Simple mechanisms for the evolution of protein complexity

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1 | GRADUALISM AND PROTEIN COMPLEXITY

To understand how living things acquired their complex features—structures and functions that arise from specific interactions among differentiated parts—has been a central aim of biology for centuries.¹ Darwin supplanted divine agency with the evolutionary view: complexity arises through “numerous successive, slight modifications” under the influence of natural selection, because each step enhances functions that contribute to fitness.²-⁴

This scenario of gradual elaboration and optimization is well-supported in numerous cases.²-⁴ The most famous is the modern vertebrate eye, which evolved from a simple light-sensitive precursor by sequentially adding cell types and more complicated relationships among tissues, each of which improved visual sensitivity or acuity.⁵,⁶

In the last half-century or so, a pageant of intricate forms has been revealed at a tiny new scale. Every protein is itself a complex system, because its physical and functional features depend on a large number of interactions among its many constituent amino acids. For example, a protein’s ability to fold into its native tertiary structure depends on complementary steric, electrostatic, and hydrophobic interactions among scores or hundreds of residues.⁷ The same is true of quaternary structure:
most proteins assemble with other molecules into specific multimeric complexes, and the interfaces that hold these complexes together often involve dozens of tightly packed residues with a high degree of electrostatic and steric complementarity.\textsuperscript{11} Another form of complexity is allostery—changes in a protein’s function caused by binding an effector molecule—which typically involves many amino acids to bind the effector and coupling of binding to the active site.\textsuperscript{12,13}

During the last $\sim$3.8 billion years, evolution has generated proteins with thousands of different folds,\textsuperscript{10} unique multimeric interactions,\textsuperscript{11,14,15} and varying modes of allosteric regulation.\textsuperscript{11,16} This diversity presents a molecular version of the classic question about the evolution of biological complexity: how did the stepwise processes of evolution repeatedly produce complicated systems from simpler precursors? Darwin’s model of gradual adaptive elaboration was developed to explain morphological and physiological complexity, but it has been assumed to apply as well to the evolution of complex molecular features.\textsuperscript{2,3,17,18} Intuitively, the view that protein complexity always evolves by long, consistently adaptive trajectories may seem sensible or even necessary, given certain assumptions. For a feature to evolve, the sequence states that encode it must arise by mutation and then be fixed in populations. Multimerization, allostery, and protein folds all involve elaborate arrays of interacting amino acids, so how else could they have been acquired if not by a long and specific series of many sequence changes? And, in turn, how could we explain the fixation of a long series of particular mutations if each step were not driven by the deterministic power of selection? It has been suggested that such features might arise neutrally,\textsuperscript{19–21} but fixation by chance alone would be vanishingly improbable if many particular mutations are required.

Recent advances in protein biochemistry and molecular evolution call into question the assumptions that underlie the argument for the gradual adaptive evolution of protein complexity. Of particular note are dramatic improvements in protein design,\textsuperscript{22–24} deep mutational scanning\textsuperscript{25–27} (which characterizes the functions of huge numbers of protein sequence variants), and ancestral protein reconstruction\textsuperscript{28,29} (which uses phylogenetics to infer the sequences of ancient proteins and experiments to determine the molecular functions and structures that existed in the deep past). This new body of work shows that just one or a few mutations can drive the acquisition of multimerization, allostery, and even new folds from natural precursors that lack these features; furthermore, it also explains why these short paths exist: simpler proteins often already possess most of the physical properties that underly these features. Moreover, the networks of sequences that yield multimerization, allostery, or a given protein fold appear to be immense, and they are closely intercalated at numerous places with the sequence networks of functional proteins that lack the feature. As a result, proteins can—and do—acquire new complex features by neutral processes. Contrary to the metaphor underlying the gradualist view, the complex features of proteins are not singular, massive mountain peaks that an evolving protein can climb only via a long trek under the deterministic engine of natural selection. Rather, many complex features are better conceived of as innumerable wrinkles, each small enough to be mounted in a single step (or just a few), which proteins repeatedly encounter as they wander through a vast multidimensional landscape of functional amino acid sequences.

2 | SHORT PATHS TO MULTIMERIZATION

A substantial body of work shows that introducing one or a few mutations into naturally occurring proteins can confer on them the capacity to form new higher-level complexes. A selection of relevant studies are listed in Table 1, and we discuss a few highlights below.

2.1 | Engineering new multimeric interactions

For years, protein engineers have been reworking the surfaces of natural proteins to yield new molecular complexes and interactions via just a few mutations.\textsuperscript{23,31–33,35} A 2008 study, for example, conferred new homomeric interactions on five different proteins by introducing between one and four mutations, which replaced polar residues on the surface with hydrophobic amino acids and improved the steric fit between the target surfaces. These mutations conferred dimerization on a monomer, tetramerization on a dimer, and octamerization on a tetramer.\textsuperscript{23} (Figure 1a).

A recent study found that large multimeric assemblies are almost shockingly easy to evolve by mutation and provided a biophysical explanation for this behavior.\textsuperscript{24} The authors were motivated by the observation that in sickle cell disease, a single glutamate-to-valine mutation on the surface of human hemoglobin-$\beta$ (Hb-$\beta$) increases the affinity of hemoglobin molecules for each other; hemoglobin is itself a tetramer that contains two Hb-$\beta$ and two Hb-$\alpha$ proteins, so this mutation appears twice in each hemoglobin complex and triggers assembly into massive disease-causing fibers.\textsuperscript{40} Inspired by this example, this study took as starting points 12 unrelated
TABLE 1  Protein engineering and evolutionary studies in which occupancy of novel oligomeric states is conferred by one or a few mutations

| Protein system                  | Initial state | Derived state | Mutations | Isologous? | Notes on mechanism                                                                 | Type of experiment          | Reference          |
|---------------------------------|---------------|---------------|-----------|------------|-------------------------------------------------------------------------------------|-----------------------------|--------------------|
| 6-phospho-Beta-galactosidase     | ○             | ○             | 4         | Yes        | Polar to hydrophobic mutations at interface                                         | Rational mutagenesis        | Grueninger et al.  |
| Urocanase                       | ○○            | ○○            | 3         | Yes        | Polar to hydrophobic mutations pack against existing hydrophobic residues. Each mutation repeated four times in the complex due to isology. | Rational mutagenesis        | Grueninger et al.  |
| 1-rhamnulose-1 phosphate        | ○○            | ○○            | 1         | Yes        | Polar to phenylalanine mutation fits into an existing hydrophobic pocket. Mutation repeated eight times in the octamer; | Rational mutagenesis        | Grueninger et al.  |
| aldolase                        | ○○            | ○○            |           |            |                                                                                      |                             |                    |
| Ancestral Hemoglobin precursor  | ○○            | ○○            | 2         | Yes        | Derived tryptophan fits into a hydrophobic pocket on ancestral surface.             | Ancestral sequence reconstruction | Pillai et al.     |
| (Anc_α/β)                       |               |               |           |            |                                                                                      |                             |                    |
| B. subtilis TRAP complex        | ○○            | ○○            | 1 (deletion of 5 residues) | No | Change in angle between neighboring subunits makes space for an additional subunit in a ring complex. | Rational mutagenesis        | Chen et al.        |
| Streptococcal Gb1               | ○             | ○             | 1         | Yes        | Phenylyalanine mutation packs against unchanged residues                             | Mutational scan             | Jee et al.         |
| Streptococcal Gb1               | ○             | ○             | 5         | Yes        | Mutations in the protein core and surface result in tetramer with interdigitated subunits. | Mutational scan             | Jee et al.         |
| IgG-binding domain of protein L | ○             | ○             | 1-3       | Yes        | Mutations stabilize a domain-swapped dimer. A single mutation produces a micromolar strength dimer. An additional two mutations yielded picomolar affinity. | Rational mutagenesis        | Kuhlman et al.     |
| PLCδ1-PH + EPOR                 | ○○            | ○○            | 2         | No         | Derived Phe in one subunit foms hydrophobic cluster with two Phes on the other;     | Rational mutagenesis        | Liu et al.         |
| Dlg-PID + Pins                  | ○○            | ○○            | 1         | No         | Mutation stabilizes open conformation, facilitating interaction with protein partner | Ancestral sequence reconstruction | Anderson et al.    |
| Isoaspartyl dipeptidase          | ○○            |               | 1         | Yes        | Derived tyrosine mediates eight isologous contacts between octamers in a fibril      | Rational mutagenesis        | Garcia-Seisdedos et al. |
| L-Fucose mutarotase             | ○○            |               | 1         | Yes        | Derived tyrosine mediates eight isologous contacts between decamers in a fibril     | Rational mutagenesis        | Garcia-Seisdedos et al. |
| AsnC                            | ○○            |               | 2         | Yes        | 2 derived tyrosines mediate isologous contacts with another octamer                 | Rational mutagenesis        | Garcia-Seisdedos et al. |

Note: The column labeled mutations indicates the number of amino acid point mutations required, unless otherwise noted.
proteins that form small soluble homomeric complexes, then introduced a few surface mutations that increase hydrophobicity (Figure 1b). They changed no more than three sites per protein, used mutations only to leucine or tyrosine, and made no attempt to design for steric or electrostatic complementarity. In all 12 proteins, the mutations triggered assembly into long fibrils, and in four cases, a single mutation was sufficient. The affinities were high enough that massive assemblies formed at physiological concentrations—in some cases in the
nanomolar range—and all the proteins remained folded in their native structures, indicating that a specific new interface, rather than unfolding and aggregation, mediated assembly. Altogether, the authors tested 73 mutants, of which 30 triggered formation of fibrous complexes. They concluded that most small multimers exist in evolutionary terms “on the edge of supramolecular assembly.”

Other studies make clear that rational design is not necessary to find short genetic paths to the acquisition of multimerization. A small library of just ~50 random mutants of the monomeric protein GB1 was found to contain a single mutation that yields a dimer, a quartet of substitutions that produced another structurally distinct domain-swapped dimer, and a quintet that produced an intertwined tetramer. And a directed evolution study using the monomeric protein αE7 carboxylesterase found that multimeric interactions can be acquired even without direct selection: just four rounds of mutation/shuffling and selection for increased thermal stability yielded a variant that assembled into a mixture of monomers, dimers, and tetramers. This new protein differed from its ancestor by seven mutations, but individual reversion experiments showed that two of these had no effect on multimerization, indicating that it takes at most five to confer multimerization.

2.2 Historical evolution of new multimers

Evolutionary case studies show that large-effect mutations have also played a causal role in the historical origin of biologically important protein complexes. One study focused on the origin of hemoglobin, the primary oxygen transport and exchange protein in jawed vertebrates. Hemoglobin’s paralogous subunits Hb-α and Hb-β are part of a larger family of globins, and those closely related to vertebrate Hbs are mostly monomeric. The authors used ancestral sequence reconstruction to infer the sequences of ancient hemoglobin progenitors and biochemical experiments to characterize their ability to multimerize. This work established that tetramerization evolved from an ancestral dimer immediately after the gene duplication that yielded separate Hb-α and Hb-β proteins. When just two historical substitutions that occurred at the tetramerization surface of Hb-β during this interval were introduced into the ancestral dimer, they conferred high-affinity assembly into tetramers (Figure 1c). As in the biochemical study of supramolecular assembly, one of these replaced a glutamine with a large hydrophobic tryptophan, which fit into a pre-existing hydrophobic cavity on the facing subunit, and this interaction occurs twice in the interfaces by which two heterodimers assemble into the hemoglobin tetramer.

Another ancestral reconstruction study addressed how a monomeric enzyme evolved to form a novel heteromeric complex that plays a key role in the organization of cells within multicellular animals. The protein–protein interaction domain (PID) of the Discs Large (Dlg) protein serves as a scaffolding molecule that orients the mitotic spindle relative to cues at the cell surface in metazoans. PID descends from a much older family of monomeric enzymes, the guanylate kinases (GKenz), which produce GTP by phosphorylating ATP and do not interact with any of Dlg-PID’s protein partners. Ancestral reconstruction experiments showed that the PID’s ability to interact with one of its protein partners was acquired just after duplication of the common the gene duplication ancestor of GKenz and Dlg-PID, which had robust
GTPase activity and no detectable protein-binding affinity. Introducing either one of two substitutions that occurred during this interval into the monomeric ancestral enzyme was found to confer micromolar protein-binding affinity. In this case, the mutations were in a hinge region and appear to increase the accessibility of the protein binding surface rather than changing its surface properties.

### 2.3 Biophysical causes of easy interface evolution

What mechanisms explain how entirely new, high-affinity interactions can be conferred on monomeric proteins by just one or a few mutations? Two fundamental features of protein biochemistry help explain this apparently counterintuitive result.

First, the scale of the nonlinear relationship between the occupancy of a state and its free energy mean that one or two favorable mutations can increase a protein’s propensity to multimerize by orders of magnitude. It has been argued that mutations that create a new hydrogen bond across an interface will contribute only weakly to the energy of binding—typically <1 kcal/mol—because the hydrogen bond also forms in the monomeric state, via interactions with water. But other kinds of mutations have much larger effects. A single mutation that satisfies or removes an unpaired hydrogen bond donor or acceptor in a buried interface can strengthen the interaction by up to 4 kcal/mol (16 kJ/mol). And a mutation that introduces a hydrophobic group that is complementary to the opposing interface can strengthen the interaction by a similar amount.

Mutations with energetic effects of this magnitude can have large effects on multimerization, because occupancy is an exponential function of energy* (Figure 2a). For a protein with an initially low propensity to dimerize (e.g., 1%) a mutation that improves the energy of binding by 4 kcal/mol will increase the occupancy of the dimeric state by about two orders of magnitude (to 75%, orange rectangle). Two mutations, each contributing 4 kcal/mol, could increase occupancy of the dimer by four orders of magnitude (e.g., from 0.01 to 90%).

The second critical factor is that most homomeric interfaces are isologous: they involve the same surface region on the two subunits, rotated 180° relative to each other around an axis of symmetry (Figure 2b). A single mutation in an isologous interface will change residues on both sides of the interface, doubling its energetic contribution and squaring the effect on occupancy. One favorable mutation can thus have the same effect as two mutations in a nonisologous interface, improving binding by up to 8 kcal/mol and occupancy by four orders of magnitude. For higher-order homomeric complexes, the effect is magnified further: a single affinity-increasing an

![Figure 2](image_url)

**Figure 2** Isology and the evolution of multimerization. (a) Non-linear relationship between the free energy of assembly into a dimer, and fraction of monomers associated into the dimeric state is shown. Boxes indicate the effect on occupancy of a mutation that modifies energy by 4 kcal/mol (orange) and 8 kcal/mol (yellow). (b) Repetition of interactions in an isologous interface. In a dihedral dimer, a single favorable mutation (red) occurs twice—once on each subunit. In a tetramer (orange), it occurs four times. In a fibril with asymmetric interactions (right), a mutation on each subunit (black stick) is repeated without limit.
isologous interface occurs four times in a homotetramer (increasing occupancy by up to 8 orders of magnitude), eight times in a homo-octamer, and a vast number of times in a fibril.\textsuperscript{23,24,30}

\subsection*{2.4 Fortuitous foundations for new multimers}

Another factor that makes it easy to acquire new multimers is that monomers fortuitously contain surface features that can contribute to the affinity of a new interaction. For example, replacing one polar surface residue with phenylalanine transforms the tetramer L-rhamnulose-1 phosphate aldolase (Rua) into an octamer, because the side-chain of Phe packs into a small hydrophobic cleft formed by several existing residues on the facing surface (Figure 1a). Similarly, the historical tryptophan substitution that played a key role in the evolution of hemoglobin tetramerization fits into a small cavity composed of hydrophobic residues that were already present in the ancestor. In both Rua and hemoglobin, these new interactions were supplemented by other favorable interactions—hydrogen bonds and hydrophobic packing—among other existing residues in the interface. On their own, these pre-existing interactions were not sufficient to drive high occupancy of the multimer, but they created a context in which the even more favorable mutation created by the focal mutation could do so.

Most monomers contain large hydrophobic patches on their surfaces that can provide a foundation for new multimeric interactions.\textsuperscript{52-54} One study of a large number of protein surfaces estimated that the average 1,000 Å\textsuperscript{2} exposed patch is just two substitutions away from being as hydrophobic as the average protein–protein interface; many such patches are just one substitution away from this level of hydrophobicity, and some are already there (Figure 1e). Because most of the affinity in multimeric interactions comes from hydrophobic interactions, these patches have the potential to mediate fortuitous new interactions, so long as electrostatic or steric clashes do not prevent them from doing so. A mutation that resolves a clash between subunits or adds a new hydrophobic residue could therefore yield a high-affinity interaction mediated by these pre-existing hydrophobic surfaces.

The surface properties that poise proteins on the edge of multimerization appear to occur fortuitously. One line of evidence comes from the finding that proteins that do not co-occur in nature often co-assemble when mixed in the laboratory. For example, when human proteins are expressed in \textit{Escherichia coli} cells and then characterized by in-cell NMR, they are more likely to form complexes with the host proteins than the \textit{E. coli} proteins do with themselves.\textsuperscript{58} The two sets of proteins have not encountered each other for billions of years, so these interactions must be entirely fortuitous. A similar lesson comes from X-ray crystallography, which is possible because many proteins fortuitously form repeating homomultimeric structures: although particular conditions must be imposed for these crystal interactions to be realized, the interfaces involved often resemble those of biological multimers in their size and complementarity.\textsuperscript{24,39,55–57} (Figure 1e). These crystallographic interactions do not occur biologically, so they could not have arisen because of selection.

\section{SHORT PATHS TO ALLOSTERY}

Allostery—changes in a protein’s activity, such as ligand-binding or catalysis, caused by binding to an effector molecule\textsuperscript{51)—arises if three necessary and sufficient conditions are present (Figure 3a): (a) a protein must bind an effector at a site that is structurally distinct from the active site; (b) the protein must be capable of occupying conformations that differ in their functional activity; and (c) effector binding must be associated with a difference in the relative stability—and therefore occupancy—of active versus inactive conformations. One might expect that the evolution of allostery from a nonallosteric precursor would require a protein to acquire each of these properties, and—because each property involves many residues in the protein—that doing so would require many mutations. In fact, recent biochemical and evolutionary studies show that allostery can be acquired via just a few genetic changes, because many natural nonallosteric proteins often already possess several of the necessary conditions for allostery. These studies are listed in Table 2, and we discuss a few highlights here.

\subsection*{3.1 Engineering allostery}

Protein engineering studies of four unrelated proteins with dramatically different folds and functions—green fluorescent protein, β-glycosidase, β-glucoronidase, and YeaZ—have shown that a single mutation can confer allostery by causing the protein to both bind an effector and become allosterically dependent on it (Figure 3b, c). In every case, the same kind of mutation was involved: replacing a tryptophan with glycine >9 Å from the active site. The mutation creates a small cavity, which destabilizes the active conformation and dramatically reduces the protein’s activity; however, the small molecule indole, which structurally resembles the side chain
of tryptophan, binds in the cavity, restabilizes the active conformation, and restores activity. There was no need to create a pathway to “transmit information” from the effector site to the active site; the active site was already sensitive to the conformation and stability at the site of the mutation.

The bacterial enzyme TEM-1 beta lactamase is nonallosteric, but mutations that create a new site for effector binding are sufficient to confer allostery (Figure 3d,e). In TEM-1’s active conformation, two structural regions of the protein separated by a flexible hinge are oriented closely together, and residues from both regions comprise the active site. Introducing just two histidine mutations into the hinge of a nonallosteric TEM-1 variant creates a binding site for metal ions, and metal binding allosterically downregulates catalytic activity by reducing occupancy of the active conformation. TEM-1 can also be mutated to acquire positive allosteric regulation: inserting a maltose-binding protein (MBP) domain interferes with the active conformation, but binding maltose...

**FIGURE 3** Legend on next page.
relieves this inhibition and restores activity.\textsuperscript{60,64} Inserting an effector-binding domain confers allosterity on other proteins, too, including alkaline phosphatase\textsuperscript{70} and dihydrofolate reductase.\textsuperscript{126} In all these cases, creating a new effector binding site by fusion or point mutation is sufficient to confer allosterity because the wild-type protein already possesses two of the three properties critical for allosterity—the capacity to occupy multiple conformations that differ in their activity, and sensitivity of those conformations to the state at a different location in the structure.

Sometimes nonallosteric proteins can already bind a ligand far from the active site, but this has no allosteric effect on the protein’s function; in these cases, a mutation that changes the relative stability of the liganded state versus the unbound state can confer allosterity. For example, in five unrelated enzymes that are noncooperative multimers with isologous interfaces, a single amino acid replacement or deletion generates strong cooperative activity.\textsuperscript{61,62,65,66,72} (Cooperativity is a form of allosterity in multimeric proteins, in which activity in one subunit increases the activity of another subunit; the ligand at the active site of one subunit serves as an effector for the other.) Homomers with isologous interfaces are particularly likely to evolve cooperativity.\textsuperscript{63} because if the active site for one subunit is conformationally coupled to the multimerization interface, then the other must be identically coupled. In some of the enzymes studied, the causal mutation was at the interface between subunits, while in others it was in the active site (Figure 3f,g). The precise structural mechanisms were not directly identified, but an interface mutation could confer cooperativity by simply changing the multimerization affinity of a liganded subunit for an unliganded subunit, relative to the affinity when both are liganded (or both unliganded). An active-site mutation could confer cooperativity by tuning the relative stability of the active conformation when the other subunit is liganded versus when it is unliganded.

There are even cases in which all three requirements for allosterity are already present in nonallosteric proteins: no effector exists in nature, but these proteins can be regulated by exogenous allosteric drugs or other effectors.\textsuperscript{73,74} For example, aminoglycoside phosphotransferase (3’)-IIIa (APH) is a bacterial kinase with no known natural allosteric effectors. One study created a large library of potential effectors by randomizing the residues along one surface of an ankyrin repeat protein (ARP), a stable
protein with a well-defined binding surface. Screening the library identified hundreds of ARPs that negatively regulate the wild-type enzyme, one of which bound with nanomolar binding affinity that completely abolished catalytic activity. An X-ray crystal structure showed that this effector bound the enzyme at a surface site >20 Å from the active site and stabilized several secondary elements of the protein in an inactive conformation.69 The

| Protein system | Initial state | Derived state | Mutations | Notes on mechanism |
|---------------|---------------|---------------|-----------|--------------------|
| cpTEM1 beta – lactamase | ![Initial State](image1) | ![Derived State](image2) | 2 | Construction of metal binding pocket stabilizes the inactive conformation of the enzyme. Mathieu et al.69 |
| Ancestral hemoglobin precursor (Ancα/β) | ![Initial State](image3) | ![Derived State](image4) | 2 | Evolution of a tetramerization interface in an ancestral dimer with high O2 affinity (R) stabilizes a lower-affinity conformation and confers cooperativity. Pillai et al.20 |
| TEM1-beta-lactamase + MBP | ![Initial State](image5) | ![Derived State](image6) | Insertion | Recombination of the two proteins yields maltose-sensitive TEM-beta lactamase activity. Guntas et al.64 |
| KSS1 | ![Initial State](image7) | ![Derived State](image8) | 2-4 | Design of novel phosphosites allows for inhibition or activation of KSS1 activity. Pincus et al.156 |
| Beta-glycosidase Beta-Glucoronidase | ![Initial State](image9) | ![Derived State](image10) | 1 | Chemical rescue of a W>G mutant by addition of indole, allows for positive regulation by indole. Deckert et al.37 |
| YeaZ | ![Initial State](image11) | ![Derived State](image12) | 1 | Chemical rescue of dimerization in a W>G mutant by addition of indole, allows for positive regulation by indole. Xia et al.58 |
| Pyruvate kinase M1 | ![Initial State](image13) | ![Derived State](image14) | One-residue deletion | Removal of a single C-terminal residue allows for FBP driven inhibition in a non-allosteric isoform of pyruvate kinase. Ikeda et al.65 |
| Ornithine transcarbamoylase | ![Initial State](image15) | ![Derived State](image16) | 1 | Substitution at the intersubunit interface allows for cooperativity. Kuo et al.66 |
| Glutathione reductase | ![Initial State](image17) | ![Derived State](image18) | 1 | Substitution at the intersubunit interface allows for cooperativity. Scrutton et al.62 |
| Aspartate transcarbamoylase | ![Initial State](image19) | ![Derived State](image20) | 1 | Substitution at the active site allows for cooperativity. Stebbins et al.61 |
| CEBPB | ![Initial State](image21) | ![Derived State](image22) | 3 | Historical substitutions abolish ancestral positive regulation and generate novel negative phospho regulation. Lynch et al.67 |
| Tryptophan synthase | ![Initial State](image23) | ![Derived State](image24) | 4 | Historical substitutions switch ancestral negative regulation to positive regulation. Schupfner et al.58 |
| Aminoglycoside phosphotransferase | ![Initial State](image25) | ![Derived State](image26) | 0 | Binding ankyrin repeat proteins regulates the wild-type nonallosteric enzyme. Kohl et al.69 |

Note: The column labeled mutations indicates the number of amino acid point mutations required, unless otherwise noted. Conformational ensembles associated with the initial (nonallosteric) and derived (allosteric) states are shown in schematic form.

Abbreviations: A, active conformations; I, inactive.
number of mutations required to confer allostery on the APH protein is therefore zero.

3.2 | Natural evolution of allostery

Just as allostery can be engineered in the lab with a few mutations, several historical case studies show that allostery has been acquired during natural evolution by very short paths (Table 2).

The hemoglobin tetramer, for example, binds oxygen cooperatively. Ancestral protein reconstruction showed that hemoglobin acquired cooperativity during the same phylogenetic interval when the tetrameric architecture evolved from an ancestral, noncooperative dimer. The two historical surface substitutions that confer tetramerization on the ancestral dimer—which was not cooperative—also confer cooperativity. The structure of hemoglobin suggests why conferring tetramerization also yields this form of allostery: the surface that mediates tetramer assembly is connected by a short helix to the protein’s active site, which subtly changes conformation when oxygen is bound, altering the shape of the binding interface. The association between the active site and this part of the surface simply arises from the globin fold; even myoglobin, a monomer that is necessarily noncooperative, undergoes a similar conformational change when it binds oxygen.75,76 Allostery arises immediately when binding to an effector is conferred via the new tetramerization interface, simply because the conformation that is optimal for oxygen binding is not optimal for tetramerization, and vice versa (Figure 3h).

A few historical studies have also documented allosteric inversions, in which positive allostery has been gained from a negatively allosteric ancestor (or vice versa), by just a few mutations.67,68 For example, ancestral reconstruction studies show that the ancestral transcription factor CEBPB was negatively regulated by phosphorylation, but allosteric activation by phosphorylation was gained on the branch leading to eutherian mammals.67 Just three historical amino acid replacements are sufficient to recapitulate this inversion. Two of these abolish phosphorylatable serines that favored the inactive conformation when phosphorylated, while the third introduces a new serine that makes the protein’s activity phosphodependent. By tuning the relative stabilities of the protein’s conformations in response to phosphorylation, these mutations abolished an ancient form of regulation and—with a single new mutation—conferred a new one.

Finally, just as some nonallosteric proteins can be regulated by allosteric drugs, the yeast protein Fus3 apparently had all the properties required for an allosteric response, before its effector evolved.77 Fus3’s activity in the mating cascade in S. cerevisiae is allosterically upregulated by the VWA domain of a protein called Ste5. Ste5 is a relatively recent evolutionary newcomer, but Fus3 proteins from a wide variety of fungal species—even those that contain no Ste5 family members—can be regulated by heterologous copies of S. cerevisiae Ste5. Phylogenetic analysis shows that Fus3’s capacity to be allosterically regulated by Ste5 existed in latent form long before Ste5 itself evolved. This suggests that when Ste5 first evolved, it bound immediately to Fus3—at a site where no other regulator is known to bind—and fortuitously upregulated its activity. As in the case of allosteric drugs, zero mutations in the protein itself were required to confer allostery on it.77

3.3 | Fortuitous foundations for acquiring allostery

Why would nonallosteric proteins possess features that are prerequisites for allostery—such as occupancy of conformations that are functionally distinct, or conformational linkages between a potential effector-binding site and the protein’s active site—if the protein is not already allosterically regulated? Like the precursors of multimerization, these features arise as fortuitous features of protein architecture.

Virtually all proteins meet the first condition for allostery: they occupy an ensemble of conformations, rather than a single rigid structure.78–83 Although physical constraints limit the number of conformations that a protein actually occupies, many degrees of freedom remain, so there are typically a vast number of subtly different conformations—differing in the angle of a helix or the rotamers of side chains, for example—with energies similar enough to allow nontrivial occupancy.84 Even proteins that almost exclusively occupy a single conformation have the capacity to occupy others, if the energy differences between the conformations are narrowed by one or a few mutations that destabilize the major conformation or stabilize others.79 Because of the exponential relationship between changes in the free energy of a fold and its occupancy, a single mutation with a small effect on stability can have a large effect on the protein’s ensemble of conformations (Figure 2a).

The second condition—that the ensemble of folded conformations includes some that are more active than others—is also a common feature of nonallosteric proteins.85,86 Many ligand-binding proteins exhibit conformational selection: they can occupy multiple conformations that vary in affinity, and the ligand binds predominantly to those with high affinity (Figure 3i).80,87,88 Similarly, many unbound enzymes transiently occupy both active
and inactive conformations, with preference for the active conformation when their substrate is bound. The functional heterogeneity of conformations arises simply because active sites have more stringent physical requirements than the fold as a whole. Affinity for a ligand can be impaired by subtle changes that disrupt the steric or electrostatic complementarity of binding surfaces, and catalytic activity involves even more precise constraints. Some of the many conformations that are occupied by a typical protein will inevitably satisfy these functional requirements better than others.

Finally, conformational changes in one region of a protein are often associated with changes elsewhere. Studies of the structural effects of mutations suggest that genetically perturbing nonallosteric proteins often changes the conformation at multiple parts of the same protein, often 10–20 Å away from the mutant sites. A recent deep mutational scan of PDZ and SH3 proteins found that a huge number of single mutations >5 Å away from the binding interface can change the proteins’ ligand affinity, often reducing but sometimes improving it. These proteins are not allosterically regulated, and the mutations do not occur in any known natural variants, so their effects cannot be attributable to selection. These couplings arise as a consequence of the constraints imposed by a protein’s structures. Although proteins have extensive conformational freedom, that freedom is not infinite. A subtle change in the position of a helix, for example, will alter the most favorable position of other residues and secondary elements that interact with it, and these changes may propagate further through the protein’s structure. Perturbing one part of a protein therefore typically changes the lowest-energy conformation at other locations. Sometimes the linked regions happen to be the active site and an effector-binding surface.

Nonallosteric proteins that have fortuitous properties like long-distance couplings or effector binding may therefore often be poised in sequence space on the evolutionary edge of allostery. A single mutation that adds the remaining requirement—such as effector binding or tuning the relative stabilities of active and inactive conformations—can trigger the acquisition of allosteric regulation.

4 | SHORT PATHS TO PROTEIN FOLDS

There are over 2,000 known protein folds in nature. Information about the mechanisms by which these folds originated—either by descent from other folds or de novo folding from unstructured precursors—is relatively scanty, because these events occur far more rarely than multimerization or allostery are acquired. Several case studies suggest, however, that simple mutations can create entirely new folds or make simpler folds suddenly more elaborate.

4.1 | Short bridges between folds

Three experimental studies show that the genetic paths between entirely different protein folds can, in some cases at least, be surprisingly short. In a classic biochemical study, it took only two point mutations to reorganize the secondary structure and tertiary fold of the N-terminal portion of the Arc repressor (Figure 4a). One of these mutations on its own resulted in partial occupancy of both folds, depending on experimental conditions. The bridge in sequence space between the folds is

![Figure 4](image-url)
therefore only two mutations long, and the journey across that bridge does require passing through an unstructured intermediate. The implication is that, even under purifying selection, one fold could rapidly evolve into the other.

A more recent study showed that a global structural reorganization of an entire globular protein can be triggered by a single mutation (Figure 4b). The A and B domains of protein G from *Streptococcus*, each ~50 amino acids long, are unrelated in sequence and structure: domain A consists of 3 α-helices, and domain B consists of 4 β strands and one helix, with a completely different tertiary fold and distinct sets of residues that make up the hydrophobic core. The authors gradually walked the two sequences towards each other in sequence space by swapping individual residues between the proteins without changing or losing the fold of either one. Ultimately, two protein sequences were achieved that differed by a single amino acid, each occupying one of the folds; the remaining mutation completely reorganized either fold into the other. Even though the “wild-type” A and B domains are as distant from each other in sequence space as is possible, a continuous network of sequences encodes each fold and, in at least this one location, the two networks are adjacent to each other.

The third example comes from two paralogous cysteine-rich domains in the cnidarian *Hydra*, which differ at 56% of their residues (Figure 4c). Unlike most homologous proteins with recognizably similar sequences, these two proteins have completely different folds, mediated by distinct patterns of disulfide bridges. Exchanging just two critical residues between the two proteins—which are not cysteines but rather favor different backbone conformations—fully switches one fold into the other. When the mutations are introduced singly, the intermediate proteins occupy both conformations. The sequence networks encoded by these folds are therefore connected by at least two short bridges, neither of which involves an unstructured intermediate. Because the proteins are paralogous, the authors commented that it is likely that one fold evolved from the other via a “smooth transition.”

Some natural proteins, too, can occupy two distinct folds, indicating that the sequence networks of the folds overlap. A recent bioinformatic study estimated that up to 4% of all proteins with solved structures have the capacity to occupy more than one distinct fold under different cellular conditions. For example, cyanobacterial protein KaiB can adopt two different tertiary structures, depending on whether it is bound or free of its binding partner KaiC. Similarly, the C-terminal domain of the transcription factor RfaH can transition between α-helical hairpin and a β-barrel fold in the cell, depending on whether it is free or bound to the protein’s NTD. In proteins like these, a single mutation that stabilizes or destabilizes one of the folds could trigger the transition to near-total occupancy of just one.

It is unknown how many protein folds in nature have originated by fold-switching mutations. Proteins with different folds are typically unalignable (though in rare cases homology between different folds can be detected), so it is impossible to trace the process of descent from potential common ancestors. But this does not necessarily mean such ancestors did not exist. Sequence evolution is constrained primarily by the protein’s structure: if a new protein family were born by transition into a new fold, it would immediately be subject to new constraints, and so the traces of their descent from a common ancestral fold would rapidly decay as the sequences diverge.

### 4.2 Elaboration of existing folds

Many proteins have complicated structures consisting of multiple layers of secondary elements packed against each other. It is clear that these complex structures can be easily acquired from simpler, more compact folds by elongation of the primary sequence and packing of the extended region against the exterior of the ancestral fold.

In the ligand-binding domain of steroid hormone receptors, for example, one group of paralogs acquired a new carboxy-terminal extension (CTE), a partially beta-containing secondary structure that creates a new layer of tertiary structure by packing against a mostly hydrophobic surface region that was exposed in the simpler ancestral structure. Similarly, a series of three nested insertion events within an ancestral loop of a beta lactamase generated an internal extension, containing both alpha and beta secondary structure, which creates a novel exterior wall by packing along one side of the ancestral fold.

Extensions and insertions are easy to acquire by mutation. C-terminal or N-terminal extensions can arise from point mutations that abolish stop or start codons, and mutations that move an intron splice site can insert new internal coding sequence. Sometimes these extensions and insertions will immediately contribute to the tertiary structure: the majority of random short peptides are predicted to have >25% secondary structure, and, as we have discussed in this review, many protein surfaces can fortuitously bind random peptides. It should be just as likely for a peptide to fortuitously bind the surface of a protein to which it is covalently attached as it is to bind the surface of a disconnected protein the two units are disconnected.
The second premise of the argument for adaptive graduality is that genotypes encoding complex features are rare. For the complex features of proteins, this assumption also turns out to be wrong. Comparative structural analyses and high-throughput mutagenesis experiments have shown that a vast number of protein sequences can encode essentially equivalent forms of multimerization, allostery, and tertiary folds. These genotypes are widely dispersed across vast connected regions of sequence space (Box 1). The bridges by which complexity can be acquired are not only short but also numerous.

5.1 | Degenerate multimerization interfaces

Deep mutational scanning experiments that the number of sequences that can encode a multimeric interaction between molecules is huge. A recent study used phage display to characterize binding by several S100 proteins to a library of random peptides 12 residues long (the same length as the region of the peptides that S100s endogenously bind). They found that each S100 protein bound 7–10% of the 30,000 peptides measured, with affinity comparable to S100’s biologically relevant binding partners. The vast majority of these peptides do not occur biologically, so these binding activities must be fortuitous rather than the result of selection.

Another study randomized 11 specificity-determining residues in the interface of the two-component signaling proteins PhoP and PhoQ and identified >500 unique functional pairs, most of which shared few or no common residues at these sites with each other or the wild-type pair. A second line of evidence comes from the record of long-term evolution in extant sequences. If interactions were subject to strict sequence constraints, the residues that mediate protein–protein interactions should be strictly constrained and therefore evolve slowly. In fact, many different amino acids are compatible with multimerization. For example, the acetyl-xylo-oligosaccharide esterase (Axe2) in thermophilic bacteria is an obligate

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**BOX 1  Multidimensional sequence space**

The concept of sequence space provides a useful organizing metaphor for protein sequence evolution. Sequence space consists of all possible sequences of a given length (the nodes), which are connected to each other by edges if they differ by a single mutation. The number of nodes in a protein space is vast: for an average protein 300 amino acids in length, there are \(20^{300} = 10^{390}\) possible sequences, far greater than the number of subatomic particles in the visible universe.

Sequence space is multidimensional and complex. Each site represents a dimension along which a protein can change independently by mut, and (leaving aside the genetic code), there are 19 possible mutations at each site. Any protein is therefore connected to \(19^{300} = 5,700\) unique neighbors via single amino acid replacements. Each of those has that many neighbors, too, so the starting protein is two steps away from about 18 million other proteins, three steps from 36 billion more, and four steps from 53 trillion others. Even the most distant pairs of proteins in this massive universe, which share no residues in common, are just 300 amino acid replacements apart.

The dense connectivity of sequence space has important consequences for evolutionary processes. Although the set of all possible protein sequences is vast, a protein has the potential to explore a huge number of possible alternative sequences via short evolutionary paths. The simplest and most common model for evolution across sequence space corresponds to drift under purifying selection and a rate of evolution slow enough that individual sequence changes are fixed sequentially rather than simultaneously. In this scenario, an evolving protein can move to any neighboring sequence, as long as the neighbor is functional. The set of functional sequences that are connected to each other by single mutations constitute a neutral network through which a protein may move over time. As long as functional proteins on average have more than one functional neighbor—which is clearly the case for real protein sequences—then neutral networks will be extensive, allowing substantial long-term divergence from the starting sequence. If the networks that encode proteins with different features or functions abut or overlap each other—as the studies that we review here establish—then a new property can evolve from an ancestral protein that lacks the feature, either by drift or positive selection, in as little as a single step.
octamer held together by several clusters of hydrogen bonds, but the states and sites used to mediate these bonds differ extensively among homologs from different species. More generally, across a large database of proteins, the rate of sequence evolution at residues buried in interfaces is only slightly slower than the mean rate over all residues, indicating that proteins can explore very large regions of sequence space while maintaining their multimeric interactions.

Not only can particular surfaces that mediate molecular interactions tolerate many different sequence states, but many different surfaces on proteins have the potential to mediate interactions. For example, members of the globin family have independently evolved to assemble into multimeric complexes multiple times in different animal lineages. On each occasion, largely nonoverlapping surfaces mediate the multimeric interactions (Figure 5a). Similarly, the homologous urease enzyme of a plant and a bacteria have independently evolved multimeric structures, but the subunits interact using different parts of their surfaces. And some proteins form multimers under crystallographic conditions that use different interfaces from those that they use to multimerize in solution.

These data show that proteins can encode a multimeric interaction using a wide variety of different amino states at the same set of sites, using different sites on the same surfaces, or use entirely different parts of the protein. A consequence of this degeneracy is to dramatically increase the number of mutational paths by which an interaction can be acquired.

### 5.2 Degeneracy of allostery

Allostery too can be encoded by a huge set of possible sequences, so nonallosteric proteins can acquire allostery by many possible mutations. Alignments of allosteric proteins indicate that the sequence networks encoding allostery are vast. For example, LacI and PurR are homologous transcription factors with similar mechanisms of effector-driven allostery, but they share <30% sequence identity.

Allostery can also be gained by a variety of mechanisms, increasing genetic degeneracy further. For example, multimeric globins in vertebrates, and mollusks (Figure 5a) independently acquired cooperative oxygen binding; these proteins differ from each other by up to 85%, and different surfaces and conformational mechanisms are involved in mediating the allosteric response to oxygen binding by other subunits.

Recent DMS studies of allostery provide a more comprehensive view of the sequence degeneracy of maintaining or gaining allostery. In a complete library of all single-site variants of the E. coli TetR protein (repressor of tetracycline resistance), 71% of all mutants maintained allostERIC regulation—far more than the number that lost allostery (while maintaining the ability to fold and function). Five non-allosteric mutants were then reintroduced and screened to identify variants that regained allosteric regulation: of 960 single-site mutants tested, 216 regained allostery. The allostery-restoring mutations were scattered throughout the protein's structure, and typically involved sites other than those that caused allostery to be lost (Figure 5b). Moreover, the sites that affect allostery in the experiments are highly variable in an alignment of TetR orthologs, indicating extensive degeneracy during the historical evolution of allostery.

Allostery can be maintained, acquired, or modified in so many different ways because active sites are fortuitously coupled to sites all across a protein's structure. A study in which a light-sensitive LOV2 domain was inserted at every surface site in DHFR found that 14 of the 61 insertions gained a significant, albeit modest, allosteric response to light. And the DMS scan of PDZ and SH3 discussed above found that ligand affinity was altered by mutations at about half of all surface sites away from the active site. This means that gaining effector binding at any one of a huge number of potential surface regions is likely to trigger an allosteric response. In a protein that already binds a potential effector, then a mutation at any third site coupled to the active site has the potential to affect the stability of active versus inactive conformations, again conferring allostery. As nonallosteric proteins evolve, they are therefore likely to encounter a lengthy menu of genetic options that, if any one of them are ordered up by mutation, can confer allostery upon them.

### 5.3 Degeneracy of folds

It is well known that a single protein fold can be encoded by a huge ensembles of variant sequences. Even the hydrophobic core of some proteins—the portion that typically evolves slowest—can be replaced with a set of entirely different hydrophobic amino acids while maintaining the native fold. Protein family members that can be structurally superimposed sometimes share no more than 15–20% sequence identity (Figure 5c). The network of sequences encoding a single fold therefore extends across almost the entire “diameter” of sequence space. Moreover, these highly divergent proteins descend from a common ancestor, indicating indicates that proteins across this span are almost certainly connected by paths on which all intermediate sequences can fold and function.
This extraordinary degeneracy means that proteins can explore vast sequence networks as they evolve under the constraints imposed by maintaining their ancestral fold. As they drift through this network, they may occasionally encounter boundaries of the networks that encode other folds, which are also vast. These bridges may be rare, but over time evolving proteins have an extraordinary number of opportunities to win the find-a-new-fold lottery without paying a price for their losing bet, because purifying selection removes mutations that cause proteins to unfold or aggregate. Moreover, gene duplication—and the functional redundancy it allows—can weaken the constraints imposed by purifying selection to maintain the ancestral function. Along with de novo origin of simple folds, evolutionary transitions from one fold to another need not have been frequent to
explain the origin of the few thousand known protein folds that exist during the course of four billion years of massively parallel evolution.

6 | EVOLUTIONARY PROCESSES AND PROTEIN COMPLEXITY

The evolution of any feature can be pictured in two steps: first, one or more mutations that produce the feature must arise in an individual, and those mutations must then become fixed in the population under the influence of selection and/or drift. The classical view of protein complexity entails interrelated assumptions about both parts of this process. With respect to mutation, the first assumption is that complex features require many mutations that gradually increase complexity from simpler ancestral forms; the second assis that there are few such sets. The final assumption pertains to fixation: complex features are assumed to be adaptive, which allows selection to drive the serial fixation of complexity-conferring mutations. The third assumption is necessary given the first two: if acquiring a complex features requires a particular set of many mutations, it would be vanishingly improbable for them all to fix by chance. If each mutation along the path were to enhance fitness, however, then selection would reliably drive the fixation of each one, given large population sizes and long enough timescales.

The problem with the classic argument is that its premises, rather than its logic, are wrong. We know now that multimerization, allostery, and new folds can be acquired from naturally occurring proteins via paths that are just one or a few mutations long, and the number of such possible paths is often very large. These findings open the door for non-classical explanations of how complexity arises and is fixed during evolution. When there are many complexity-conferring genotypes, and each requires just one or a few of many possible mutations, then persistent, long-term selection is not necessary to drive the smaller number of fixation events that are required. It becomes much more plausible that any one of the many possible paths to complexity would be followed under the influence of drift alone. Scenarios involving weak or transient selection, alone or in tandem with drift — linkages, also become plausible.

6.1 | Neutral acquisition of complex features

Neutral mutations that confer complex features would go to fixation by the same stochastic processes that cause other neutral sequence substitutions, a process that is well understood and essentially universal. Most mutations—whether neutral or beneficial—are lost by chance in the first few generations after they arise, because they occur in a single copy of a gene, and that copy is often unlucky in the lottery by which genes are transmitted to the next generation. Some mutations escape this fate and increase in frequency, and some — even neutral mutations—eventually reach fixation, simply because the frequency of an allele in a population fluctuates stochastically across generations. Any particular mutation is more likely to fix if it is beneficial than if it is neutral—and it takes less time for it to do so—but there are so many possible neutral mutations, and they arise so regularly, that a parade of neutral substitutions is constantly evolving over time in natural populations. Neutral sequence evolution takes place at a constant rate irrespective of population size, because any neutral mutation is more likely to fix in a small population, but more such mutations are generated in large populations, and the effect of population size therefore cancels out. In the end, the critical factor determining the rate of neutral substitution in a protein is the neutral mutation rate—the number of new neutral mutations in each copy of the gene coding for the protein in each generation. The resulting neutral sequence drift is the primary cause of diversity among protein sequences that have similar.

The dynamics of neutral sequence evolution can be extended to understand the evolution of neutral functional or biochemical features, like multimerization, allostery, or a new fold (see Ref. 135). If it takes only one neutral mutation to confer the feature, then the critical factor determining the rate at which the feature will evolve (and thus the probability that it will evolve in any particular period of time) is the overall mutation rate and the degeneracy of the feature—the number of possible amino acid replacements that have the potential to confer it. If instead it takes several mutations to confer the feature, the process will take longer, but the probability that such a path will be followed again increases with degeneracy—which is now the number of sets of mutations that can confer the feature. If the mutations are beneficial, this would speed the process, whether the selection pressure is strong and sustained—as in the classic model—or if it is weak or intermittent. Even paths involving weakly deleterious intermediate steps can be followed, but the plausibility of this scenario depends on the population size and recombination rate.
selection is always required to drive a long series of rare mutations. Complexity may sometimes be acquired neutrally via one or a few mutations; it may sometimes evolve by short paths driven by selection; and in other cases, it may involve long paths in which each step is adaptive, as envisioned by the classic model.

Supporting the idea that protein complexity sometimes evolves neutrally is the fact that many orthologous or paralogous multimers vary in these features but are no more efficient biochemically or functionally than their relatives. Experiments show that some multimers can be replaced by simpler versions without any detectable costs to fitness or function. Allostery, too, can apparently be acquired neutrally, as demonstrated by the existence of allosteric drugs, which can affect protein activity even though the protein’s intrinsic capacity for allostery has no biological function.

### 6.2 Entrenchment of complex features

A rejoinder to our view might be that complex features often persist for long periods of time, and we might expect that a neutrally acquired feature would be lost as readily as it was gained. Consistent with a neutral explanation, however, complex features have in fact been repeatedly gained and lost in many protein families. For example, many multimeric enzymes exist in a wide array of stoichiometries, both between and within species, as expected if this feature continues to be neutral. And allosteric regulation has been gained and lost within protein families, as well.

But some complex features are in fact conserved in protein families over very long periods of time. Some of these may have been adaptive as soon as they were acquired, but purifying selection can preserve even neutrally acquired features. For example, complex features that are inconsequential when they originate can become almost impossible to lose if they become functionally important later—a process that clearly occurs in some cases.

Complex features that confer no functional benefit can also persist for long periods of time if they become entrenched and are then preserved by purifying selection. Entrenchment is caused by substitutions that are compatible with the complex state but not with the simpler form; reverting to the simpler ancestral state is then deleterious and is prevented by purifying selection. An apparently widespread mechanism for the entrenchment of multimerization is a hydrophobic ratchet: sites buried in the interfaces of multimers can neutrally accrue hydrophobic substitutions, but if multimerization were subsequently lost these residues would be exposed to solvent, causing instability and/or aggregation. This process is expected to be nearly universal, because the mutational process, owing to the genetic code, has a propensity to produce hydrophobic residues at a frequency far higher than exposed surfaces on monom can tolerate. A recent analysis showed that a large majority of known multimers are likely entrenched by this mechanism and would persist whether or not they have a useful function.

Functionally inconsequential elaborations of a protein’s fold can also be entrenched by a similar ratchet: hydrophobic substitutions can occur neutrally in the buried regions that mediate contact between the ancestral surface and the novel decoration, which makes loss of the decoration subsequently deleterious. Activation by an allosteric effector could also easily become entrenched, because mutations that reduce the stability of the active conformation in the effector’s absence could accrue neutrally but make reversion to a constitutively active ancestor deleterious.

### 6.3 Purifying selection against complexity

If complex features can be so easily acquired and then entrenched, why doesn’t every protein contain end layers of tertiary structure, multimerize into 256-mers, and respond allosterically to hundreds of effectors at sites all across its surface? One answer is that new complex features may often be removed by purifying selection because they are deleterious. New multimeric interactions may obscure active sites, compete against other biologically important interactions, or produce toxic aggregates or fibrils, as in sickle cell disease. Purifying selection clearly removes many fortuitous interactions, as demonstrated by the finding that E. coli proteins form more complexes with heterologously expressed human proteins than they do with themselves. Similarly, mutations that confer allostery may inhibit essential protein activities or upregulate them in biologically disruptive times or places. And mutations that make a protein’s tertiary structure more elaborate or confer an entirely new fold will by necessity create new surfaces, potentially leading to aggregation or disruptive interactions with other molecules.

The limiting factor in the evolution of protein complexity therefore appears not to be how difficult it is for mutation to produce complex features. Rather, the major brake on the evolution of complexity may be incompatibility between new complex features and the biology of the cell or organism in which they arise. Proteins are constantly bombarded by a hail of mutations, some of which
create new interactions, modes of regulation, or changes in fold and conformation. Occasionally these may immediately enhance a biological process, but more often they will be deleterious—and so removed by purifying selection—or neutral. If the neutral features are not rapidly lost, they can become entrenched despite being functionally gratuitous or, through additional mutations in the future, become incorporated into some biologically significant function.

7 | FUTURE DIRECTIONS

The evidence that we have reviewed here establishes that proteins can acquire complex features by short paths, and in numerous historical cases they have actually done so. But the frequency with which this scenario actually occurs is not empirically known. Several lines of experimental inquiry can help to better sketch a broadly applicable account of the genetic and biophysical mechanisms underlying the evolution of protein complexity.

First, we should more thoroughly catalog historical gains and losses in complex protein features. We have a handful of case-studies in which complexity was gained along some lineage, but our understanding of natural variation in multimerization and allostery among related proteins in a phylogenetic context is very sparse. Vertebrates account for most of the crystallographic data in the PDB, despite making up <5% of animal diversity and a tiny fraction of all species. With improved taxonomic sampling and biochemical characterization of proteins, it should be possible to identify far more cases in which multimerization, allostery, and changes in tertiary structure were acquired, and to isolate those changes on relatively short phylogenetic branches. Doing so would set the stage for detailed biochemical and genetic analyses of parent and child ancestral proteins to identify the mechanisms by which the complex feature was gained. There may even be variation in multimeric state, allosterey, and protein structure within natural populations, but there has been virtually no effort to address this possibility outside of humans.

Second, we need more data on the degeneracy of complex protein features and the number of mutational paths by which they can be acquired. A handful of studies have used deep mutational scanning and experimental evolution techniques to understand the sequence-function map around extant and ancestral proteins for a few kinds of molecular interactions and forms of allostery. But we know of no DMS scanning methods at present for homo-oligomerization or for allosteric control of most functions. Developing high-throughput bulk assays in which protein sequence can be linked to these features would help to illuminate these issues. An additional limitation is that most of these mutational scanning studies probe only the immediate sequence neighborhood of one or a few proteins. Advances in the throughput and speed of these assay techniques—and improvements in computational prediction of protein structure and function—may soon allow us to characterize larger fractions of sequence space. New computational approaches to predicting protein folds may also help to address a similar question about protein folds: how extensive are the sequence networks encoding individual folds, and how far are their borders from those of other folds?

Finally, there are other complex protein features, the evolutionary origins of which we would like to understand mechanistically. Historical and directed evolution studies have begun to identify relatively simple means by which catalysis can be acquired de novo. There are many other protein features that may also be tractable to evolutionary and biochemical analysis, including how channels and pores evolved the capacity to pass substrates through membranes and how proteins evolved fluorescent and light-sensing capacities. Particularly exciting would be to reveal how multimeric molecular machines acquire their capacity to transform chemical energy into organized kinetic work. In each case, if a phylogenetic approach can identify intervals in which a property of interest first emerged, an historical biochemical analysis, together with DMS, may be able to identify the causal genetic and biophysical changes that mediated these events.

Tracing the origins of complexity is challenging, but it is, in part and at least for some protein systems, experimentally tractable. Studying complexity at this most fundamental level of biological organization may reveal principles that pertain at higher levels, such as the cell, tissue, organism, society, and ecosystem. Also, higher levels of biology are themselves built upon molecular innovations; understanding how and why proteins came to have their present-day properties is therefore a key step in building a mechanistic account of the evolution of the more visible forms of biological complexity. Given the pace of progress in recent years, there is reason to hope that, by working together, biochemists and evolutionary biologists can advance our understanding of one of biology’s great and enduring sources of wonder.

AUTHOR CONTRIBUTIONS
Arvind S. Pillai: Conceptualization (equal); investigation (equal); writing – original draft (equal); writing – review and editing (equal). Georg K.A. Hochberg: Conceptualization (equal); investigation (equal); writing – original draft (equal); writing – review and editing (equal).
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No primary data were generated for this review.

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**ENDNOTE**
* For a protein that can homodimerize, the fraction of molecules in the bound state is given by the second-order equation:

\[
f = \frac{2}{c} \frac{e^{-\frac{\Delta G}{RT}}}{e^{-\frac{\Delta G}{RT}} + \sqrt{e^{-\frac{2\Delta G}{RT}} + \frac{8c}{4} e^{-\frac{\Delta G}{RT}}}}^2,
\]

where \( \Delta G \) is the difference between the Gibbs free energies of the bound and unbound states, \( R \) is the universal gas constant, \( c \) is the concentration, and \( T \) is the temperature.

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