Fine Tuning of Spatial Arrangement of Enzymes in a PCNA-Mediated Multienzyme Complex Using a Rigid Poly-L-Proline Linker

Tomoaki Haga¹, Hidehiko Hirakawa¹*, Teruyuki Nagamune¹,²*

1 Department of Chemistry and Biotechnology, School of Engineering, the University of Tokyo, Hongo, Tokyo, Japan, 2 Department of Bioengineering, School of Engineering, the University of Tokyo, Hongo, Tokyo, Japan

Abstract

Inspired by natural multienzyme complexes, many types of artificial multienzyme complexes have recently been constructed. We previously constructed a self-assembled complex of a bacterial cytochrome P450 and its ferredoxin and ferredoxin reductase partners using heterotrimerization of proliferating cell nuclear antigen (PCNA) from Sulfolobus solfataricus. In this study, we inserted different peptide linkers between ferredoxin and the PCNA subunit, and examined the effect on activity of the self-assembled multienzyme complex. Although the activity was affected by the lengths of both the rigid poly-L-proline-rich linkers and the flexible Gly₄-Ser repeating linkers, the poly-L-proline-rich linkers provided the greatest activity enhancement. The optimized poly-L-proline-rich linker enhanced the activity 1.9-fold compared with the GGGGLVPGRSGGGGS linker used in the previously reported complex, while the Gly₄-Ser repeating linkers, (G₄S)ₙ (n = 1–6), did not yield higher activity than the maximum activity by the optimized poly-L-proline linker. Both the rigidity/flexibility and length of the peptide linker were found to be important for enhancing the overall activity of the multienzyme complex.

Citation: Haga T, Hirakawa H, Nagamune T (2013) Fine Tuning of Spatial Arrangement of Enzymes in a PCNA-Mediated Multienzyme Complex Using a Rigid Poly-L-Proline Linker. PLoS ONE 8(9): e75114. doi:10.1371/journal.pone.0075114

Introduction

In nature, enzymes often work together in cascade or coupled reactions by forming multienzyme complexes. This helps to prevent degradation of unstable intermediate products and to enhance mass transfer between enzymes. Inspired by natural multienzyme complexes, artificial multienzyme complexes have been constructed by gene fusion, co-immobilization, co-entrapment, and scaffold-mediated assembly [1]. Scaffold-mediated multienzyme complexes have been constructed using biomaterials such as RNA [2], DNA [3–7] and protein [8–12]. In these complexes, enzymes are connected to single scaffolds by nucleic acids or peptide linkers. The linkers regulate the relative spatial arrangements of the enzymes, which affect the overall activities of the multienzyme complexes. However, except for a few reports [8,9], the linkers in artificial multienzyme complexes have not been examined in detail.

Recently, we demonstrated selective complex formation of Pseudomonas putida cytochrome P450 (P450) with its redox partner proteins, putidaredoxin (PdX) and putidaredoxin reductase (PdR), using proliferating cell nuclear antigen (PCNA) subunits from Sulfolobus solfataricus as a self-assembling scaffold [13] (Figure 1A). S. solfataricus PCNA is a heterotrimeric ring-shaped protein that provides a scaffold for DNA-related enzymes by encircling dsDNA [14–16]. The PCNA heterotrimer is composed of three distinct subunits, PCNA1, PCNA2 and PCNA3, which can be separately expressed as monomer proteins and then assembled to form the heterotrimer. Thus, three genetic fusion proteins of the PCNA subunits with PdR, PdX and P450cam, i.e. PCNA1-PdR, PCNA2-PdX and PCNA3-P450cam, respectively, were used to form a heterotrimeric complex termed “PUPPET” (PCNA-utilized protein complex of P450 and its two electron transfer-related proteins). PUPPET showed much higher monooxygenase activity than an equimolar mixture of free PdR, PdX and P450cam. This is because PdR, PdX and P450cam are located in close proximity to each other on the PCNA heterotrimer and thus electrons are more efficiently relayed from PdR to P450cam via PdX. However, the peptide linkers between the PCNA subunits and the three enzymes,
which are likely to affect the overall activity of PUPPET, were not examined in this earlier study.

In this report, we examined the effect of the peptide linker between PCNA2 and PdX on the monooxygenase activity of PUPPET. An optimum poly-L-proline-rich linker increased the activity 1.9-fold compared with the GGGGSLVPRGSGGGGS linker used in the previous report. The increased activity conferred by the poly-L-proline-rich linker could not be achieved using the optimized Gly4-Ser repeating linker. The poly-L-proline-rich linker provided a rigid spacer between PdX and PCNA and thus may have enabled a spatially-appropriate arrangement for more efficient interactions between PdX and P450cam. Therefore, linkers comprising a rigid peptide sequence are likely to be the best candidates for fusing enzymes with protein scaffolds for the construction of multienzyme complexes.

**Materials and Methods**

**Plasmid construction**

The plasmids expressing PCNA2 protein fused to Pdx were constructed by inserting DNA sequences encoding peptide linkers between the C-terminus of PCNA2 and the N-terminus of the Cys73Ser/Cys85Ser mutant of PdX. First, two SapI sites were inserted by PCR between the genes encoding PCNA2 and the Cys73Ser/Cys85Ser mutant of PdX in pHSG+PCNA2-PdX [17], using two primers, 5'-GGATCCGCTCTTCAGGAGGTGGTGGCTCTATGTCTAAAG-
3’ (forward) and 5’-GAATTCGCTTCTTACCCGCACGCACG-3’ (reverse). The DNA fragment, containing two Sapl sites between PCNA2 and PdX, was amplified from the resulting plasmid by PCR using two primers, 5’-TATTCCAGGGCCATATGAAAGCGCAATGTACCG-3’ (forward) and 5’-ATGCGCTTGAAATACAGG-3’ (reverse). A modified pET-15b (+) (Novagen, Darmstadt, Germany), termed pET15T, which lacked a Sapl site and had a TEV protease recognition site between a thrombin recognition site and an Ndel site, was linearized by PCR using two primers, 5’-GGATCCGGCTGCTAACAAAG-3’ (forward) and 5’-GAATTCGCTCTTCTACCGCCGTCCGCGCTTACCG-3’ (reverse). The amplified DNA fragment was fused to the linearized plasmid using the In-Fusion enzyme (Clontech, Mountain View, CA, USA), to obtain p2SSX. The oligonucleotide sequences encoding (GnS)n (n = 1–3) were inserted between the sequences encoding PCNA2 and PdX into p2SSX by PCR. The oligonucleotide sequences encoding (GnS)n (n = 4–6) and GnS(Pn)GnS (n = 1–5) were inserted by ligating hybridized oligonucleotides with sticky ends into Sapl-digested p2SSX. The oligonucleotides used to insert the linkers are summarized in Table S1.

A gene encoding the PCNA3-P450cam fusion protein was inserted between the Ndel and BamHI sites of pET-15b (+). The resulting plasmid, pET15O+O3C, was mutated by PCR using two primers, 5’-AGCACCCGTCGTGAATTTACGTCGC-3’ (forward) and 5’-TTAAGCCGAGGGTGCTGGCATAATGGT-3’ (reverse). The resulting plasmid, pET15O+P3 N106C, expressed and purified as described in a previous report [13], PCNA1-PdR and PCNA2-PdX fusion proteins were expressed and purified as described in a previous report [13], except that BL21 Star (DE3) pLysS (Invitrogen, Carlsbad, CA, USA) was used. PCNA3-P450cam fusion protein, in which PCNA3 had a N106R mutation that improved the solubility of the fusion protein but did not affect PUPPET formation (unpublished data), was expressed and purified as described in the above report, except that T7 Express (EMD Millipore, Billerica, MA, USA) was transformed with pET15O+P3 N106C.

An approximately equimolar ratio of PCNA1-PdR and PCNA2-PdX and a slightly excess amount of PCNA3-P450cam were mixed and incubated on ice for 1 h. The mixture was subjected to size-exclusion chromatography on a Superdex 200 10/300 GL column (1.0 x 30 cm; GE Healthcare, Little Chalfont, Buckinghamshire, UK) with 50 mM potassium phosphate buffer, pH7.4, containing 150 mM potassium chloride and 2 mM d-camphor (Wako Pure Chemical Industries, Osaka, Japan). The fractions with an Abs392/Abs455 ratio of 0.31–0.34 were collected. The collected fractions were combined and concentrated using an Amicon Ultra-15 Centrifugal Unit (30,000 NMWL; Millipore, Billerica, MA, USA). The d-camphor was then removed using a HiTrap desalting column pre-equilibrated with 50 mM potassium phosphate buffer, pH7.4, containing 150 mM potassium chloride.

**Enzyme assays**

Cytochrome c reduction rates were determined from the absorption changes at 550 nm. The extinction coefficient difference (cytochrome c red – cytochrome c ox) used to calculate the rates was 21.1 mM-1 cm-1. The assays were performed in 50 mM potassium phosphate buffer, pH7.4, containing 150 mM potassium chloride, 20 µM cytochrome c (Sigma-Aldrich, St. Louis, MS, USA), 50 µM NADH (Sigma-Aldrich) and 4.5 mM heterotrimeric complex. P450cam catalytic activities were determined from O2 consumption rates, which were directly measured using a Clark-type oxygen electrode, by subtracting the values obtained in the absence of d-camphor from those in the presence of d-camphor. The assays were performed in 50 mM potassium phosphate buffer, pH7.4, containing 150 mM potassium chloride, 500 µM d-camphor, 100 µM NADH and various concentrations of the heterotrimers. All assays were conducted at 25°C.

**Results**

**Construction of PCNA2-PdX fusion proteins**

To enhance the activity of PUPPET, we optimized the peptide linker connecting PdX and PCNA2, because PdX interacts with both PdR and P450cam, and the spatial arrangement of PdX relative to these enzymes affects their interactions in the PUPPET complex. We therefore introduced various lengths of the poly-L-proline or Gly2-Ser repeating linkers between PCNA2 and PdX. Rigid poly-L-proline peptides have often been used as spacers [18–22]. Flexible Gly2-Ser repeating peptides have also frequently been used for antibody engineering [23] and bifunctional fusion protein construction [24–26]. PCNA2-PdX fusion proteins, PCNA2-GnS(Pn)GnS-PdX (n = 1–5) and PCNA2-(GnS)n-PdX (n = 1–6), which have proline-rich linkers, GnS(Pn)GnS, and Gly–Ser-rich linkers, (GnS)n, respectively (Figure 1B), were successfully expressed and purified (Figure 2A). The fusion proteins retained characteristic UV–Vis spectra for PdX (Figure 3A and Figure S1).

**Preparation of the heterotrimers**

Heterotrimers of PdR, PdX and P450cam were prepared by mixing PCNA1-PdR, PCNA2-PdX and PCNA3-P450cam. Each resulting protein complex, PUPPET-Pn (n = 1–5) and PUPPET-GnS (n = 1–6), comprising PCNA2-GnS(Pn)GnS-PdX (n = 1–5) and PCNA2-(GnS)n-PdX (n = 1–6), respectively (Figure 2B), showed similar UV–Vis spectra to a linear combination of PCNA1-PdR, PCNA2-PdX and PCNA3-P450cam, as previously reported [13] (Figure 3B and Figure S2). The spectrum of each PUPPET had a slightly lower absorbance around 390 nm and slightly higher absorbance around 420 nm, compared with the linear combination spectra of PCNA1-PdR, PCNA2-PdX and PCNA3-P450cam as observed in the previous study. The linker between PCNA2 and PdX did not significantly affect the PUPPET spectrum (Figure 3B and Figure S2).
Likewise, in PUPPET-G respectively. Proteins and PUPPETs.

Cytochrome c reduction activities of PUPPET (Figure 2).

The linker between PCNA2 and PdX was not observed to significantly affect the cytochrome c reduction activity of PUPPET (Figure 4).

Effect of the linker on PUPPET activity

The effect of the linker on the monooxygenase activities of PUPPET was evaluated by measuring the substrate-dependent O₂ consumption rate at the complex concentration of 18 nM (Figure 5). In PUPPET-Pₙ, the activity increased with increasing linker length in the range of n = 1–3, although PUPPET-P₉ (n = 3–5) showed similar activity (Figure 5A). Likewise, in PUPPET-Gₙ, the activity increased with increasing linker length in the range of n = 1–3, although PUPPET-G₉ (n = 3–6) showed similar activity (Figure 5B). Although the linker length had an effect on the monooxygenase activity of both PUPPET-Pₙ and PUPPET-Gₙ, the poly-L-proline-rich linkers, G₄S₅P₃₅G₄Sₕ(n = 2–5), were more effective than the Gly₄Ser repeating linkers in enhancing the activity of PUPPET. The G₄S₅P₃₅G₄Sₕ linker gave the highest activity (6.5 ± 0.3 μM.min⁻¹); a level not achieved by the Gly₄Ser repeating linkers, (G₄Sₕ) (n = 1–6). This activity is 1.9-fold greater than that of the previously reported PUPPET complex that had a GGGGSSLVPRGSGGGGS linker (3.4 ± 0.1 μM.min⁻¹).

We compared the monooxygenase activities of PUPPET-P₄ and PUPPET-G₉ to the original PUPPET complex (Figure 6), using various enzyme concentrations. In agreement with the previous report [13], the initial activity of each PUPPET did not have a linear relationship with the PUPPET concentration (Figure 6A), and the apparent specific activity decreased according to the decrease in PUPPET concentration (Figure 6B), probably because of the dissociation of PCNA3-P450cam from PUPPET at lower PUPPET concentrations. At each enzyme concentration, PUPPET-P₄ showed higher activity compared with PUPPET-G₉. The apparent specific activities of PUPPET-P₄ and PUPPET-G₉ showed similar PUPPET concentration dependency to that of the original PUPPET, and therefore the enhanced activity levels of PUPPET-P₄ and PUPPET-G₅ compared with the original PUPPET complex are independent of the PUPPET concentration.

Discussion

The functions of natural multi-domain proteins are often regulated by properties of the interdomain-connecting peptide linkers, such as their hydrophobicity and rigidity (reviewed in 28). The functions of artificial chimeric proteins are often similarly dependent on the linker properties [29]. Recently, artificial scaffold-mediated multienzyme complexes, in which linkers connect enzymes with single scaffolds, have attracted much attention because scaffolds can provide a basis for the precise control of the relative position and orientation of multiple enzymes [1]. Although the effect of linker length on protein-ligand and protein-protein associations [30,31] and bifunctional multienzyme complexes [26,32,33] has been well studied, only a few reports have investigated the effect of the linker length on artificial scaffold-mediated multienzyme complexes [8,9]. Moreover, the effect of the type of linker on the overall activities of artificial scaffold-mediated multienzyme complexes has never been examined. Linkers in artificial multienzyme complexes [3,10–12], which catalyze cascade reactions, might not have been considered to significantly affect the whole catalytic activities because cascade reactions do not depend on direct physical contact between enzymes. In this study, we examined the effect of the type and length of the peptide linker connecting PdX with the protein scaffold on the overall activity of the artificial protein complex comprising P450cam, PdX and PdR.

The monooxygenase activity of PUPPET was observed to depend on the characteristics of the linker between PCNA2 and PdX (Figure 5). This implies that the linker affected the electron transfer process from PdR to P450cam, because the electron transfer from PdX to P450cam during their transient complex formation is a rate limiting step in the P450cam...
Figure 3. UV–Vis spectra of PCNA2-G₄S-PdX and PUPPET-Gₙ. (A) UV–Vis spectrum of PCNA2-G₄S-PdX. The peaks at 314, 412, and 460 nm, and the broad band in the long wavelength region, are specific to [2Fe-2S] cluster-containing proteins. UV–Vis spectra of PCNA2-(G₄S)ₙ-PdX (n = 2–6) and PCNA2-G₄S(P₅)ₙG₄S-PdX (n = 1–5) are shown in Figure S1. (B) UV–Vis spectra of PUPPET-Gₙ (solid line), and a linear combination of the individual component protein spectra (broken line). PUPPET-Gₙ (n = 2–6) and PUPPET-Pₙ (n = 1–5) are shown in Figure S2.

doi: 10.1371/journal.pone.0075114.g003
Figure 4. Cytochrome c reduction activities of the PUPPET linker variants. The activities of (A) PUPPET-Pr (n = 1–5) and (B) PUPPET-Gpr (n = 1–6) were evaluated in a reaction mixture containing 4.5 nM enzymes and 20 µM cytochrome c. Error bars represent the standard errors of three replicates.

doi: 10.1371/journal.pone.0075114.g004
Figure 5. P450cam oxidation activities of the PUPPET linker variants. The activities of (A) PUPPET-P<sub>n</sub> (n = 1–5) and (B) PUPPET-G<sub>n</sub> (n = 1–6) were evaluated in reaction mixtures containing 18 nM enzyme. The activity of the previously reported PUPPET, containing a GGGSLVPRGSGGGS linker (indicated as "O"), and the activity of an equimolar mixture of PdR, PdX and P450cam (indicated as "R"), were also evaluated under the same reaction conditions. Error bars represent the standard error of three replicates; *P < 0.05, ***P < 0.001 (Student's t-test).

doi: 10.1371/journal.pone.0075114.g005
Figure 6. PUPPET concentration-dependent monooxygenase activities. (A) Initial rates and (B) specific activities were plotted against the protein concentration of PUPPET-P₄ (open squares), PUPPET-G₅ (open triangles) and original PUPPET (closed circles). Each specific activity was normalized using the activity at 72 nM in Figure 6B. Error bars represent the standard error of three replicates.

doi: 10.1371/journal.pone.0075114.g006
catalytic cycle [34]. The electron transfer process from PdR to P450cam is divided into two steps: reduction of PdX by PdR, and electron donation from reduced PdX to P450cam. The cytochrome c reduction activity, which reflects the reduction rate of PdX by PdR, was not significantly affected by changing the PCNA2-PdX linker (Figure 4). The dependence of cytochrome c reduction activity on the linker was not identical to that of the monooxygenase activity. This result indicates that the linker mostly affected the electron donation step from reduced PdX to P450cam.

The linker length of the poly-L-proline-rich linker, G\textsubscript{4}S(P\textsubscript{5})\textsubscript{n}G\textsubscript{4}S (n = 1–5), had a marked effect on the monooxygenase activity of PUPPET (Figure 5). Though the dissociation of PCNA3-P450cam from PUPPET is known to have a negative effect on the activity [13,17], the effect of the linker length was not involved in this dissociation, because the relationship between concentration and activity of PUPPET was not affected by the linker variants (Figure 6B). The poly-L-proline linker has been used as a “molecular ruler”, because poly-L-proline is expected to form an all-trans type II helix and the length of the helix can be predicted as approximately 3 Å per residue [21,35]. Recent studies demonstrated that the end-to-end distances of the poly-L-proline peptides are shorter than those of perfectly rigid type II proline helices, probably because of the small number of cis isomers interspersed in aqueous solutions of poly-L-proline peptides [36–39]. Nevertheless, the end-to-end distance was reported to increase with an increase in the number of proline residues, up to at least 24 amino acids [38]. Therefore, the repeat number of the Pro residue in the linker is expected to control the distance between PCNA2 and PdX, and to subsequently affect the spatial arrangement of PdX relative to P450cam in PUPPET-P\textsubscript{n}. The strong dependence of the monooxygenase activity on the repeat number of the Pro residue in the G\textsubscript{2}S(P\textsubscript{3}),G\textsubscript{2}S linker indicates that the relative spatial arrangement of PdX governs the electron donation step from reduced PdX to P450cam in the PUPPETs.

A previously reported docking model of the P450cam-PdX electron transfer complex [40], demonstrated that the distance between Leu11 of P450cam (of which residues Met1–Asn10 are undefined in the crystal structure), and the N-terminus of PdX, is approximately 65 Å in the electron transfer complex (Figure 7A). We also estimated that the mean end-to-end distance of helical linker Pro\textsubscript{20} was approximately 50 Å according to a previous study [38]. PdX and P450cam are located on the same side of the PCNA ring because they are fused to the C-termini of PCNA2 and PCNA3, respectively. By combining the above structural information, we suggest a model of the electron transfer complex, in which the PdX-binding site of P450cam faces in a perpendicular direction towards the PCNA ring (Figure 7B). In this model, the G\textsubscript{4}S(P\textsubscript{5})\textsubscript{n}G\textsubscript{4}S linker between the C-terminus of PCNA2 and the N-terminus of PdX is the most effective in allowing PdX close to the P450cam binding site. On the contrary, the speculation that the binding site faces in a parallel direction to the PCNA ring as shown in Figure 7C, conflicts with the experimental data showing that the activity of PUPPET-P\textsubscript{n} increased with an increase in the distance between the N-terminus of PdX and the C-terminus of PCNA2, in the range of n = 1–4. Therefore, the PdX-binding site of P450cam should preferably face in the perpendicular direction to the PCNA ring in the complex.

Although the length of the Gly-Ser repeating linker, (G\textsubscript{3}S\textsubscript{n})\textsubscript{1–6}, also affected the monooxygenase activity, these linkers did not allow these PUPPET complexes to achieve the activity levels of even PUPPET-P\textsubscript{2} (Figure 5). This result suggests that the mean end-to-end distances of the (G\textsubscript{3}S\textsubscript{n}) linkers are shorter than that of G\textsubscript{3}S(P\textsubscript{5})\textsubscript{10}G\textsubscript{2}S. Evers et al. demonstrated that the elongation of a Gly- and Ser-rich (GGSGGS)\textsubscript{n} linker from n =
1–6 only increased the distance between the two linked proteins by approximately 6 Å [41]. The Gly_{n}\text{-Ser} repeating linker probably has similar characteristics to the (GGS)_{n} linker. Therefore, the flexible (G_{4}S_{4}) linkers do not enable PdX to be located at the appropriate distance away from the PCNA ring in order for the electron transfer complex to form, as shown in Figure 7B, and may instead cause PdX to interact with P450cam in an unfavorable manner, as shown in Figure 7C. The difference in the possible spatial arrangements of PdX in PUPPET-P_{n} and PUPPET-G_{n} might therefore result in the differences observed in their activities.

Conclusion

In this study, we successfully enhanced the monooxygenase activity of the PCNA-mediated protein complex of PdR, PdX and P450cam by optimizing the peptide linker between PCNA2 and PdX. The linker affected the electron donation step from reduced PdX to P450cam in the complex. The G_{2}S_{2}P_{4}S linker most effectively enhanced the activity, most likely because the rigid Pro_{20} sequence contributed to PdX positioning close to the PdX-binding site of P450cam. On the other hand, the high activity obtained with this linker was not achieved with the flexible (G_{n}S_{n}) linkers in the range of n = 1–6, owing to the short end-to-end distances of these flexible linkers. Recently, we have demonstrated that introduction of disulfide bonds between the PCNA subunits prevented the dissociation of PCNA3-P450cam from PUPPET and enhanced the activity [17]. Therefore, introduction of disulfide bonds can further enhance the activity of PUPPET-P_{20}. Our findings suggest that linkers containing rigid peptides are more suitable for the control of the spatial arrangement of enzymes in artificial multienzyme complexes in which component enzymes/proteins need to interact with each other, and can thereby result in improved catalytic activity.

Supporting Information

Figure S1. UV–Vis spectra of PCNA2-PdX linker variants. UV–Vis spectra of PCNA2-G_{n}S(P_{4})_{n}G_{4}S-PdX (n = 1–5, P_{1–P_{5}}) and PCNA2-(G_{n}S)_{n}-PdX (n = 2–6, G_{2–G_{6}}) are listed. (TIF)

Figure S2. UV–Vis spectra of PUPPET linker variants. Solid lines indicate each PUPPET-P_{n} (n = 1–5, P_{1–P_{5}}) and PUPPET-G_{n} (n = 2–6, G_{2–G_{6}}) spectrum. Broken lines indicate a linear combination of the individual component protein spectra. (TIF)

Table S1. Oligo DNA sequences to insert peptide linkers between PCNA2 and PdX. (DOC)

Author Contributions

Conceived and designed the experiments: TH HH. Performed the experiments: TH HH. Analyzed the data: TH HH. Contributed reagents/materials/analysis tools: TH HH. Wrote the manuscript: TH HH TN. Interpreted the data: TH HH TN.

References

1. Schoffelen S, Van Hest JCM (2012) Multi-enzyme systems: bringing enzymes together in vitro. Soft Matter 8: 1736–1746. doi: 10.1039/c1sm06452e.
2. Delebecque CJ, Lindner AB, Silver PA, Aldaye FA (2011) Organization of intracellular reactions with rationally designed RNA assemblies. Science 333: 470–474. doi: 10.1126/science.1206938. PubMed: 21708039.
3. Conrado RJ, Wu GC, Bocko JT, Xu H, Chen SY et al. (2012) DNA-guided assembly of biosynthetic pathways promotes improved catalytic efficiency. Nucleic Acids Res 40: 1879–1889. doi: 10.1093/nar/gkr888. PubMed: 22021385.
4. Ermel Hon Y, Kuo CH, Hsu FY, Lin CY, Lee WJ et al. (2012) DNA-mediated assembly of cytochrome P450 BM3 subdomains. J Am Chem Soc 133: 16111–16118. doi: 10.1021/ja204993s. PubMed: 21919448.
5. Fu J, Liu M, Yu Y, Woodbury NW, Yan H (2012) Interezyme substrate diffusion for an enzyme cascade organized on spatially addressable DNA nanostructures. J Am Chem Soc 134: 5516–5519. doi: 10.1021/ja300897h. PubMed: 22414276.
6. Müller J, Niemeyer CM (2008) DNA-directed assembly of artificial multienzyme complexes. Biochem Biophys Res Commun 377: 62–67. doi: 10.1016/j.bbrc.2008.07.078. PubMed: 18623945.
7. Wilner Ol, Weizmann Y, Gill R, Lioubashevski O, Freeman R et al. (2009) Enzyme cascades activated on topologically programmed DNA scaffolds. Nat Nanotechnol 4: 249–254. doi: 10.1038/nnano.2009.50. PubMed: 19350036.
8. Agapakis CM, Ducat DC, Boyle PM, Wintemute EH, Way JC et al. (2010) Insulation of a synthetic hydrogen metabolism circuit in bacteria. J Biol Eng 4: 3. doi: 10.1186/1754-1611-4-3. PubMed: 20184755.
9. Caspi J, Barak Y, Haimovitz R, Irwin D, Lamed R et al. (2009) Effect of linker length and dockerin position on conversion of a Thermobifida fusca endoglucanase to the cellulosomal mode. Appl Environ Microbiol 75: 7335–7342. doi: 10.1128/AEM.01241-09. PubMed: 19820150.
10. Dueber JE, Wu GC, Malmirchegini GR, Moon TS, Petzold CJ et al. (2009) Synthetic protein scaffolds provide modular control over metabolic flux. Nat Biotechnol 27: 753–759. doi: 10.1038/nbt.1557. PubMed: 19648909.
11. You C, Myung S, Zhang Y-HP (2012) Facilitated substrate channeling in a self-assembled trifunctional enzyme complex. Angew Chem Int Ed 51: 8787–8790. doi: 10.1002/anie.201202441.
12. You C, Zhang Y-HP (2013) Self-Assembly of Synthetic Metabolons through Synthetic Protein Scaffolds: One-Step Purification, Co-immobilization, and Substrate Channeling. ACS Synth Biol 2: 102–110. doi: 10.1021/sb300066y. PubMed: 23656373.
13. Hirakawa H, Nagamune T (2010) Molecular assembly of P450 with ferredoxin and ferredoxin reductase by fusion to PCNA. Chembiochem 11: 1517–1520. doi: 10.1002/cbic.2010000226. PubMed: 20607777.
14. Dionne I, Nookala RK, Jackson SP, Doherty AJ, Bell SD (2003) A heterotrimetric PCNA in the hyperthermophilic archaeon Sulfolobus solfataricus. Mol Cell 11: 275–282. doi: 10.1016/S0977-2765(02)00824-9. PubMed: 12535540.
15. Dionne I, Brown NJ, Woodgate R, Bell SD (2008) On the mechanism of loading the PCNA sliding clamp by RFC. Mol Microbiol 68: 216–222. doi: 10.1111/j.1365-2958.2008.06150.x. PubMed: 18312273.
16. Hirakawa H, Kakitani A, Nagamune T (2013) Introduction of selective intersubunit disulfide bonds into self-assembly protein scaffold to enhance an artificial multienzyme complex's activity. Biotechnol Bioeng 110: 1858–1864. doi: 10.1002/bit.24861. PubMed: 23404255.
17. Arora PS, Ansari AZ, Best TP, Ptashne M, Dervan PB (2002) Design of artificial transcriptional activators with rigid poly-L-proline linkers. J Am Chem Soc 124: 13067–13071. doi: 10.1021/ja0208355. PubMed: 12405833.
18. Bianco-Canosa JB, Medintz IL, Farrell D, Mattoussi H, Dawson PE (2010) Rapid covalent ligation of fluorescent peptides to water
solvabilized quantum dots. J Am Chem Soc 132: 10027–10033. doi: 10.1021/ja107447w. PubMed: 20973474.

20. Sato S, Kwon Y, Kamisuki S, Srivastava N, Mao Q et al. (2007) Polyproline-rod approach to isolating target proteins of bioactive small molecules: isolation of a new target of indomethacin. J Am Chem Soc 129: 783–788. doi: 10.1021/ja065566x. PubMed: 17243824.

21. Streyer L, Haugland RP (1967) Energy transfer: a spectroscopic ruler. Proc Natl Acad Sci U S A 58: 719–726. doi: 10.1073/pnas.58.2.719. PubMed: 5233469.

22. Tanaka T, Nomura W, Narumi T, Masuda A, Tamamura H (2010) Bivalent ligands of CXCR4 with rigid linkers for elucidation of the dimerization state in cells. J Am Chem Soc 132: 15899–15901. doi: 10.1021/ja107447w. PubMed: 20973474.

23. Huston JS, Levinson D, Mudgett-Hunter M, Tai MS, Novotny J et al. (1988) Protein engineering of antibody binding sites: recovery of specific activity in an anti-digoxin single-chain Fv analogue produced in Esherichia coli. Proc Natl Acad Sci U S A 85: 5879–5883. doi: 10.1073/pnas.85.16.5879. PubMed: 3045807.

24. Arai R, Ueda H, Kiyama A, Kamiya N, Nagamura T (2001) Design of the linkers which effectively separate domains of a bifunctional fusion protein. Protein Eng 14: 529–532. doi: 10.1093/protein/14.8.529. PubMed: 11579220.

25. Arai R, Wiggers W, Nishikawa Y, Nagamura T, Fujisawa T (2004) Conformations of variably linked chimeric proteins evaluated by synchrotron X-ray small-angle scattering. Proteins 57: 829–838. doi: 10.1002/prot.20244. PubMed: 15390267.

26. Lu P, Feng MG (2008) Bifunctional enhancement of a beta-glucanase-xylanase fusion enzyme by optimization of peptide linkers. Appl Microbiol Biotechnol 79: 579–587. doi: 10.1007/s00253-008-1468-4. PubMed: 16415085.

27. Roome PW, Peterson JA (1988) The oxidation of reduced putidaredoxin reductase by oxidized putidaredoxin. Arch Biochem Biophys 266: 41–50. doi: 10.1016/0003-9861(88)90234-2. PubMed: 2845865.

28. Wiggers W, Chakravarty S, Jennings PA (2005) Control of protein functional dynamics by peptide linkers. Biopolymers 80: 736–746. doi: 10.1002/bip.20291. PubMed: 15880774.

29. Chen X, Zaro JL, Shen W-C (2013) Fusion protein linkers: Property, design and functionality. Adv Drug Deliv Rev (2012). doi: 10.1016/j.addr.2012.09.039. PubMed: 23026637.

30. Anthis NJ, Clore GM (2013) The length of the calmodulin linker determines the extent of transient interdomain association and target affinity. J Am Chem Soc 135: 9648–9651. doi: 10.1021/ja405142z. PubMed: 23782151.

31. Krishnamurthy VM, Semetye V, Bracher PJ, Shen N, Whitesides GM (2007) Dependence of effective molarity on linker length for an intramolecular protein-ligand system. J Am Chem Soc 129: 1312–1320. doi: 10.1021/ja066796e. PubMed: 17253415.

32. Albertsen L, Chen Y, Bach LS, Rattleff S, Maury J et al. (2011) Diversion of flux toward sesquiterpene production in Saccharomyces cerevisiae by fusion of host and heterologous enzymes. Appl Environ Microbiol 77: 1033–1040. doi: 10.1128/AEM.01361-10. PubMed: 21146687.

33. Carlsson H, Ljung S, Bubow L (1996) Physical and kinetic effects on induction of various linker regions in beta-galactosidase/galactose dehydrogenase fusion enzymes. Biochim Biophys Acta 1293: 154–160. doi: 10.1016/0167-4838(95)00240-5. PubMed: 8652521.

34. Brewer CB, Peterson JA (1988) Single turnover kinetics of the reaction between oxyctochrome P-450cam and reduced putidaredoxin. J Biol Chem 263: 791–798. PubMed: 2826462.

35. Cowan PM, McGavin S (1955) Structure of Poly-L-Proline. Nature 176: 501–503. doi: 10.1038/176501a0. PubMed: 15880774.

36. Evers TH, Van Dongen EMWM, Faesen AC, Meijer EW, Merkx M (2006) Structural distributions from single-molecule measurements as a tool for molecular mechanics. Chem Phys 396: 61–71. doi: 10.1016/j.chemphys.2011.06.014. PubMed: 22681822.

37. Watkins LP, Chang H, Yang H (2008) Quantitative single-molecule conformational distributions: a case study with poly-(L-proline). J Phys Chem A 110: 5191–5203. doi: 10.1021/jp075588d. PubMed: 16610843.

38. Kuznetsov VY, Poulos TL, Sverrioukova IF (2006) Putidaredoxin-to-cytochrome P450cam electron transfer: differences between the two reductive steps required for catalysis. Biochemistry 45: 11933–11944. doi: 10.1021/bi0611154. PubMed: 17002293.

39. Tochigrechko A, Vakser I A (2006) GRAMM-X public web server for protein-protein docking. Nucleic Acids Res 34: W310–W314. doi: 10.1093/nar/gkl099. PubMed: 16845016.