

# Protein engineering for molecular electronics

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Recombinant DNA technology allows the manipulation of the physical properties of proteins that perform electron transport and photochemical processes. Recent work is reviewed that has a potential impact on the development of molecular electronic devices within a general framework outlining strategies for device fabrication. This review is also published in *Current Opinion in Biotechnology* 1992, 3:388–394.

*Current Opinion in Structural Biology* 1992, 2:587–592

## Introduction

This review is prompted by the recent enthusiasm for the use of biological macromolecules as building blocks for the construction of molecular electronic devices (MEDs). The term molecular electronics has two popular definitions. One refers to electronic devices that use materials whose unique properties result from their molecular structure. In this regard, proteins can be used in MEDs to sense, respond to, or record chemical, electrical or physical stimuli. Examples include the use of the bacterial photopigment bacteriorhodopsin as a three-dimensional medium for the storage and readout of optically encoded information [1,2,3]. The second definition of molecular electronics embodies the concepts that individual molecules are functional units responding to stimuli, and that they have the potential to be interconnected in order to functionally replicate electronic circuits. Good examples of such MEDs do not yet exist, but the principles necessary for their design and assembly are beginning to emerge from a variety of areas.

From the perspective of practical device design, there are several levels at which the ability to modify protein molecules using recombinant DNA technology could have an impact on the fabrication of MEDs. These include the alteration of intrinsic optical and electronic properties, increasing protein stability for operation in non-biological environments, and the modification of surface properties to facilitate *de novo* design of mesoscale molecular assemblies or molecular circuits. Current activities in each of these categories are described below.

## Modification of protein electronic and physical properties

Naturally occurring proteins incorporate a wide range of physical properties that are of potential use in electronic devices. In most cases, the useful properties derive from a prosthetic group such as a metal center, organic redox cofactor, or chromophore. Although naturally oc-

curing redox and photoactive proteins exhibit great functional diversity, this is achieved through a limited number of prosthetic groups whose properties are modulated through interaction with amino acid side chains of the surrounding protein moiety. Attempts to engineer proteins in order to modify their properties in useful ways have followed precedents suggested by the study of structure–function relationships in natural systems. In this respect, heme-containing proteins, which comprise a diverse set of molecules with electron-transfer, ligand-binding or catalytic function, are of particular interest. Alterations in the electronics or chemistry of the heme iron center frequently produce large changes in optical spectroscopic or magnetic properties which are useful as indicators for state assignment or device readout. Following pioneering work on cytochrome *b5* [4], myoglobin [5–8], and cytochrome *p450* [9], recent work reporting site-directed modifications of residues that are heme iron ligands, or can otherwise influence heme spectral properties, included additional studies on cytochrome *p450* [10] and myoglobin [11], together with studies on cytochrome *c* peroxidase [12,13] and iso-1-cytochrome *c* [14]. Although the study of the structure–property relationships that affect heme proteins remains an area of active interest [15], few systematic efforts have been directed at producing unusually stable molecules that might increase the reliability of state assignment. However, a serendipitous enhancement of protein stability was reported for a site-directed mutant of iso-1-cytochrome *c* [14], in which an internal asparagine had been changed to a hydrophobic isoleucine residue. This alteration resulted in the loss of an internal water molecule whose location in the interior of the native protein was postulated to be energetically unfavorable. Similar hydrophobic enhancements might direct further efforts towards the important goal of stabilizing redox proteins in non-aqueous environments.

A property central to the function of many MED devices is the regulation of electron-transfer rate between proteins or between proteins and an external oxidoreductant. Physical aspects of the electron-transfer pro-

## Abbreviation

MED—molecular electronic device.

cesses in proteins are now relatively well understood, and suggest that electron-transfer rates depend primarily upon prosthetic group separation, differences in oxidation-reduction potential and molecular reorganization energy [16,17]. This is consistent with recent site-directed modifications of the invariant Phe82 residue in yeast iso-1-cytochrome *c*, which suggested that intermolecular electron transfer does not require the participation of specific aromatic amino acids as electron wires between protein prosthetic groups [18,19]. Nevertheless, translation of the physical requirements for efficient electron transfer into a specific modification strategy remains complicated, as shown by the results obtained by Baker *et al.* for yeast iso-1-cytochrome *c* [20]. In addition to heme-containing redox proteins, the introduction or regeneration of copper-binding sites in proteins has also been investigated. In one study, both type I and type II copper-site properties were obtained by addition of an appropriate external ligand to an azurin mutant in which one of the native histidine copper ligands had been deleted by site-directed mutagenesis [21]. In a second study, site-directed modifications of cytochrome *c* were performed in order to introduce a copper ligand site [22]. In both cases, the modified proteins appeared to lack the intrinsic stability of the native molecule, but may point the way to successive generations of molecules with useful functions.

### Interaction specificity

Because rates of intermolecular transfer depend very strongly on prosthetic group separation distance [16,17], reactions between reversibly binding electron-transfer proteins depend on interactions that ensure the proper relative intermolecular orientation. Early work that modeled the interactions of cytochromes *c* and *b5* established the importance of solvent exclusion and complementary electrostatic interactions in redox protein interactions [23]. This relationship has since been extended to many other biological electron-transfer interactions. With the advent of site-directed mutagenesis methods, investigation of intermolecular recognition interactions has intensified [24–29]. Most notably, methods have recently been developed that discriminate between the relative contributions of the principal components of intermolecular recognition: polar interactions that result from hydrogen bonds and salt bridges forming in the interaction domain; and, non-polar van der Waals interactions that occur when protein surfaces dehydrate upon formation of the complementary complex [30]. This methodology also facilitates mapping the interaction domain between two associating macromolecules, a key step in the development of generalized strategies for both studying natural systems and engineering alternative interactions among molecules suitable for use in MEDs.

### Bacteriorhodopsin in MED applications

Bacteriorhodopsin (molecular weight  $\approx 26.0$  kD) functions as a light-driven proton pump in the purple membrane of the salt-marsh micro-organism *Halobacterium halobium* [1,2]. Bacteriorhodopsin incorporates a reti-

nal chromophore, which is covalently bound as a protonated Schiff base in an all-*trans* conformation in the resting state. Upon absorbing a light photon, the retinal photoisomerizes and subsequently undergoes a multi-step photocycle which spans much of the visible spectrum. The photocycle can be stopped at specific intermediates by low-temperature trapping, and recycled by thermalization or the application of a second light pulse of the appropriate wavelength [1,2]. Photoactivated retinal switching occurs with high quantum yields at disparate wavelengths, which facilitates state assignment, and produces both changes in protein refractive index and a photoelectric potential in oriented assemblies. These properties, together with the excellent stability of the protein when immobilized in polymer films or gels, form the basis for a variety of prototypes for data storage media, holographic memory, and electro-optic applications [1,2,3,31,32]. Three-dimensional photochromic memories have incorporated bacteriorhodopsin that has been oriented by an external electric field and then immobilized in a solid polyacrylamide matrix [2]. This application exploits the two-photon absorption cross section of bacteriorhodopsin and requires a small volume of the matrix to be simultaneously illuminated by intersecting laser beams for data storage.

Readout is achieved by reillumination, which produces an electrical signal which depends on the state of the irradiated volume in the data-storage matrix. In this context, it is interesting to note recent experiments demonstrating the maintenance of redox-protein properties when immobilized in silicate glasses [33], as well as the preservation and control of proteolytic enzyme activity in a photochromic azobenzene copolymer [34]. Although most readily applied in biosensor applications, both schemes could potentially be useful in three-dimensional optical memory applications which make use of photochromic properties of incorporated proteins, or couple protein catalytic properties to the photochromic matrix.

Current work on bacteriorhodopsin with relevance to molecular electronics involves finding additives, alternative pigments, or site-directed mutants of the protein that will alter the relative stabilities of the photo-intermediates [35], enhance stabilization of the final intermediate at room temperature [36], or alter other aspects of photocycle-coupled proton translocation [37,38] that might facilitate readout from information-storage applications. Consequently, investigations of the Asp96Asn modification [39], which alters coupling of retinal photoisomerization to proton translocation, continue to have substantial practical interest [40]. The physical changes in the modified protein include an increase in the lifetime of the intermediate from 10 to 750 ms, together with improved diffraction efficiency and photochromic sensitivity relative to the native protein. Considerable latitude for property improvement exists using a combination of site-directed modifications at the chromophore binding site, introduction of alternative chromophores, or the engineering of modified ion-binding sites, all of which can affect intermediate lifetimes and spectral properties [1,2]. In this regard, the recent structure determination of bac-

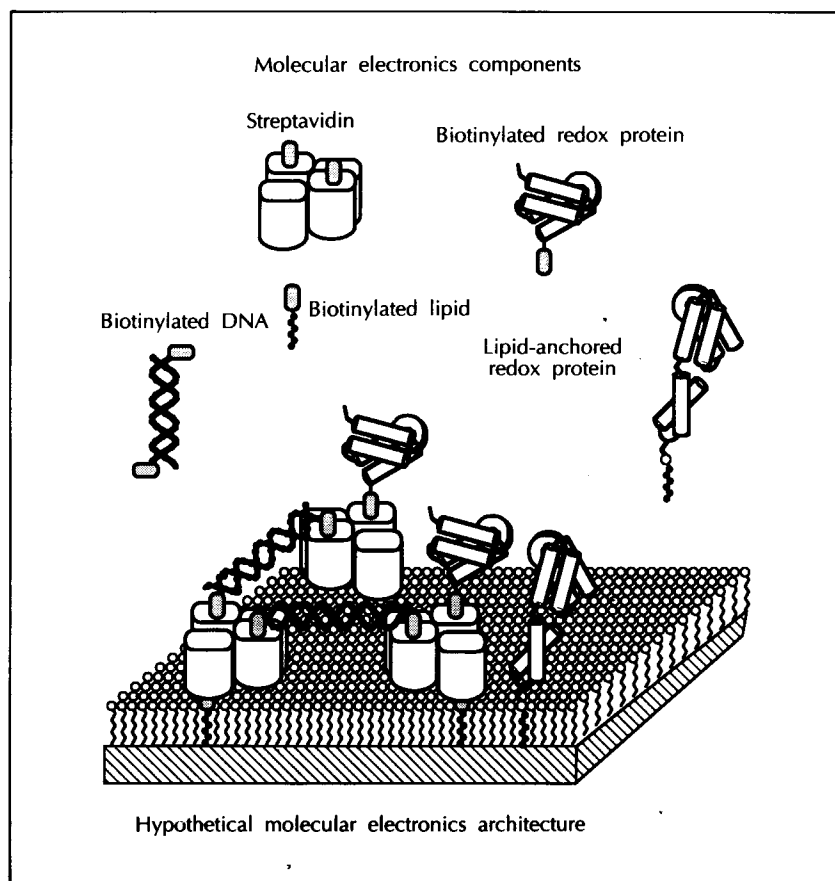
teriorhodopsin by electron diffraction methods [41•], coupled with rapidly advancing methods of computational simulation [42•], may provide new insights into the engineering of bacteriorhodopsin for MED applications.

### Oriented protein immobilization on surfaces

Bacteriorhodopsin is not unique in its ability to spontaneously order in two-dimensional films that resemble its native membrane state and, indeed, other membrane protein systems have been suggested for use or have potential in a variety of MED or related biosensor roles. Recent work includes studies of engineered bacterial transmembrane pores [43] and isolated bacterial reaction centers immobilized on electrode surfaces [44]. In these cases, surface orientation of the protein complexes results from preexisting interactions between the protein complexes and/or their membrane environments. However, a number of experiments have been carried out that involve tethering proteins to electrode surfaces through polymeric 'wires' [45,46] or through surface electrostatic interactions [47] in order to enhance electron-transfer efficiencies. Related studies of interest have examined those properties of soluble proteins conjugated to photo- or redox-active organic substituents that enhance enzyme catalytic function in the absence of natural cofactors or regeneration systems [48–50]. It is easy to envisage hybrid systems that would incorporate proteins with conjugated cofactors immobilized on electrode surfaces. Although the studies described

above have emphasized electron conduction between device components, the efficiency of many devices incorporating prosthetic groups in sensor or non-linear optical applications depends critically on the ability to precisely control prosthetic group orientation relative to the electrode or optical waveguide surface. The site-specific introduction of anchoring points in a protein of known three-dimensional structure should provide a straightforward solution to this problem. As an example, recent site-directed modifications of the heme protein cytochrome *b5* carried out in the Biomolecular Electronics Group in the Beckman Institute at the University of Illinois introduced cysteine residues at specified positions on the molecular surface. This allowed oriented coupling to a silane substrate and produced immobilized molecular arrays demonstrating a high level of heme prosthetic group orientation, as determined by linear dichroism spectroscopy.

Methods to introduce two-dimensional periodicity onto surface molecular arrays depend either on engineering both intermolecular interactions [30•] and anchoring sites for surface immobilization, or on schemes that erect molecular assemblies on a scaffold already possessing periodic order on the few tens of Ångstroms scale. Scaffold possibilities include natural two-dimensional lattices [43], DNA lattices [51•], or the very highly ordered synthetic lattices formed from streptavidin anchored to surfaces through biotinylated phospholipids [52•]. As streptavidin is a tetrameric protein with four biotin-binding sites arranged with approximate tetrahedral geometry, two biotin-binding sites per tetramer remain free in



**Fig. 1.** Illustration of molecular electronics components and potential architectural features of self-assembling systems restricted to diffusion in two-dimensional lipid films.

the two-dimensional arrays and would provide a regular lattice for attaching additional molecules (Fig. 1).

### Strategies for protein patterning and circuit fabrication

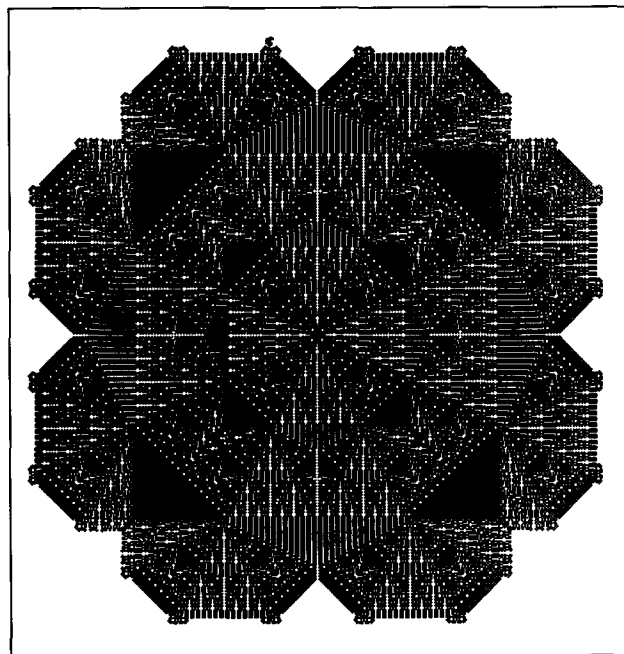
Although it appears possible that quite complicated molecular assemblies could be erected on two-dimensional protein lattices using a combination of chemical conjugation, site-directed introduction of anchor points, and molecular fusions at the DNA level (Nilsson *et al.*, pp 569–575), the materials described thus far all come under our first definition of molecular electronics as bulk materials. Although these substrates could undoubtedly be patterned using lithography methods similar to those used in fabricating integrated circuits, optical diffraction would limit detail to scales substantially larger than individual molecular assemblies. Such devices would not realize the ultimate in miniaturization of individually assignable devices.

Molecular electronics of the second definition, where individual molecular assemblies have unique addressability and specific connectivity between components, would be a major advance. Nevertheless, it is interesting to consider how it might be possible to construct true molecular circuits, as well as the dual problem at the molecular level of how such devices could be addressed and read. Atomic probe microscopy could potentially be used to manipulate and assemble molecules as a means of constructing devices, as well as a means of setting molecular states or reading out information [53•]. The limitations of manipulating only one or a few components at a time, however, would seem ultimately to defeat the objectives of creating a molecular device.

Alternatively, it may be possible to 'condition' a pre-existing molecular lattice of uniformly interconnected molecules so that it is able to store or process information as a neural network analog. Information processing using a distributed approach requires each molecular assembly to sense the state of its nearest neighbors and alter its configuration accordingly. With phasing to an external clock, this scheme has the formal structure of a cellular automata operating with individual macromolecules as building blocks. In such a highly cooperative device, it may not be necessary to directly manipulate individual molecules; rather, one might use intermolecular communications common to many cooperative multimeric proteins to perform some analog computing function.

Fabrication possibilities also exist using defect tessellation automata [54•], where very complicated structures (Fig. 2) can be generated by defect introduction into a periodic lattice subject to a regular site-replacement protocol. Although presently only the subject of computer simulation, it seems possible to embody the required properties in ligand-capture and displacement schemes using symmetric protein molecules with multiple binding sites. Access and readout could potentially be achieved by any of the methods outlined above. Although these devices are presently wholly conceptual, and may ultimately be limited by molecular noise that will necessitate distributed or redundant processing to insure reliability, it

seems clear that genuine approaches exist to investigate the limits of this technology.



**Fig. 2.** A computer simulation showing a finite tessellation automata derived from a single defect. This displays connectivity reminiscent of integrated circuits. Published with permission [54•].

### Conclusion

The practical application of engineered proteins in molecular electronics and sensor materials applications are clearly on the near horizon. Protein molecules represent the ultimate miniaturization possible in individual devices whose state can potentially be controlled independently of its neighbors. This property, together with the emergence of a knowledge base that allows protein properties and interaction specificity to be engineered with relative ease, will undoubtedly lead to progressively more sophisticated assemblies, whose functional limitations can only now be guessed at. Multidisciplinary programs that pioneered development of bacteriorhodopsin-based devices [1•,3], or the Frontier Research Program of the Riken Institute in Japan [55•], represent focused efforts to realize the potential of this technology.

### References and recommended reading

Papers of particular interest, published within the annual period of review, have been highlighted as:

- of special interest
  - of outstanding interest
1. OESTERHELT D, BRAUCHLE C, HAMPP N: **Bacteriorhodopsin: a Biological Material for Information Processing.** *Q Rev Biophys* 1991, 4:425–478.
  2. BIRGE RR: **Photophysics and Molecular Electronic Applications of the Rhodopsins.** *Annu Rev Phys Chem* 1990, 41:683–733.

A recent review that summarizes structure–function relationships in bacteriorhodopsin and includes a discussion of site-specific mutants.

This review emphasizes MED applications.

3. BELCHINSKY GY, BOGATOV PN, GRESKO AP, KONENKO AA, LUKASHEV EP, CHAMOROVSKY SK: Application of Pigment-Protein Complexes as Carriers of Optic Information. *Biofizika* 1991, 36:248-251.
  4. SLIGAR S, EGERBERG K, SAGE T, MORIKIS D, CHAMPION P: Alteration of Heme Axial Ligands by Site Directed Mutagenesis: a Cytochrome Becomes a Catalytic Demethylase. *J Am Chem Soc* 1987, 109:7896-7897.
  5. LAMBRIGHT DG, BALASUBRAMANIAN S, BOXER SG: Ligand and Proton Exchange Dynamics in Recombinant Human Myoglobin Mutants. *J Mol Biol* 1989, 207:289-299.
  6. VARADARAJAN R, ZEWERT TE, GRAY HB, BOXER SG: Effects of Buried Ionizable Amino Acids on the Reduction Potential of Recombinant Myoglobin. *Science* 1989, 243:69-72.
  7. VARADARAJAN R, LAMBRIGHT DG, BOXER SG: Electrostatic Interactions in Wild-type and Mutant Recombinant Human Myoglobins. *Biochemistry* 1989, 28:3771-3781.
  8. EGERBERG KD, SPRINGER BA, MARTINIS SA, SLIGAR SG, MORIKIS D, CHAMPION PM: Alteration of Sperm Whale Myoglobin Axial Ligation by Site-directed Mutagenesis. *Biochemistry* 1990, 29:9783-9791.
  9. DI PRIMO C, HUI BON HOA, DOUZOU P, SLIGAR S: Mutagenesis of a Single Hydrogen Bond in Cytochrome P-450 Alters Cation Binding and Heme Solvation. *J Biol Chem* 1990, 265:5361-5363.
  10. IWASAKI M, JUVONEN R, LINDBERG R, NEGISHI M: Alteration of High and Low Spin Equilibrium by a Single Mutation of Amino Acid-209 in Mouse Cytochrome-P450. *J Biol Chem* 1991, 266:3380-3382.
  11. ADACHI S, NAGANO S, WATANABE Y, ISHIMORI K, MORISHIMA I: Alteration of Human Myoglobin Proximal Histidine to Cysteine or Tyrosine by Site-directed Mutagenesis—Characterization and Their Catalytic Activities. *Biochem Biophys Res Commun* 1991, 180:138-144.
  12. GOODIN DB, DAVIDSON MG, ROE JA, MAUK AG, SMITH M: Amino Acid Substitutions at Tryptophan-51 of Cytochrome-C Peroxidase—Effects on Coordination, Species Preference for Cytochrome-c, and Electron Transfer. *Biochemistry* 1991, 30:4953-4962.
  13. SMULEVICH G, MILLER MA, KRAUT J, SPIRO TG: Conformational Change and Histidine Control of Heme Chemistry in Cytochrome-c Peroxidase—Resonance Raman Evidence from Leu-52 and Gly-181 Mutants of Cytochrome-c Peroxidase. *Biochemistry* 1991, 30:9546-9558.
  14. HICKEY DR, BERGHUIS AM, LAFOND G, JAEGER JA, CARDILLO TS, MCLENDON D, DAS G, SHERMAN F, BRAYER GD: Enhanced Thermodynamic Stabilities of Yeast Iso-1-cytochrome-c with Amino Acid Replacement at Position-52 and Position-102. *J Biol Chem* 1991, 266:11686-11694.
  15. FRAUENFELDER H, SLIGAR SG, WOLYNES PG: The Energy Landscapes and Motions of Proteins. *Science* 1991, 254:1598-1603.
- The authors describe current concepts about dynamic interconversion among conformational substates in proteins and the resulting physical manifestations. This is of fundamental importance in understanding how thermal phenomena will affect switching reliability in protein molecules when excited by external stimuli.
16. MOSER CC, KESKE JM, WARNCHE K, FARID RS, DUTTON PL: Nature of Biological Electron Transfer. *Nature* 1992, 355:796-802.
- This paper summarizes much experimental data showing that electron-transfer rates effectively depend on prosthetic group separation distance, differences in prosthetic-group free energy and reorganization energies upon oxidoreduction. The treatment suggests that the protein presents a uniform electronic tunnelling barrier that derives from glass-like protein properties also revealed in dynamic studies.
17. BERATAN DN, ONUCHI JN, BETTS JN, BOWLER BE, GRAY HB: Electron Tunneling Pathways in Ruthenated Proteins. *J Am Chem Soc* 1990, 112:7915-7921.
  18. EVEREST AM, WALLIN SA, STEMP ED, NOCEK JM, MAUK AG, HOFFMAN B: Aromatic Hole Superexchange Through Position-82 in Cytochrome C is Not Required for Intracomplex Electron Transfer to Zn-Cytochrome C Peroxidase. *J Am Chem Soc* 1991, 113:4337-4338.
  19. INGLIS SC, GUILLETTE JG, JOHNSON JA, SMITH M: Analysis of the Invariant Phe82 Residue of Yeast Iso-1-cytochrome c by Site-directed Mutagenesis Using a Phagemid Yeast Shuttle Vector. *Protein Eng* 1991, 4:569-574.
  20. BARKER PD, MAUK MR, MAUK AG: Proton Titration Curve of Yeast Iso-1-Ferricytochrome-c—Electrostatic and Conformational Effects of Point Mutations. *Biochemistry* 1991, 30:2377-2383.
  21. DENBLAAUWEN T, VANDEKAMP M, CANTER GW: Type-I and Type-II Copper Sites Obtained by External Addition of Ligands to a His117Gly Azurin Mutant. *J Am Chem Soc* 1991, 113:5050-5052.
  22. TODD RJ, VANDAM ME, CASIMIRO D, HAYMORE BL, ARNOLD FH: Cu(II)-binding Properties of a Cytochrome-c with a Synthetic Metal-binding Site—His-X<sub>3</sub>-His in an Alpha-helix. *Proteins* 1991, 10:156-161.
  23. SALEMM FR: A Hypothetical Structure for an Intermolecular Electron Transfer Complex of Cytochrome c and b5. *J Mol Biol* 1976, 102:563-568.
  24. RODGERS K, POCHAPSKY T, SLIGAR S: Probing the Mechanisms of Macromolecular Recognition: the Cytochrome b5-Cytochrome C Complex. *Science* 1988, 240:1657-1659.
  25. STAYTON PS, SLIGAR SG: The Cytochrome P-450cam Binding Surface as Defined by Site-directed Mutagenesis and Electrostatic Modeling. *Biochemistry* 1990, 29:7381-7386.
  26. SHIMIZU T, TATEISHI T, HATANO M, FUJIKURIYAMA Y: Probing the Role of Lysines and Arginines in the Catalytic Function of Cytochrome-P450d by Site Directed Mutagenesis—Interaction with NADPH-cytochrome-P450 Reductase. *J Biol Chem* 1991, 266:3372-3375.
  27. COGHIAN VM, VICKERY LE: Site-specific Mutations in Human Ferredoxin that Affect Binding to Ferredoxin Reductase and Cytochrome-P450<sub>osc</sub>. *J Biol Chem* 1991, 266:18606-18612.
  28. SHIMIZU T, TATEISHI T, HATANO M, FUJIKURIYAMA Y: Probing the Role of Lysines and Arginines in the Catalytic Function of Cytochrome-P450D by Site-directed Mutagenesis—Interaction with NADPH-cytochrome-P450 Reductase. *J Biol Chem* 1991, 266:3372-3375.
  29. SHIRABE K, YUBISUI T, NISHINO T, TAKESHITA M: Role of Cysteine Residues in Human NADH-cytochrome-b5 Reductase Studies by Site-directed Mutagenesis—Cys-273 and Cys-283 Are Located Close to the NADH-binding Site but Are Not Catalytically Essential. *J Biol Chem* 1991, 266:7531-7536.
  30. RODGERS KK, SLIGAR SG: Mapping Electrostatic Interactions in Macromolecular Associations. *J Mol Biol* 1991, 221:1453-1460.
- This paper describes how the relative contributions of complementary electrostatic and hydrophobic interactions can be extracted from a variety of physical measurements on reversibly binding electron-transfer proteins.
31. MIYASAKA T, KOYAMA K, ITOH I: Quantum Conversion and Image Detection by a Bacteriorhodopsin-based Artificial Photoreceptor. *Science* 1992, 255:342-243.
  32. TAKEI H, LEWIS A, CHEN Z, NEBENZAHL: Implementing Receptive Fields with Excitatory and Inhibitory Optoelectrical Responses of Bacteriorhodopsin Films. *Appl Optics* 1991, 30:500-509.

This paper draws an interesting parallel between bacteriorhodopsin film response and visual receptor fields, and illustrates the basic properties of cooperative, distributed information-processing networks.

33. ELLERBY LM, NISHIDA CR, NISHIDA F, YAMANAKA SA, DUNN B, VALENTINE JS, ZINK JI: Encapsulation of Proteins in Transparent Porous Silicate Glasses Prepared by the Sol-gel Method. *Science* 1992, 255:1113-1115.

The redox proteins copper-zinc superoxide dismutase, cytochrome *c* and myoglobin are immobilized in a glass matrix under gentle conditions that maintain the functional integrity of the protein. This matrix allows the diffusion of small molecules, making the composite material potentially useful for materials or biosensor applications.

34. WILLNER I, RUBIN S, ZOR T: Photoregulation of Alpha-chymotrypsin by Its Immobilization in a Photochromic Azobenzene Copolymer. *J Am Chem Soc* 1991, 113:4013-4014.
35. SUBRAMANIAM S, GREENHALGH DA, RATH P, ROTHSCHILD KJ, KHORANA HG: Replacement of Leucine-93 by Alanine or Threonine Slows Down the Decay of the N-Intermediate and O-Intermediate in the Photocycle of Bacteriorhodopsin—Implications for Proton Uptake and 13-*cis*-retinal → All-*trans*-Retinal Reversion. *Proc Natl Acad Sci USA* 1991, 88:6873-6877.
36. THORGEIRSSON TE, MILDER SJ, MIERCKE IJW, BETLACH MC, SHAND RF, STROUD RM, KLUGER DS: Effects of Asp-96 → Asn, Asp-85 → Asn, and Arg-82 → Gln Single-site Substitutions on the Photocycle of Bacteriorhodopsin. *Biochemistry* 1991, 30:9133-9142.
37. SUBRAMANIAM S, MARTI T, ROSSELET SJ, ROTHSCHILD KJ, KHORANA HG: The Reaction of Hydroxylamine with Bacteriorhodopsin Studied with Mutants that Have Altered Photocycles: Selective Reactivity of Different Photointermediates. *Proc Natl Acad Sci USA* 1991, 88:2583-2587.
38. MARTI T, OTTO H, MOGI T, ROSSELET SJ, HEYN MP, KHORANA HG: Bacteriorhodopsin Mutants Containing Single Substitutions of Serine or Threonine Residues Are All Active in Proton Translocation. *J Biol Chem* 1991, 266:6919-6927.
39. HOLZ M, DRACHEV IA, MOGI T, OTTO H, KAULEN AD, HEYN MP, SKULACHEV VP, KHORANA HG: Replacement of Aspartic Acid-96 by Asparagine in Bacteriorhodopsin Slows Both the Decay of the M Intermediate and the Associated Proton Movement. *Proc Natl Acad Sci USA* 1989, 86:2167-2171.
40. HAMPP N, BRAUCHLE C, OESTERHELT D: Bacteriorhodopsin Wildtype and Variant Aspartate-96 to Asparagine as Reversible Holographic Media. *Biophys J* 1990, 58:83-93.
41. HENDERSON R, BALDWIN JM, CESKA TA, ZEMLIN F, BECKMANN E, DOWNING KH: Model for the Structure of Bacteriorhodopsin Based on High-resolution Electron Cryo-microscopy. *J Mol Biol* 1990, 213:899-929.

A breakthrough paper describing the bacteriorhodopsin three-dimensional structure at the level of detail required for systematic protein engineering.

42. NONELLA M, WINDEMUTH A, SHULTEN K: Structure of Bacteriorhodopsin and *in situ* Isomerization of Retinal: a Molecular Dynamics Study. *Photochem Photobiol* 1991, 54:937-948.

A state-of-the-art study of dynamical processes that are key to bacteriorhodopsin function.

43. BAYLEY H: Monolayers from Genetically Engineered Protein Pores. *Mater Res Soc Symp Proc* 1991, 218:69-74.
44. PICOREL R, HOLT RE, HEALD R, COTTON TM, SEIBERT M: Stability of Isolated Bacterial and Photosystem-II Reaction Center Complexes on Ag Electrode Surfaces—a Surface-enhanced Resonance Raman Study. *J Am Chem Soc* 1991, 113:2839-2843.
45. SCHUHMAN W, OHARA TJ, SCHMIDT HL, HELLER A: Electron Transfer Between Glucose Oxidase and Electrodes via Redox Mediators Bound with Flexible Chains to the Enzyme Surface. *J Am Chem Soc* 1991, 113:1394-1397.

46. FINKLEA HO, HANSEW DD: Electron Transfer Kinetics in Organized Thiol Monolayers with Attached Pentammine(pyridine)ruthenium Redox Centers. *J Am Chem Soc* 1992, 114:3173-3182.

47. TARLOV MJ, BOWDEN EF: Electron-transfer Reaction of Cytochrome-c Adsorbed on Carboxylic Acid Terminated Alkanethiol Monolayer Electrodes. *J Am Chem Soc* 1991, 113:1847-1849.

48. WILLNER I, LAPIDOT N: Electrically Wired Glutathione Reductase—a Biocatalyst for the Photochemical Reduction of Glutathione. *J Am Chem Soc* 1991, 113:3625-3626.

49. YOMO T, URABE I, OKADA H: Preparation and Kinetic Properties of 5-Ethylphenazine Polyethylene-glycol Glutamate-dehydrogenase Conjugate—a Semisynthetic NADH Oxidase. *Eur J Biochem* 1991, 196:343-348.

50. PERSSON M, MAANSSON MO, BÜLOW L, MOSBACH K: Continuous Regeneration of NAD(H) Covalently Bound to an Engineered Cysteine Residue of Glucose Dehydrogenase. *Biotechnology* 1991, 9:280-285.

51. SEEMAN NC: DNA Structural Engineering Using Immobile Junctions. *Curr Opin Struct Biol* 1991, 1:653-661.

DNA lattices offer extraordinary potential for two- and three-dimensional scaffolding for MED devices, particularly when they additionally incorporate internal or terminal biotinylation sites. These sites can interconnect DNA lattices with avidin or streptavidin molecules to produce additional symmetric, multivalent biotin attachment sites.

52. DARST SA, AHLERS M, MELLER PH, KUBALEK EW, BLAKENBURG R, RIBI HO, RINGSDORF H, KORNBERG RD: Two-dimensional Crystals of Streptavidin on Biotinylated Lipid Layers and Their Interactions with Biotinylated Macromolecules. *Biophys J* 1991, 59:387-396.

Streptavidin monolayers are demonstrated to have extraordinary surface order which, together with their ability to irreversibly bind virtually anything that can be conjugated to biotin, suggest a central role for the system in MED-device fabrication.

53. EIGLER DM, SCHWEIZER EK: Positioning Single Atoms with a Scanning Tunnelling Microscope. *Nature* 1990, 344:524-526.

An initial demonstration of the possibility of patterning surfaces by manipulation of single atoms.

54. PICKOVER C: Mathematics and Beauty VIII: Tessellation Automata Derived from a Single Defect. *Comp Math Appl* 1989, 17:321-336.

A remarkable paper illustrating the temporal evolution of complex interconnected lattices bearing a striking resemblance to integrated circuits.

55. SASABE H (ED): *Proceedings of the Frontier Research Forum on Bioelectronic Materials*. Wako-Shi: RIKEN Institute; 1991. Creative efforts to capitalize on the use of proteins as biomolecular devices has come from extensive work by the Frontier Research Program at the RIKEN Institute in Japan under the leadership of Hiroyuki Sasabe. This has included ordered assembly of such photoactive systems as nitrile hydratase, photosynthetic reaction centers and bacteriorhodopsin, as well as non-linear optical properties of polymer protein systems.

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