A high throughput approach for the generation of orthogonally interacting protein pairs

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In contrast to the nearly error-free self-assembly of protein architectures in nature, artificial assembly of protein complexes with pre-defined structure and function in vitro is still challenging. To mimic nature’s strategy to construct pre-defined three-dimensional protein architectures, highly specific protein-protein interacting pairs are needed. Here we report an effort to create an orthogonally interacting protein pair from its parental pair using a bacteria-based in vivo directed evolution strategy. This high throughput approach features a combination of a negative and a positive selection. The newly developed negative selection from this work was used to remove any protein mutants that retain effective interaction with their parents. The positive selection was used to identify mutant pairs that can engage in effective mutual interaction. By using the cohesin-dockerin protein pair that is responsible for the self-assembly of cellulosome as a model system, we demonstrated that a protein pair that is orthogonal to its parent pair could be readily generated using our strategy. This approach could open new avenues to a wide range of protein-based assembly, such as biocatalysis or nanomaterials, with pre-determined architecture and potentially novel functions and properties.

Although significant progress has been made in recent years1–14, the precise manipulation of artificial self-assembly of protein complexes in vitro remains a great challenge. In contrast, highly ordered permanent or transient protein complexes widely exist in nature and participate in virtually every type of cellular function, including catalysis, structural support, bodily movement, signal transduction, transport, etc. Nature’s error-free self-assembly of protein architectures, such as virus capsids15, bacterial carboxysomes16, and cellulosomes (Fig. 1)17–19 is driven by many weak, noncovalent interactions at protein-protein interfaces20. The geometry of subunits in a protein complex is precisely defined by those specific noncovalent interactions13. In order to mimic nature’s strategy to construct highly defined three-dimensional protein architectures, we need to have highly specific protein-protein interacting pairs, analogous to G-C and A-T base-pairing interactions in DNA. One potential solution is to explore naturally occurring protein pairs, such as the barnase and barstar pair21. The other potential approach is to artificially generate mutually orthogonal protein pairs from a known parent protein pair. This could further expand the repertoire of highly specific protein pairs that are available for the assembly of protein complexes. In addition, such evolved protein pairs are orthogonal but consist of high sequence homology to the parent protein pair and, therefore, have similar physical/chemical properties. This may minimize certain complications when protein pairs with very different properties are used in the assembly of a protein complex.

To demonstrate the feasibility of the aforementioned approach, we selected a type-I cohesin-dockerin pair from Clostridium thermocellum as our model system. High affinity cohesin-dockerin interactions are the basis of self-assembly of cellulosomes17,18, which are multi-protein complexes from certain anaerobic bacteria and fungi for a highly efficient degradation of cellulotic material (Fig. 1). A cellulosome consists of a core structural protein (scaffoldin) that serves as a scaffold to connect multiple catalytic enzymes through the interaction between the type I cohesin domains on itself and the type I dockerin domains of catalytic enzymes. Due to the indiscriminatory nature of cohesin–dockerin recognition within a microorganism species, the assembled cellulosomes have diverse molecular composition and structure, which corresponds to heterogeneous catalytic activities for cellulotic material degradation. In this work, we seek to generate a mutant cohesin-dockerin pair (Fig. 2A) that is derived from but orthogonal to the naturally occurring (wild-type) one. The generation of orthogonal

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cohesin-dockerin pairs will allow controlled assembly of cellulosomes (Fig. 1), which will facilitate current studies of synergistic actions among cellulosomal enzymes. The second cohesin domain from the scaffoldin protein (CipA; residues 182–328) and the dockerin domain from a glycoside hydrolase (xylanase 10B; residues 733–791) were used in this study. The crystal structure of the protein complex of these two domains has been reported.

A few methods have been developed for a high-throughput engineering of protein-protein interactions. Phage, yeast, and bacterial displayed protein libraries are generally screened with either panning or fluorescence-activated cell sorting (FACS). In contrast, the yeast two-hybrid system links protein-protein interactions to a phenotype (e.g., cell growth) that confers a selective advantage to the host, which simplifies the selection process. The bacterial two-hybrid system has also been developed. In comparison to the yeast system, the bacterial system has the advantage of higher transformation efficiency and faster cell growth rate. However, the current bacterial system lacks a negative (counter) selection scheme. In the present study, we devised a bacterial negative selection scheme that is analogous to the yeast one. We subsequently demonstrated that an orthogonal mutant pair could be readily obtained through a combination of positive and negative selections. Throughout the article, we will use Cohwt-Docwt and Coh1-Doc1 to represent the parent (wild-type) and mutant (evolved) cohesin-dockerin pairs, respectively.

Results and Discussion

General approach. In order to generate a mutant cohesin-dockerin pair that is orthogonal to its parent pair, we explored a structure-guided, semi-rational protein engineering approach (Fig. 2). This approach consists of two essential steps: (1) mutagenesis. The important amino acid residues at the protein-protein interface of the parent pair are randomized. Presumably, such modification would completely abolish or significantly weaken the interaction between mutants and their parents; and (2) selection. This process identifies mutant protein pairs that interact to each other, but do not have significant cross interaction with the parent protein pair. Our selection system consists of both a positive selection and a negative selection. The positive selection selects mutant cohesin-dockerin pairs that can engage in effective interaction. The negative selection removes any dockerin or cohesin mutants that retain effective interaction with the parent cohesin or dockerin. The combination of negative and positive selection should yield interacting protein pairs that are orthogonal to their parent. This selection scheme can likely be generalized and used to create orthogonal pairs for other proteins as well.
Positive selection system. The BacterioMatch II two-hybrid system was used as the molecular biology platform for the positive selection (Fig. 3A). Two reasons promoted us to choose bacteria (E. coli) two-hybrid over yeast two-hybrid as the positive selection system: (1) E. coli grows much faster than yeast; (2) E. coli is transformed with higher efficiency so larger libraries can be readily constructed and selected/screened. BacterioMatch II selection is built upon the genetic complementation of the chromosomal hisB gene deletion by the episomal expression of the S. cerevisiae HIS3 gene in an E. coli host strain. Both genes encode imidazoleglycerol-phosphate dehydratase, which is an essential enzyme in the L-histidine biosynthesis. To study the interaction between a cohesin and a dockerin protein, the cohesin is expressed as a C-terminal fusion protein to the full-length bacteriophage λ repressor protein (λcI), and the dockerin is fused to the N-terminal domain of the α-subunit of RNA polymerase (RNAPα). When both fusion proteins are co-expressed in E. coli selection host, if the cohesin and the dockerin variants interact, they recruit and stabilize the binding of RNA polymerase at the promoter and activate the transcription of the HIS3 reporter gene, which allows cells to grow in the presence of 3-amino-1,2,4-triazole (3-AT), a competitive inhibitor of HIS3 gene product. In general, a stronger interaction confers the cells resistant to higher concentrations of 3-AT, while lack of interaction only permits cells to survive on media without 3-AT. It should be noted that other factors, such as the protein expression level, may affect the cell growth as well. For example, a protein pair with higher expression level can likely survive higher concentrations of 3-AT than a protein pair with lower expression level. If one would like to compare the interaction strength between two different protein pairs by using the positive selection system, the expression levels of the two pairs need to be adjusted to a similar level (e.g., to manipulate gene transcription level using different concentrations of inducer such as IPTG).

To test if the positive selection works, we examined the interaction between the Cohwt-Docwt pair. To this end, two plasmids were constructed, including pBT-Cohwt (containing the gene that encodes the λcI-Cohwt fusion protein) and pTRG-Docwt (containing the gene that encodes the RNAPα-Docwt fusion protein). We initially examined a construct in which Docwt was directly fused to RNAPα. However, we observed poor cell growth in a two-hybrid study of the Cohwt-Docwt interaction. We hypothesized that such poor cell growth was resulted from the degradation of the RNAPα-Docwt fusion protein since it was known that dockerin domain is prone to degradation in Escherichia coli. According to literature, dockerin-containing enzymes could be expressed as full-length proteins in E. coli; we decided to improve the stability of the dockerin protein by inserting the X6b carbohydrate-binding domain between RNAPα and the dockerin domain. The X6b domain is naturally fused to the N-terminus of the type I dockerin domain from C. thermocellum and does not interact with the cohesin domain. The X6b domain was included in all dockerin constructs in this work. As shown in Table 1, cell growth was observed in the presence of 5 mM of 3-AT when both pBT-Cohwt and pTRG-Docwt were co-transformed into the E. coli selection host (entry 1; Table 1). As negative controls, no cell growth was detected when either pBT-Cohwt was co-transformed with empty pTRG vector or the pTRG-Docwt was co-transformed...
with the empty pBT vector (entry 2, 3; Table 1). The results confirmed that cells only grew when both Coh wt and Doc wt proteins were present. In comparison to the positive control provided by the BacterioMatch® II kit, the interaction between Coh wt and Doc wt (entry 1; Table 1) led to similar level of cell growth as the interaction between Gal11P and LGF2 (entry 4; Table 1), which were co-expressed from the pTRG-Gal11P and the pBT-LGF2 positive control plasmids.

**Negative selection system.** As a critical component to enable the generation of orthogonally interacting protein pairs, we developed a negative selection method (Fig. 2B). We modified the yeast URA3/5-FOA counter selection system into the bacterial two-hybrid system. UR3 encodes orotidine 5′-phosphate decarboxylase, which catalyzes the transformation of 5-fluoroorotic acid (5-FOA) into a highly toxic compound (5-fluorouracil) and causes cell death. A similar approach was demonstrated in a bacterial one-hybrid system to select for Zn finger proteins.

To enable the selection, we first deleted the pyrF gene (encodes orotidine 5′-phosphate decarboxylase) on the chromosome of the BacterioMatch II reporter strain to generate strain WNPPI7. As a result, E. coli WNPPI7 lost the ability to convert 5-fluoroorotic acid (5-FOA) into a cytotoxic compound, and therefore survives on solid minimal media containing 5-FOA and uracil supplementation (Entry 3, 4; Table 2). We then modified the F′ plasmid of WNPPI7 to replace the HIS3 reporter gene with a copy of the URA3 gene, resulting in strain WNPPI5. However, the 5-FOA tolerance test showed that the basal expression level of URA3 protein in strain WNPPI5 is high enough to lead to cell death of the host strain itself on plates containing 0.5 mM of 5-FOA (Entry 5, 6; Table 2). To solve the problem, we constructed strain WNPPI8, in which the URA3 gene was inserted behind the HIS3 reporter gene. As the second gene in an operon, the reduced basal expression level of URA3 in strain WNPPI8 allowed the cells to survive on plates containing 2.5 mM of 5-FOA (Entry 7; Table 2). When an interacting protein pair (LGF2 and Gal11P) was expressed in strain WNPPI8, the increased transcription level of URA3 resulted in cell death in the presence of as low as 0.5 mM of 5-FOA (Entry 8; Table 2). The negative selection system was further evaluated when pBT-Coh wt and pTRG-Doc wt were co-transformed into WNPPI8. 5-FOA
Identification of an orthogonal cohesin-dockerin pair through library selections. The dockerin library was first subjected to one round of negative selection against Cohwt in order to eliminate dockerin mutants that retained effective interaction with Cohwt. Surviving cells should contain dockerin mutants that were either non-functional or lost the ability to recognize Docwt. Similarly, the cohesin library was subjected to one round of negative selection against Docwt in order to eliminate cohesin mutants that retained effective interaction with Docwt. Surviving cells should contain cohesin mutants that were either non-functional or lost the ability to recognize Docwt. It should also be noted that some mutants that can engage in effective interaction with Cohwt or Docwt might survive the negative selection if their expression levels were too low to induce a sufficient level of URA3 expression. While such possibility exists, these mutants will likely be eliminated in the positive selection due to their low expression levels.

By using plates containing 2.5 mM FOA, the survival rate of the dockerin and cohesin mutants was estimated to be 10–20%. This number was based on the comparison between the control plate (no FOA) and the selection plate (with 2.5 mM FOA; Figure S3). We subsequently examined if we could identify cohesin mutants from the reduced cohesin library after negative selection to engage functional interaction with the aforementioned mutant dockerin variants.

| host                  | protein pair              | cell growth (cfu) with or without 5-FOA (mM) |
|-----------------------|---------------------------|---------------------------------------------|
|                       |                           | 0     | 0.5 | 2.5 |
| 1 BacterioMatch II    | none                      | ~1 × 10⁵ | 0 | 0 |
| 2 BacterioMatch II    | Gag11P and LGF2           | ~1 × 10⁵ | 0 | 0 |
| 3 WNPPI7              | none                      | ~1 × 10⁵ | ~1 × 10³ | ~1 × 10⁵ |
| 4 WNPPI7              | Gag11P and LGF2           | ~1 × 10⁵ | ~1 × 10³ | ~1 × 10⁵ |
| 5 WNPPI5              | none                      | ~1 × 10⁵ | ~1 × 10⁵ (timy⁴) | 0 |
| 6 WNPPI5              | Gag11P and LGF2           | ~1 × 10⁵ | 0 | 0 |
| 7 WNPPI5              | none                      | ~1 × 10⁵ | ~1 × 10³ | ~1 × 10⁵ |
| 8 WNPPI5              | Gag11P and LGF2           | ~1 × 10⁵ | 0 | 0 |
| 9 WNPPI5              | Cohwt and Docwt           | ~1 × 10⁵ | 0 | 0 |
| 10 WNPPI5             | Cohwt and DocAL           | ~1 × 10⁵ | ~1 × 10³ | ~1 × 10⁵ |

Table 2. Examination of negative selection host strains. a cfu, colony-forming unit. b In comparison to the regular colony size (~1 mm) from other tests, these colonies (<0.1 mm) were barely seen by eye.
DocAL mutant. To our delight, a large number of colonies were obtained after one round of positive selection between DocAL and the reduced cohesin library. We arbitrarily picked eight colonies with different sizes for DNA sequencing analysis. Seven distinct sequences were obtained, while cohesin mutants 3 and 7 converged to the same sequence (Table S2). To estimate the effectiveness of the negative selection and to eliminate false positives (e.g., beneficial host mutations) identified in the positive selection, the pBT-Coh plasmids of the seven distinctive mutants were isolated and reintroduced into the positive selection hosts that harbored either pTRG-DocAL or pTRG-Docwt. We observed that all seven mutants engaged in strong interactions with DocAL and supported cell growth in the presence of 7.5 mM 3-AT. On the other hand, six out of the seven mutants were not able to support cell growth when Docwt was co-expressed. Apparently, these cohesin mutants did not interact with Docwt, which indicated that our established negative selection protocol was highly effective.

We arbitrarily picked cohesin mutant 1 (Coh1; Asn37Leu, Asp39Thr, Gly121Leu, and Ala125Leu; Table S2) for the subsequent selection against the reduced dockerin library. Among a few hundred survived colonies, five were picked and the corresponding pTRG-Doc plasmids were isolated and reintroduced into the positive selection hosts that harbored either pBT-Cohwt or pBT-Coh1. Cell growth test confirmed that all five dockerin mutants engaged in strong interactions with Coh1 and four out of five did not interact with Cohwt. One dockerin mutant displayed moderate interaction with Cohwt. Based on the colony size and growth rate on the positive selection plate, we chose a dockerin mutant (named as Doc1; Ser11Arg, Thr12Pro, Ser45Pro, and Thr46Ala) for the following in vitro characterization.

### In vitro characterization.

Previous reports suggested that dockerin domain could not be produced as a discrete entity due to its degradation in *Escherichia coli*. On the other hand, large quantity of dockerin domain could be obtained when it was co-expressed with cohesin. In addition, many reports showed that good expression of dockerin could be achieved when it was fused to well-folded proteins. To this end, dockerin domain variants (including its N-terminal X6b domain) were expressed as a C-terminal fusion to the maltose binding protein (MBP). A His6 tag was added to the C-terminus of the fusion protein to facilitate the purification and ELISA experiments. Cohesin domain variants were purified as a C-terminal fusion to the glutathione S-transferase (GST). The GST tag improved expression of cohesin domains, facilitated protein purification, and did not interfere with the ELISA experiments using anti-His6 antibody. We have also verified that GST does not interact with MBP.

To estimate the strength of interaction between cohesin and dockerin, we conducted semi-quantitative ELISA experiments. Briefly, wells of microtiter plates were coated with a GST-tagged cohesin. Different concentrations of the His6-tagged dockerin of interest were then applied into each well. Following washing steps, the amounts of interacting dockerin were determined immunochemically using anti-His6 antibody and HRP-labeled secondary antibody. As shown in Table 3, Doc1 and Coh1 displayed a strong mutual interaction with a $K_d$ value of $4.57 \pm 1.61$ nM, which is comparable to that of the parent Docwt-Cohwt pair ($K_d = 0.77 \pm 0.10$ nM). On the other hand, the Doc1-Coh1 pair did not show obvious cross-interaction with the Docwt-Cohwt pair. The $K_d$ values of the Doc1-Cohwt and Docwt-Coh1 cross pairs were too large to be accurately measured, which were estimated to be larger than 500 nM (Table 3 and Figure S4). The ELISA experiments confirmed that the Doc1-Coh1 pair is orthogonal to the parental Docwt-Cohwt pair. To verify that the evolved Doc1 mutant does not have increased level of non-specific interaction with other proteins due to a few hydrophilic-to-hydrophobic mutations, we conducted ELISA experiments between Doc1 and a control protein, BSA. The $K_d$ value of the Doc1-BSA interaction was too large to be accurately measured. It is estimated to be larger than 1,500 nM, which is similar to that of the Docwt-BSA pair ($K_d > 1000$ nM; Figure S4). We therefore concluded that mutations in Doc1 do not promote non-specific binding.

### Conclusion.

In summary, we have developed an approach to generate an interacting protein pair that is derived from but orthogonal to the parent protein pair. This is achieved by engineering the protein-protein interacting interface and facilitated by a combination of positive and negative selections in bacteria. To the best of our knowledge, our negative selection is the first example of applying URA3/5-FOA selection in a bacterial two-hybrid system. Presumably, more than one orthogonal protein pair can be generated with multiple cycles of positive and negative selections. The approach and tools that were developed in the current work can also potentially be applied to the generation of other orthogonal proteins pairs of one’s interest. These orthogonal protein pairs can potentially be applied to the assembly of artificial protein complexes both in vitro and in vivo. The precise control of relative contents and positions of building blocks within a protein assembly will likely facilitate the construction of protein complexes for protein-based nanomaterials or for efficient catalytic synthetase of bio-based chemicals through co-localization of enzymes.

### Methods

**Materials and General Methods.** Primers were ordered from Sigma. DNA sequencing services were provided by Eurofins MWG Operon. Restriction enzymes, antarctic phosphatase (AP) and T4 DNA ligase were purchased from New England Biolabs. KOD hot start DNA polymerase was purchased from EMD Millipore. Standard molecular biology techniques were used throughout. Site-directed mutagenesis was carried out using overlapping PCR. *E. coli* XL1-Blue MR' was used in the construction and DNA propagation of all plasmids that were derived from pBT and pTRG vectors. *E. coli* GeneHogs were used for routine cloning and DNA propagation.

| $K_d$ (nM) | Docwt-Cohwt | Doc1-Coh1 | Docwt-Coh1 | Doc1-Cohwt |
|-----------|-------------|-----------|------------|------------|
| $0.77 \pm 0.10$ | $4.57 \pm 1.61$ | $>500$ | $>500$ |

Table 3. $K_d$ values of cohesin-dockerin pairs.
of all other plasms. All solutions were prepared in deionized water that was further treated by Barnstead Nanopure® ultrapure water purification system (Thermo Fisher Scientific Inc). LB medium (1 L) contained Bacto tryptone (10 g), Bacto yeast extract (5 g), and NaCl (10 g). M9 salts (1 L) contained Na₂HPO₄ (6 g), KH₂PO₄ (3 g), NH₄Cl (1 g), and NaCl (0.5 g). M9 glucose medium contained glucose (10 g), MgSO₄ (0.12 g), CaCl₂ (0.028 g) and thiamine hydrochloride (0.001 g) in 1 L of M9 salts. Antibiotics were added where appropriate to following final concentrations: ampicillin, 100 mg L⁻¹; kanamycin, 50 mg L⁻¹; chloramphenicol, 25 mg L⁻¹; tetracycline, 12.5 mg L⁻¹. The 6xHis tag monoclonal antibody was purchased from Thermo Fisher Scientific. The goat anti-mouse IgG-HRP conjugate was purchased from BioRad.

Plasmid construction. Plasmid pBT-Coh₃₇₄₉ was constructed by inserting Coh₂⁻encoding gene (the second cohesin domain from CipA; residues 182–328) between the EcoRI and BamHI sites of the pBT vector. Coh₂⁻encoding gene was PCR amplified using primers P1 and P2 (Table S1) from the chromosomal DNA of ATCC 27405.

Plasmid pTRG-Doc₃₇₄₉ was constructed by inserting Doc₂⁻encoding gene (the dockerin domain from xylanase 10B; residues 733–791) between the BamHI and Xhol sites of the pTRG vector. Doc₂⁻encoding gene was PCR amplified using primers P3 and P4 (Table S1) from the chromosomal DNA of ATCC 27405. DNA sequence that encodes X6b carbohydrate binding domain was included at the 5’ end of the Doc₂⁻-encoding gene.

Plasmid pTRG-Doc₃₇₄₉ was constructed by site-directed mutagenesis of plasmid pTRG-Doc₃₇₄₉. Mutations (Ser11Ala, Thr12Leu, Ser45Ala, and Thr46Leu) were introduced by overlapping PCR using primers P5, P6, P7, P8, and P9 (Table S1). The resulting DNA fragment was inserted into the pTRG vector using Ligation Independent Cloning (SLIC)⁴³.

Plasmids pGEX-Coh₁ and pGEX-Coh₁ were constructed by inserting Coh₁⁻encoding gene and Coh₁⁻encoding gene between the EcoRI and Xhol sites of pGEX vector, respectively. As a result, the cohesin variants were expressed as fusion protein to the C-terminus of GST. The Coh₁⁻encoding gene and Coh₁⁻encoding gene were amplified using primers P10 and P11 (Table S1).

Plasmids pMAL-Doc₁ and pMAL-Doc₁ were constructed by inserting a dockerin gene of interest between the BamHI and Xhol sites of pMAL vector, respectively. The dockerin variants were expressed as C-terminus fusion of MBP. The dockerin encoding genes were amplified using primers P3 and P12 (Table S1).

Construction of a dockerin library. Four residues (Ser11, Thr12, Ser45, and Thr46) of dockerin were randomized. Primers P29 and P30 (Table S1) were used to introduce these mutations. The full-length dockerin mutant fragments were assembled by overlapping PCR of the above two DNA fragments using primers P3 and P4 (Table S1). The resulting DNA was digested with BamHI and Xhol, and subsequently cloned into the pTRG vector that was digested with the same pair of restriction enzymes to produce the dockerin library.

Construction of a cohesin library. Four residues (Asn37, Asp39, Gly123, and Ala125) of cohesin that are in contact with the Ser/Thr motif of dockerin were randomized. Primers P13 and P14 (Table S1) were used to amplify a DNA fragment that contains mutations at positions Gly123 and Ala125. Primers P15 and P16 were used to amplify a DNA fragment that contains mutation at positions Asn37 and Asp39. The full-length cohesin mutant fragments were assembled by overlapping PCR of the above two DNA fragments using primers P1 and P2 (Table S1). The resulting DNA was digested with EcoRI and BamHI, and subsequently cloned into the pBT vector that was digested with the same pair of restriction enzymes to produce the cohesin library.

Construction of host strain for negative selection. Chromosomal deletion and modification of the F’ plasmid was carried out using the phage λ Red-mediated homologous recombination⁴⁴. In brief, an appropriate E. coli host strain was first transformed with plasmid pRed-ET (Gene Bridges). A single colony of transformed cells was cultured in LB medium containing Ap at 30 °C. Expression of the E. coli host strain was first transformed with plasmid pRed-ET (Gene Bridges). A single colony of transformed host strain was cultured in LB medium containing Ap at 30 °C. Expression of the E. coli host strain was first transformed with plasmid pRed-ET (Gene Bridges). A single colony of transformed host strain was cultured in LB medium containing Ap at 30 °C. Expression of the E. coli host strain was first transformed with plasmid pRed-ET (Gene Bridges). A single colony of transformed host strain was cultured in LB medium containing Ap at 30 °C. Expression of the E. coli host strain was first transformed with plasmid pRed-ET (Gene Bridges). A single colony of transformed host strain was cultured in LB medium containing Ap at 30 °C. Expression of the E. coli host strain was first transformed with plasmid pRed-ET (Gene Bridges). A single colony of transformed host strain was cultured in LB medium containing Ap at 30 °C. Expression of the E. coli host strain was first transformed with plasmid pRed-ET (Gene Bridges). A single colony of transformed host strain was cultured in LB medium containing Ap at 30 °C. Expression of the E. coli host strain was first transformed with plasmid pRed-ET (Gene Bridges). A single colony of transformed host strain was cultured in LB medium containing Ap at 30 °C. Expression of the E. coli host strain was first transformed with plasmid pRed-ET (Gene Bridges). A single colony of transformed host strain was cultured in LB medium containing Ap at 30 °C. Expression of the E. coli host strain was first transformed with plasmid pRed-ET (Gene Bridges). A single colony of transformed host strain was cultured in LB medium containing Ap at 30 °C. Expression of the E. coli host strain was first transformed with plasmid pRed-ET (Gene Bridges). A single colony of transformed host strain was cultured in LB medium containing Ap at 30 °C. Expression of the E. col
with shaking (225 rpm), which allowed cells to adapt to the growth in minimal medium before plating. Around 5 × 10^6 cells were plated on M9 complete medium plates containing 2.5 mM 5-FOA, 0.05 mM IPTG, and appropriate concentrations of chloramphenicol and tetracycline. After 36 h of incubation at 37 °C, cells were collected from the plates into 1 mL M9 complete medium. Plasmid DNA was extracted from the collected cells. Mixture of the pBT-CohΔh and pTRG- DocΔh plasmids was first treated by restriction enzyme to linearize the pTRG-DocΔh plasmid. The pBT-CohΔh plasmids were then isolated by DNA gel electrophoresis purification. Negative selection with the dockerin library was conducted using the same procedure.

**Positive selection.** For positive selections with the reduced cohesin library, the pBT-CohΔh from the negative selection was transformed into the BacterioMatch II reporter strain containing a pTRG-Doc variant of interest. The positive selection followed instructions of the BacterioMatch II two-hybrid kit. In brief, overnight bacterial culture in LB medium (1 mL) was collected by centrifugation. Cell pellets were washed with M9-His drop medium (1 × M9 salts, 1 mM MgSO_4, 0.1 mM CaCl_2, 10 mg/L thiamine, 0.001 mM ZnSO_4, 1 × His-drop supplement, 0.2 mM adenine, 0.4% D-glucose) to completely remove the residue LB medium. Cells were then re-suspended in 1 mL M9-His drop medium and incubated for 2 h at 37 °C with shaking (225 rpm). An aliquot of cells (3 × 10^6) was plated on M9-His drop medium plates containing 5 mM 3-AT, 0.05 mM IPTG, and appropriate concentrations of chloramphenicol and tetracycline. Positive selections with the reduced dockerin library flowed the same procedure.

**Enzyme-linked immunosorbent assay (ELISA).** MaxiSorp 96 well ELISA plates were coated overnight at 4 °C with 100 μL of the desired protein (30 nM in 0.1 M Na_2CO_3 (pH 9.6)). Following steps were all performed at a volume of 100 μL/well at room temperature unless otherwise stated. After removal of the coating solution, blocking buffer (1 mM CaCl_2 and 1% BSA in TBS buffer) was added and the plates were incubated for 1 h. The plates were washed three times with washing buffer (blocking buffer supplemented with 0.05% Tween 20 without BSA; 200 μL/well per wash). Potential binding partner of the coating protein (10 pM–1 μM in blocking buffer) was applied to each well. The plates were again incubated for 1 h followed by washing for three times. Primary antibody (anti-6xHis, 1:500 dilution in blocking buffer) was added and incubated for 1 h. Plates were washed. Secondary antibody (goat anti-mouse IgG-HRP conjugate, 1:500 dilution in blocking buffer) was added for the final incubation of 1 h followed by washing for three times. Quantification of the HRP activity used the 3, 3′, 5′-tetramethylbenzidine (TMB) substrate. The reaction was initiated by the addition of 100 μL/well TMB. After 5 min of reaction or until the desired color development achieved, 50 μL/well of 1 M H_2SO_4 was added to terminate the reaction. Absorbance at 450 nm was measured. Dissociation constants were calculated by curve fitting with Hill Equation using MATLAB (R2016b).

**References**

1. Lovejoy, B. et al. Crystal structure of a synthetic triple-stranded alpha-helical bundle. Science 259, 1288–1293 (1993).
2. Harbury, P. B., Plecs, J. J., Tidor, B., Alber, T. & Kim, P. S. High-resolution protein design with backbone freedom. Science 282, 1462–1467 (1998).
3. Gribbon, C. & et al. MagicWand: A single, designed peptide that assembles to stable, ordered α-helical fibers. Biochemistry 47, 10365–10371 (2008).
4. Zaccai, N. R. et al. A de novo peptide hexamer with a mutable channel. Nat. Chem. Biol. 7, 935–941 (2011).
5. Kodier, R. L. et al. Design and engineering of an O2 transport protein. Nature 458, 305–309 (2009).
6. Ballister, E. R., Lai, A. H., Zuckermann, R. N., Cheng, Y. & Mougous, J. D. In vitro self-assembly of tailorable nanotubes from a simple protein building block. Proc. Natl. Acad. Sci. USA 105, 3733–3738 (2008).
7. Uusi, K. et al. Nanoscale elongating control of the self-assembled protein filament with the cysteine-introduced building blocks. Protein Sci. 18, 960–969 (2009).
8. Ringler, P. & Schulz, G. E. Self-assembly of proteins into designed networks. Science 302, 106–109 (2003).
9. Walz, G., Engst, M., Haas, D. & Emmerling, M. Recombinant expression of the Bacillus subtilis FtsS peptidoglycan lipoprotein. J. Biol. Chem. 278, 42806–42813 (2003).
10. Pellock, J. A., Alber, T. & Kim, P. S. Design and engineering of a biologically active α-helix. Science 305, 1568–1572 (2004).
11. Draper, R. J. et al. Skin-derived cell adhesion molecules and the regulation of tissue repair. J. Biol. Chem. 279, 474–480 (2004).
12. Coussens, L. M. et al. The role of the tumor microenvironment in tumor evasion of the immune system. J. Clin. Invest. 116, 1381–1389 (2006).
13. Padilla, J. E., Colovos, C. & Yeates, T. O. Nanohedra: Using symmetry to design self-assembling protein cages, layers, crystals, and filaments. Proc. Natl. Acad. Sci. USA 98, 2217–2221 (2001).
14. Sinclair, J. C., Davies, K. M., Venien-Bryan, C. & Noble, M. E. Generation of protein lattices by fusing proteins with matching rotational symmetry. Nat. Nanotechnol. 6, 558–562 (2011).
15. Sinha, A. et al. Synthetic viruses: Making friends with old foes. Science 340, 1071–1076 (2011).
16. Padilla, J. E., Colovos, C. & Yeates, T. O. Nanohedra: Using symmetry to design self-assembling protein cages, layers, crystals, and filaments. Proc. Natl. Acad. Sci. USA 98, 2217–2221 (2001).
17. Sinclair, J. C., Davies, K. M., Venien-Bryan, C. & Noble, M. E. Generation of protein lattices by fusing proteins with matching rotational symmetry. Nat. Nanotechnol. 6, 558–562 (2011).
18. Vyas, S., Alber, T., Engst, M. & Emmerling, M. Recombinant expression of the Bacillus subtilis FtsS peptidoglycan lipoprotein. J. Biol. Chem. 278, 42806–42813 (2003).
19. Paterson, A. D., Ito, T. & Lomax, T. A. The P1N1 polypeptide vaccine promotes T cell mediated immune responses. J. Immunol. 184, 5097–5106 (2010).
20. Bickle, C. et al. Atomic-level models of the bacterial carboxysome shell. Science 318, 1083–1086 (2008).
21. Mayer, A. A., Gelbart, W. M. & Harvey, M. J. Cellulosomes: Multienzyme machines for degradation of plant cell wall polysaccharides. Annu. Rev. Microbiol. 58, 521–554 (2004).
22. Cristea, I. M. et al. Crystal structure of the cohesin-dockerin complex. Proc. Natl. Acad. Sci. USA 100, 13809–13814 (2003).
23. Smith, G. P. Filamentous fusion phage: novel expression vectors that display cloned antigens on the virion surface. Science 228, 1315–1317 (1985).
24. Boder, E. T. & Wittrup, K. D. Yeast surface display for screening combinatorial polypeptide libraries. Nat. Biotechnol. 15, 553–557 (1997).
25. Arndt, J. P. & Bode, W. 3D molecular graphics with PyMOL. Curr. Opin. Struct. Biol. 17, 474–480 (2007).
26. Fields, S. & Song, O. A novel genetic system to detect protein–protein interactions. Nature 340, 245–246 (1989).
27. Dove, S. L. & Hochschild, A. Conversion of the ω subunit of Escherichia coli RNA polymerase into a transcriptional activator or an activation target. *Genes Dev.* 12, 745–754 (1998).
28. Dove, S. L., Joung, J. K. & Hochschild, A. Activation of prokaryotic transcription through arbitrary protein–protein contacts. *Nature* 386, 627–630 (1997).
29. Joung, J. K., Ramm, E. I. & Pabo, C. O. A bacterial two-hybrid selection system for studying protein-DNA and protein–protein interactions. *Proc. Natl. Acad. Sci. USA* 97, 7382–7387 (2000).
30. Vidal, M., Brachmann, R. K., Fattaey, A., Harlow, E. & Boeke, J. D. Reverse two-hybrid and one-hybrid systems to detect dissociation of protein–protein and DNA–protein interactions. *Proc. Natl. Acad. Sci. USA* 93, 10315–10320 (1996).
31. Boeke, J. D., LaCroute, F. & Fink, G. R. A positive selection for mutants lacking orotidine-5′-phosphate decarboxylase activity in yeast: 5-fluoro-orotic acid resistance. *Mol. Gen. Genet.* 197, 345–346 (1984).
32. Haimovitz, R. *et al.* Cohesin-dockerin microarray: diverse specificities between two complementary families of interacting protein modules. *Proteomics* 8, 968–979 (2008).
33. Karpol, A., Barak, Y., Lamed, R., Shoham, Y. & Bayer, E. A. Functional asymmetry in cohesin binding belies inherent symmetry of the dockerin module: insight into cellulose assembly revealed by systematic mutagenesis. *Biochem. J.* 410, 331–338 (2008).
34. Schaeffer, F. *et al.* Matching fusion protein systems for affinity analysis of two interacting families of proteins: The cohesin-dockerin interaction. *J. Mol. Recognit.* 18, 491–501 (2005).
35. Fierobe, H.-P. *et al.* Design and production of active cellulose chimeras. Selective incorporation of dockerin–containing enzymes into defined functional complexes. *J. Biol. Chem.* 276, 21257–21261 (2001).
36. Meng, X. & Wolfe, S. A. Identifying DNA sequences recognized by a transcription factor using a bacterial one-hybrid system. *Nat. Protoc.* 1, 30–46 (2006).
37. Schaeffer, F. *et al.* Duplicated dockerin subdomains of *Clostridium thermocellum* endoglucanase CelD bind to a cohesin domain of the scaffolding protein CipA with distinct thermodynamic parameters and a negative cooperativity. *Biochemistry* 41, 2106–2114 (2002).
38. Mechaly, A. *et al.* Cohesin-dockerin recognition in cellulose assembly: experiment versus hypothesis. *Proteins: Struct., Funct., Genet.* 39, 170–177 (2000).
39. Mechaly, A. *et al.* Cohesin-dockerin interaction in cellulose assembly: a single hydroxyl group of a dockerin domain distinguishes between nonrecognition and high affinity recognition. *J. Biol. Chem.* 276, 9883–9888 (2001).
40. Schaeffer, F. *et al.* Duplicated dockerin subdomains of *Clostridium thermocellum* endoglucanase CelD bind to a cohesin domain of the scaffolding protein CipA with distinct thermodynamic parameters and a negative cooperativity. *Biochemistry* 41, 2106–2114 (2002).
41. Carvalho, A. L., Dias, F. M. V., Nagy, T., Prates, J. A. M. & Proctor, M. R. Evidence for a dual binding mode of dockerin modules to cohesins. *Proc. Natl. Acad. Sci. USA* 104, 3089–3094 (2007).
42. Sambrook, J. E., Russell, D. W. & Editors. *Molecular cloning: A laboratory manual, third edition.* (Cold Spring HarborLaboratory Press, 2000).
43. Barak, Y. *et al.* Harnessing homologous recombination in *vivo* to generate recombinant DNA via SLIC. *Nat. Methods* 4, 251–256 (2007).
44. Datsenko, K. A. & Wanner, B. L. One-step inactivation of chromosomal genes in *Escherichia coli* K-12 using PCR products. *Proc. Natl. Acad. Sci. USA* 97, 6640–6645 (2000).

Acknowledgements
This work is supported by the New Faculty Startup Fund to J. Guo from the Chemistry Department of University of Nebraska – Lincoln, by cycle-5 grant from the Nebraska Center For Energy Sciences Research (to J.G. and W.N.), and by Grant CBET 1264708 (to J.G. and W.N.) from NSF.

Author Contributions
J.L. constructed plasmids, performed protein purification and *in vitro* characterization. X.S. constructed plasmids, performed selection and protein purification. W.N. constructed libraries, devised the negative selection, and wrote the manuscript. J.G. designed the study and wrote the manuscript.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-018-19281-6.

Competing Interests: The authors declare that they have no competing interests.

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