - 12G

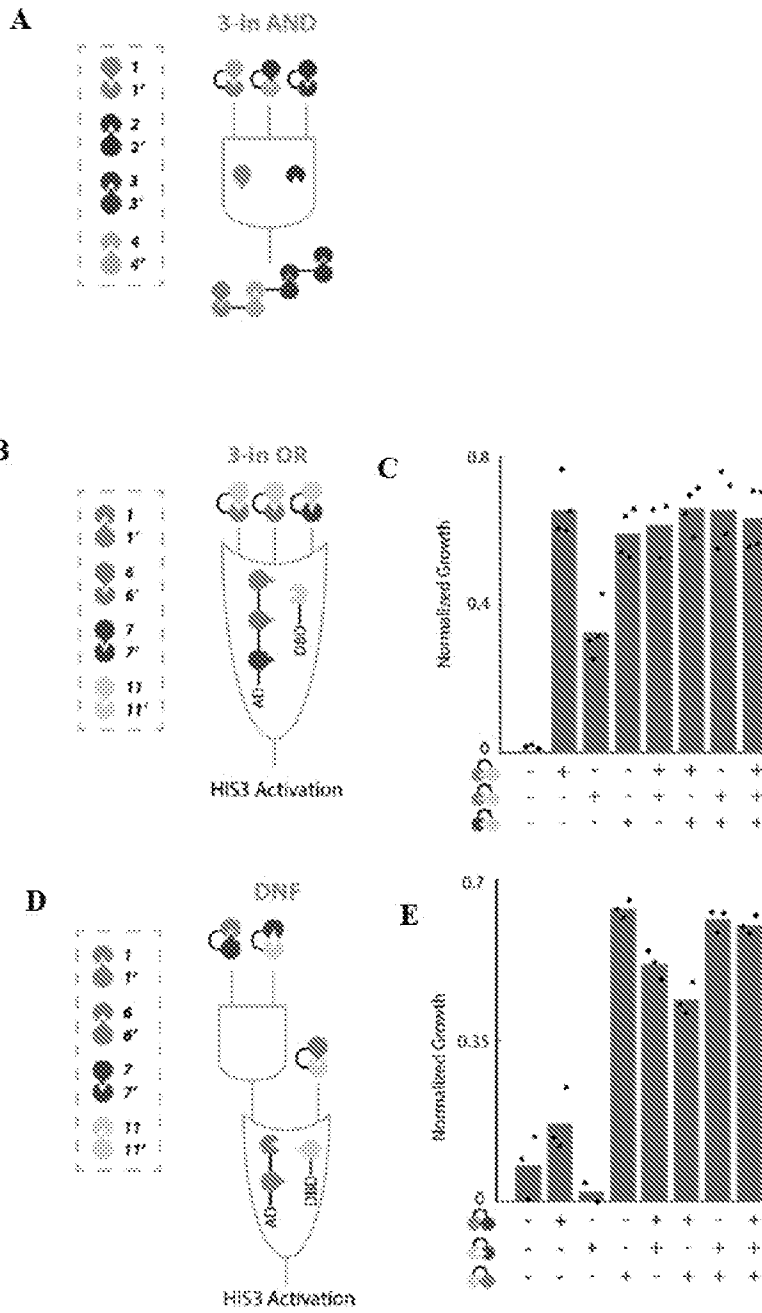


FIGURE 13A – 13E

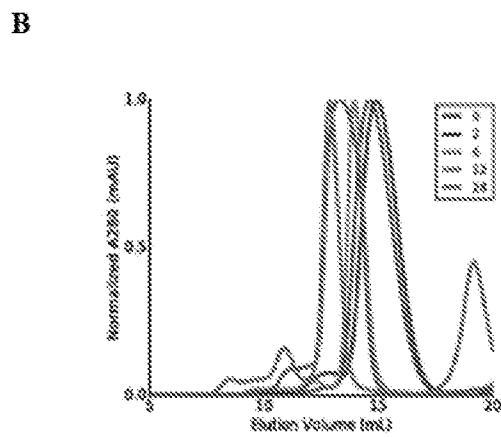
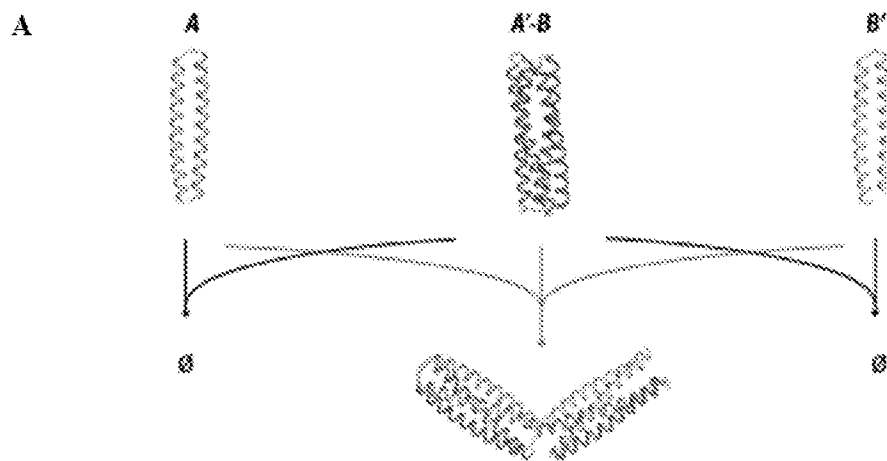


FIGURE 14A – 14B

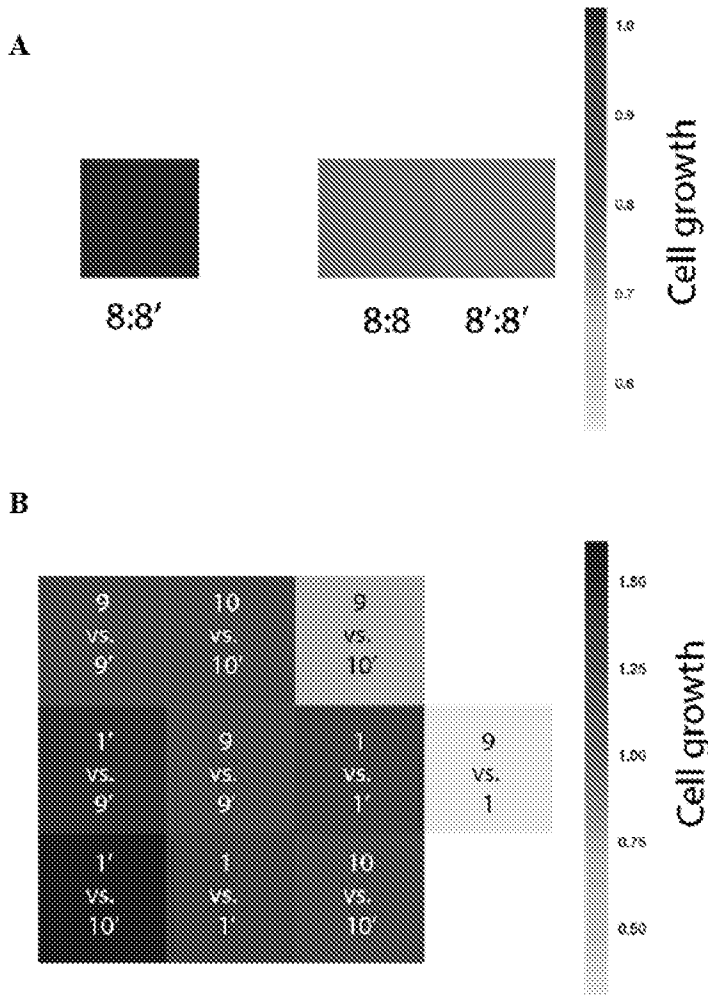


FIGURE 15A – 15B

Monomer A SEQ ID No.	Monomer B SEQ ID NO:	Monomer A SEQ ID No.	Monomer B SEQ ID NO:	Monomer A SEQ ID No.	Monomer B SEQ ID NO:
1	2	55	56	99	100
331	2	55	360	99	402
3	4	57	58	101	102
5	6	57	362	341	404
5	332	59	60	103	104
7	8	59	364	103	406
7	334	61	62	105	106
9	10	61	366	343	408
11	12	63	64	107	108
13	14	65	66	107	410
13	336	65	368	109	110
15	16	67	68	109	412
15	338	67	370	111	112
17	18	69	70	111	424
19	20	69	372	113	114
21	22	71	72	113	416
23	24	71	374	115	116
25	26	73	74	459	420
25	340	73	376	117	118
27	28	75	76	345	422
29	30	75	378	119	120
29	342	77	78	347	424
31	32	77	380	121	122
31	344	79	80	349	426
33	34	79	382	123	124
33	346	81	82	351	428
35	36	337	384	125	126
35	348	83	84	353	126
37	38	339	386	127	128
37	418	85	86	355	430
39	40	85	388	129	130
39	350	87	88	357	432
41	42	87	390	131	132
41	352	89	90	359	434
43	44	89	392	133	134
45	46	91	92	361	436
45	354	91	394	135	136
47	48	93	94	363	438
47	356	93	396	137	138
49	50	95	96	365	440
51	52	95	398	139	140
53	54	97	98	367	442
53	358	97	400	141	142

FIGURE 16

SUBSTITUTE SHEET (RULE 26)

Monomer A SEQ ID No.	Monomer B SEQ ID NO:	Monomer A SEQ ID No.	Monomer B SEQ ID NO:	Monomer A SEQ ID No.	Monomer B SEQ ID NO:
369	444	411	484	259	260
143	144	185	186	261	262
		413	486	263	264
371	446	187	188	265	266
145	146	415	488	267	268
373	448	189	190	269	270
147	148	417	490	271	272
375	450	191	192	273	274
149	150	419	492	275	276
377	452	193	194	277	278
151	152	421	494	279	280
379	454	195	196	281	282
153	154	197	198	283	284
381	456	199	200	285	286
155	156	201	202	287	288
383	458	203	204	289	290
157	158	205	206		
385	460	207	208		
159	160	209	210		
387	462	211	212		
161	162	213	214		
389	464	215	216		
163	164	217	218		
391	466	219	220		
165	166	221	222		
393	468	223	224		
167	168	225	226		
395	470	227	228		
169	170	229	230		
397	472	231	232		
171	172	233	234		
399	474	235	236		
173	174	237	238		
401	174	239	240		
175	176	241	242		
403	476	243	244		
177	178	245	246		
405	478	247	248		
179	180	249	250		
407	480	251	252		
181	182	253	254		
409	482	255	256		
183	184	257	258		

FIGURE 16 (cont.)

SUBSTITUTE SHEET (RULE 26)

INTERNATIONAL SEARCH REPORT

International application No
PCT/US2019/059654

A. CLASSIFICATION OF SUBJECT MATTER
INV. C07K14/435
ADD.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED
Minimum documentation searched (classification system followed by classification symbols)
C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
EPO-Internal, BIOSIS, Sequence Search, EMBASE, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO 2009/030780 A2 (COMPLIX NV [BE]; DESMET JOHAN [BE]; LASTERS IGNACE [BE]) 12 March 2009 (2009-03-12) abstract page 15, line 16 - page 16, line 26 page 23 - page 26 page 32 - page 34 claims 1-17 ----- -/--	1-116

Further documents are listed in the continuation of Box C.

See patent family annex.

* Special categories of cited documents :

<p>"A" document defining the general state of the art which is not considered to be of particular relevance</p> <p>"E" earlier application or patent but published on or after the international filing date</p> <p>"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)</p> <p>"O" document referring to an oral disclosure, use, exhibition or other means</p> <p>"P" document published prior to the international filing date but later than the priority date claimed</p>	<p>"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention</p> <p>"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone</p> <p>"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art</p> <p>"&" document member of the same patent family</p>
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Date of the actual completion of the international search 18 December 2019	Date of mailing of the international search report 09/03/2020
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Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016	Authorized officer Keller, Yves
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INTERNATIONAL SEARCH REPORT

International application No
PCT/US2019/059654

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO 97/12988 A1 (PENCE INC [CA]; HOUSTON MICHAEL E JR [CA] ET AL.) 10 April 1997 (1997-04-10) abstract page 3 - page 4 figure 3D figure 4A page 14 - page 16 figures 1-25 figures 1A-1F	1-116
X	----- O'SHEA E K ET AL: "Peptide 'Velcro': Design of a heterodimeric coiled coil", CURRENT BIOLOGY, CURRENT SCIENCE, GB, vol. 3, no. 10, 1 October 1993 (1993-10-01), pages 658-667, XP024248213, ISSN: 0960-9822, DOI: 10.1016/0960-9822(93)90063-T [retrieved on 1993-10-01] the whole document	1-116
X	----- HELENA GRADISAR ET AL: "De novo design of orthogonal peptide pairs forming parallel coiled-coil heterodimers", JOURNAL OF PEPTIDE SCIENCE, vol. 17, no. 2, 28 December 2010 (2010-12-28), pages 100-106, XP055066656, ISSN: 1075-2617, DOI: 10.1002/psc.1331 the whole document	1-116
X	----- CROOKS RICHARD O ET AL: "Deriving Heterospecific Self-Assembling Protein-Protein Interactions Using a Computational Interactome Screen", JOURNAL OF MOLECULAR BIOLOGY, ACADEMIC PRESS, UNITED KINGDOM, vol. 428, no. 2, 2 December 2015 (2015-12-02), pages 385-398, XP029407028, ISSN: 0022-2836, DOI: 10.1016/J.JMB.2015.11.022 the whole document	1-116
X	----- AARON W. REINKE ET AL: "A Synthetic Coiled-Coil Interactome Provides Heterospecific Modules for Molecular Engineering", JOURNAL OF THE AMERICAN CHEMICAL SOCIETY, vol. 132, no. 17, 5 May 2010 (2010-05-05), pages 6025-6031, XP055652786, ISSN: 0002-7863, DOI: 10.1021/ja907617a the whole document	1-116
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INTERNATIONAL SEARCH REPORT

International application No

PCT/US2019/059654

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
T	<p>AMANDA L. EDWARDS ET AL: "Challenges in Targeting a Basic Helix-Loop-Helix Transcription Factor with Hydrocarbon-Stapled Peptides", ACS CHEMICAL BIOLOGY, vol. 11, no. 11, 19 September 2016 (2016-09-19), pages 3146-3153, XP055373864, ISSN: 1554-8929, DOI: 10.1021/acscchembio.6b00465</p> <p>-----</p>	
T	<p>HELENA GRADISAR ET AL: "Design of a single-chain polypeptide tetrahedron assembled from coiled-coil segments", NATURE CHEMICAL BIOLOGY, vol. 9, no. 6, 28 April 2013 (2013-04-28), pages 362-366, XP055293813, Basingstoke ISSN: 1552-4450, DOI: 10.1038/nchembio.1248 the whole document</p> <p>-----</p>	
X,P	<p>CHEN ZIBO ET AL: "Programmable design of orthogonal protein heterodimers", NATURE, MACMILLAN JOURNALS LTD., ETC., LONDON, vol. 565, no. 7737, 19 December 2018 (2018-12-19), pages 106-111, XP036664328, ISSN: 0028-0836, DOI: 10.1038/S41586-018-0802-Y [retrieved on 2018-12-19] the whole document</p> <p>-----</p>	1-116

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US2019/059654

Box No. II Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. Claims Nos.:
because they relate to subject matter not required to be searched by this Authority, namely:

2. Claims Nos.:
because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3. Claims Nos.:
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box No. III Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

see additional sheet

1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.

2. As all searchable claims could be searched without effort justifying an additional fees, this Authority did not invite payment of additional fees.

3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:

4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

1-116(partially)

Remark on Protest

- The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.
- The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.
- No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. claims: 1-116(partially)

A heterodimer protein, comprising:(a) a monomer A polypeptide, wherein the monomer A polypeptide comprising 1-5 alpha helices and(b) a monomer B polypeptide, wherein the monomer B polypeptide 1-5 alpha helices,wherein monomer A and monomer B non-covalently interact to form the designed heterodimer protein and wherein the alpha helice of monomer A is SEQ ID. No. 1 (as well as subject matter to the monomer A as defined above (SEQ ID No 1))

2-290. claims: 1-116(partially)

A heterodimer protein, comprising:(a) a monomer A polypeptide, wherein the monomer A polypeptide comprising 1-5 alpha helices and(b) a monomer B polypeptide, wherein the monomer B polypeptide 1-5 alpha helices,wherein monomer A and monomer B non-covalently interact to form the designed heterodimer protein and wherein the alpha helice of monomer A is SEQ ID. No. 2-290 (as well as subject matter to the monomer A as defined above (SEQ ID No 2-290)) each SEQ ID No representing a single invention

INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No

PCT/US2019/059654

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 2009030780	A2	12-03-2009	AT 538129 T 15-01-2012
			EP 2188303 A2 26-05-2010
			US 2010305304 A1 02-12-2010
			WO 2009030780 A2 12-03-2009

WO 9712988	A1	10-04-1997	AU 695679 B2 20-08-1998
			EP 0854931 A1 29-07-1998
			JP H11512620 A 02-11-1999
			WO 9712988 A1 10-04-1997
