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De novo protein design: how do we expand into the universe of possible protein structures?
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Protein scientists are paving the way to a new phase in protein design and engineering. Approaches and methods are being developed that could allow the design of proteins beyond the confines of natural protein structures. This possibility of designing entirely new proteins opens new questions: What do we build? How do we build into protein-structure space where there are few, if any, natural structures to guide us? To what uses can the resulting proteins be put? And, what, if anything, does this pursuit tell us about how natural proteins fold, function and evolve? We describe the origins of this emerging area of fully de novo protein design, how it could be developed, where it might lead, and what challenges lie ahead.

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Introduction and scope of this review
Why study protein structures beyond those presented to us by nature?
It is now clearly established that the number of protein folds evolved through and used by biology is limited, and might comprise 1000–10000 different types [1–4]. (Herein, a protein fold is defined as the arrangement of secondary structure elements (SSEs) relative to each other in space.) That natural protein structures are limited stands to reason from the following argument.

Nature could not have explored all of the possible protein sequences or structures over the course of evolution. Many calculations have attempted to illustrate this, but all have caveats [5]. Even with many orders of magnitude shaved off such calculations, which attempt to enumerate the possible permutations of sequences and SSEs, we would be left with a mind-boggling number of molecules compared with say the estimated number of atoms available in the observable universe. In short, natural proteins potentially represent a tiny amount of the possible sequence and fold space. Thus, it is unlikely that nature evolved proteins by sampling this space exhaustively, and that this process was directed in some way; indeed, modern proteins likely arose through assembly and concatenation of smaller fragments [6,7]. On this basis, the vast majority of the possible protein sequences and structures have not been tested by evolution. However, some of these could be evaluated by de novo protein design, and potentially provide solutions to new protein-structure/function targets.

In the context of protein redesign, natural proteins do provide an extremely powerful toolkit and starting points for engineering new attributes and functions into protein architectures [8–12]. However, in terms of genuinely de novo protein design they put up borders between the known and immediately accessible protein world, and what might be out there to explore; rather like the perimeters of the cities in Logan’s Run [13] or The Truman Show [14]. For the majority of this review, we focus on two related questions: First, if natural protein structures are not the only possibilities, what other protein folds are there? Second, how can we access these computationally and experimentally? We refer to such proteins that may only be accessible through design as fully de novo proteins. That said, we do this in the context of what has been achieved in de novo design thus far, and many of these designs should still be considered as being close to observed natural protein folds.

Current estimates of the number of natural protein structures
By the end of 2014, the RCSB Protein Data Bank (PDB) held >105,000 high-resolution protein structures [15]. The most widely used protein-structure classification systems, CATH and SCOP [16–18], suggest that these are accounted for by ≤1400 different protein folds, usually defined as the arrangement of secondary structural elements in space. The rate of discovery of new folds appears to be low: no new folds from CATH have been deposited in the PDB since 2012 [15], and an extended version of
SCOP [18] has reported just 15 new folds since 2009. There are problems with such analyses, however, for instance: how should protein folds and domains be defined [19]? Is the PDB subject to skewed sampling effects? And even that new folds are no longer being registered. As a result, there are disagreements on how many natural protein folds there might be, with estimates of up to 10,000 [1–4]. Nonetheless, the message is clear: natural proteins employ a limited set of 3D protein folds over again.

**Enumerating and organising the protein-space that is possible**

The above realisation leads to our first question: *what protein structures are possible beyond those presented to us by nature?* There are different ways to frame and consider this question [5]. One straightforward approach is to enumerate how many ways multiple SSEs — that is, α-helices and β-strands — can be combined in a linear chain. Figure 1a–d. For n SSEs, there are \((n - 1)! \times 2^{2(n - 1)}\) such permutations. We recognise that this is somewhat naïve, not least because it ignores 3D arrangements of secondary structures — that is, the overall protein fold and domain organisation — and also isolated β-strands tend not to be stable. Nonetheless, the calculation serves a purpose.

The resulting numbers of permutations would be experimentally manageable for chains with very few SSEs \((n \leq 3)\). However, even for modest polypeptide chains with six SSEs, of which the 76-residue ubiquitin is an example, there are 245,760 permutations, with ubiquitin as just one of these. The same exercise on a chain with 16 SSEs, as in the (α/β)₄-helices or TIM-barrels, gives \(>2.8 \times 10^{21}\) possible permutations. Again, the TIM-barrel is just one of these. True, this is a beautiful structure of alternating α-helices and β-strands. It is likely that its meandering topology and consolidated β-barrel aid its folding, stabilization and functionalization. In turn, this may help explain the predominance of the TIM barrel in nature, where it accounts for 10% of all of the known enzyme structures [20,21]. In these respects of folding kinetics and adaptability for function, it may be a privileged fold, or at least one very good solution to the protein folding/function problem. However, it is probable that within the galaxy of the 2.8 \(\times 10^{21}\) 16-element chains there will be other stable folds. The distribution of numbers of SSEs in known protein structures and domains is shown in Figure 1e.

Attempts have been made to rationalise and organise this space for specific protein folds such as four-helix bundles [22,23]; β-sandwich folds [24–26]; and structures that have similar arrangements of secondary structures neglecting the path or topology of the protein chain through these [27]. The most ambitious of these comes from Taylor, who has organised many of the possible 3D arrangements of secondary structures, referred to as basic forms, into a periodic table of protein structures [28].

**Unexplored protein space**

Figure 1

![](image)

Possible permutations and observed numbers of the two main secondary structure elements (SSEs) in protein chains. (a–d) A simple calculation of the number of possible permutations, illustrated in (a–c) for three SSEs. In these models, the left-most SSE is the N-terminal one. (a) In a sequence of n SSEs, each one can be either α-helix (red cylinder) or β-strand (blue arrow), and there are \(2^n\) different orderings of these. (b) For a given ordering (e.g., circled in a), there are \(2^n\) orientations. Half of these are symmetry related, giving \(2^{n-1}\) unique orientations. (c) For a specified ordering and orientation (e.g., circled in b), there are \((n-1)!\) possible paths through the remaining \((n-1)\) SSEs. (d) Schematic for the general case of n SSEs. At each position there are two choices of SSE, and each adjacent pair of SSEs can be arranged in a parallel or antiparallel orientation. With the left-most SSE defined as the N-terminal element, there are \((n-1)!\) ways that the remaining \((n-1)\) elements can be arranged in the primary structure of the protein. This calculation gives a total of \((n-1)! \times 2^{2(n-1)}\) permutations. (e) Observed numbers of SSEs in non-redundant (<=40% sequence identity) protein chains of the PDB (blue), protein domains in SCOP (green), and those domains classified in SCOP as TIM barrel (red). α-Helices and β-strands were identified by Promotif [29].
The possibilities available to protein sequences and structures are often referred to synonymously as the protein universe [3,30]. We focus here on 3D protein folds rather than sequences. The difference between what is possible in protein-fold space, and what has been explored by nature — or at least what we have glimpsed of this so far — has been referred to by different names. Luisi calls these never born proteins. However, he is mainly concerned with generating random de novo sequences, and then finding structure and function within these [31]. Thus, this is more related to the studies of Szostak and others who select functional sequences from random libraries of proteins, than the structural resolution that we seek herein [32]. With this structural focus, Taylor calls the difference the dark matter of protein space [33]; Baker considers it part of post-evolution biology, which might be considered analogous to synthetic biology [34]; and we prefer fully de novo proteins.

Several groups are delving into this new space computationally. Taylor et al. have generated a large number of three-layer α–β–α structures in silico [33]. The starting points are known structures with this overall architecture, and multiple-sequence alignments made from these. Ambiguities in the latter, along with introduced variations, allow new protein topologies to be generated. Herein, protein topology refers to the string of secondary structures and the connectivities between these. In this way, protein-fold space is explored through the loss or gain of SSEs relative to the parent structure. The new topologies are then used to generate 3D models, which are compared in various ways to the known protein structures. Although this searches protein structures locally, the vast majority of the generated structures are new and unrelated to any known protein structures. Moreover, they are protein-like, and potential candidates for experimental designs.

In a different approach, Cossio et al. explore compact protein-like conformations accessible to a polypeptide chain of 60 valine residues through molecular-dynamics simulations [35]. This produces a large number (≈7000) of tangible, independent protein topologies. Although all of the known topologies for natural proteins of a similar size appear, they represent only a small fraction (≈300) of the full set. The simplicity of these models precludes immediate experimental validation, but, again, it illustrates that considerable structural complexity is possible beyond natural structures observed so far.

Moving from in silico design to experimental testing: parameterisation of protein structures

A key question for this new area of fully de novo protein design is: how do we move from the theoretically anticipated dark matter of protein-fold space to its experimental exploration? To put this into perspective, there are two clear advantages of designing de novo proteins that mimic, or closely resemble natural proteins: First, natural backbone structures can be used as templates to start the design process. Second, sequence-to-structure relationships and/or statistical forcefields can be gleaned from natural proteins to guide the design of sequences. There are no such templates or relationships for complete de novo design; although, the statistical forcefields should still be useful for assessing any designs in silico.

Therefore, how do we kick-start the design process to move into the dark matter of protein space? There is hope: protein structures are modular, which opens possibilities for design approaches that employ secondary or supersecondary structures as building blocks. It helps that the two main secondary structures have regular backbone geometries and are scalable; that is, they are readily parameterised. Thus, in principle, larger protein structures and assemblies can be built through non-covalent association, or single-chain concatenation of standardised designs for regular smaller elements. Indeed, considerable progress is being made here, in what might be considered a combination of supramolecular chemistry and protein design/engineering. Several groups have achieved impressive, beautiful, and in some cases functional peptide and protein-based suprastructures based on de novo peptides [36,37,38*], and engineered proteins [39–41,42*, 43].

Returning to generating completely de novo protein folds — where, arguably there is only one example to date, namely TOP7 from the Baker lab [44] — there are three main challenges: first, how do we position elements of secondary structure in space to produce new structures? That is, is this space reducible and parameterizable in any way? Second, how do we cement specific and stable interactions between elements of secondary structure? Third, how do we link the building blocks up with turns and loops to make single-chain polypeptides that direct folding as desired? The latter, loop-design problem often thwarts both protein engineering and design projects and solutions are actively being sought [45,46], but we will not address it directly herein.

Turning to the first challenge, the parametric description of protein folds has been a goal of protein scientists for decades [28,47–49]. These offer tangible routes to limit the structural space that must be searched to achieve novel protein folds. Similarly, the second challenge has been addressed in terms of understanding helix–helix and strand–strand interactions in natural protein structures for some time [50–52]. However, in both cases, there has been less effort and success in translating this understanding into de novo contexts. One way forward initially is to consider structures of high symmetry. Several protein structural types present as clear candidates here, for instance all-β (porin) and αβ-type (TIM) barrels, solenoid structures such as the leucine-rich repeats and so on.
Arguably, the α-helical coiled coil is the best-understood repeat structure [53,54], and, as such offers an ideal starting point for parametric design. In α-helical coiled coils two or more α-helices associate into rope-like bundles. This is programmed at the sequence level by variations on the conspicuous signature of coiled coils; namely, the heptad repeat of hydrophobic (H) and polar (P) residues, HPPPPP. Coiled coils have been the subjects of many protein design studies [54,55]. Much of this work has focused on sequence-based designs, which use and embellish the heptad repeat [53]. Therefore, it is easy to forget that Crick’s original postulate was as much about the geometry of helices and helical packing, as it was about the sequence patterns that might direct helix–helix interactions in coiled coils. Moreover, Crick’s parametric equations, which elegantly describe classical coiled coils using just four parameters, have been demonstrated to reproduce the vast majority of subsequently experimentally determined coiled-coil structures very faithfully indeed [56,57]. As such, Crick’s parameters provide an excellent basis for the modelling and design of de novo coiled coils in silico [58–60]. Most recently, two web-accessible and user-friendly tools have become available to make coiled-coil modelling and design accessible to all. These are CCCP and CCBuilder [56,57]. They widen the possibilities for modelling and designing new coiled-coil structures to the complete periodic table of natural coiled coils [61] and beyond.

Before moving onto more-recent protein design studies that are pertinent to our line of reasoning in the remainder of this review, we should like to make one thing clear: we do not regard most of the following examples as fully de novo protein designs as defined above. Some of them directly mimic natural protein folds; others are clear variations on natural themes; and only one is, as far as we can tell, unprecedented in the natural protein-fold space. That said, the studies are the state of the art, and they encourage us that computational routes into the dark matter of protein-fold space will follow.

**Parametric protein designs achieved so far**

Parametric structure-based designs have been realised before now for coiled-coil proteins, including a right-handed structure that incorporates non-natural amino acids to satisfy unusual packing in the interior of the helical assembly [62], Figure 2a. However, CCCP, CCBuilder and methods developed in parallel in the Baker lab [56,57,63] allow structure-based designs of coiled coils and helical bundles to be tackled by more groups. A key feature of all of these methods is that they implicitly incorporate backbone variations, which has always been a challenge in computational protein design [56,57,62,63].

Grigoryan, DeGrado and co-workers have used CCCP to generate helices predicted to make barrel-like assemblies around carbon nanotubes [37]. Though still waiting for full structural characterisation, this is an exciting achievement with potential applications in solubilising carbon nanotubes for applications in bionanotechnology. More recently, the same group applied the parametric approach impressively to design functional membrane-spanning helical bundles [65]. ‘Rocker’ is a four-helix bundle with two zinc-binding sites. The rationale is that zinc ions presented to one side of the membrane (the cis side) are transported to the other side (trans) via the two sites facilitated by the rocking of the structure such that it opens on the cis side, binds zinc at the first site passes it onto the second, and then exits the trans side of the membrane, Figure 3a. The group have characterised the designs to a high level of detail with a combination of liposome ion-flux assays, X-ray crystallography (of an apo, dimeric form) and NMR spectroscopy; although a high-resolution structure of the complete, functional, four-helix target remains elusive. DeGrado’s group has also attempted to design peptides that switch between water-soluble and membrane-spanning peptides on lowering pH from 7.4 to 5.5 [66], Figure 3b. In assays with red blood cells, the low-pH states facilitate the release of ATP and miRNAs, but not haemoglobin. Further structural details are needed to validate the designs, and it is curious why the oligomer states of these peptides collapse in membranes, but still conduct small and macromolecules.

Baker’s group has achieved parametric designs for hyper-stable coiled coils [63]. These are water-soluble assemblies including single-chain, three-helix and four-helix bundles that include both parallel and antiparallel helix–helix contacts, and a non-covalent, parallel five-helix bundle, Figure 2b. A key design feature is the structural focus on layers of hydrophobic residues that define the hydrophobic core, rather than more-traditional sequence-based repeats. By considering two-layer, three-layer and five-layer structures, which correspond to traditional seven-residue, 11-residue and 18-residue sequence repeats, the designs are for a non-covalent pentamer with a left-handed supercoil, and of single-chain constructs for a right-handed four-helix bundle, and a three-helix assembly with straight helices, respectively. These are confirmed by X-ray crystallography, Figure 2b. Interestingly, although designed using computationally using Rosetta, some of the packing solutions mimic those for natural and canonical designed coiled coils [54]; though not all have complete knobs-into-holes packing between helices as judged by SOCKET [57].

The third team is our own. Following the serendipitous discovery of a non-covalent coiled-coil hexamer (CC-Hex [68]), we have built parallel pentamers and a heptamer, CC-Pent and CC-Hept, and other hexamers by computational design as follows [64]. The new designs are based on the realisation that successive helical interfaces in the cyclic structures can be approximated as hetero-
Structurally resolved, computational designs of coiled-coil assemblies and helical bundles. (a) A left-handed tetrameric assembly that incorporates non-natural side chains to achieve unusual packing in the hydrophobic core (PDB identifier 1RH4 [62]). (b) Hyperstable structures for three-helix and four-helix single-chain designs and a pentamer from the Baker laboratory (4TQL, 4UOS, and 4UOT, respectively [63**]). (c) Three designed \( \alpha \)-helical barrels, CC-Pent, CC-Hex and CC-Hept (4PN8, 4PN9, and 4PNA, respectively [64**]), which have clear central channels.
dimeric faces encoded within the same peptide chain. In short, the helical interface is extended to span two helix–helix interfaces, in a so-called ‘Type-II’ bifaceted coiled coil [53,69]. Next, sequences are selected computationally from one million alternatives. Then, ahead of experimental validation, the selected sequences are modelled in CCBUILDER to predict the preferred oligomer state in the range tetramer to octamer. In this way, we have extended the reach of de novo coiled-coil design past canonical dimers, trimers and tetramers [70], which are plentiful in nature, into fully de novo pentamers and above for which there are few or no natural examples, Figure 2c. The structures are also noteworthy, and potentially useful, because they all have central channels the diameters of which scale with the number of helices in the assembly — the channels of CC-Pent, CC-Hex and CC-Hept are approximately 5, 6 and 7 Å across, respectively, [64**] Figure 2c. Thus, there is potential for such structures in the rational design and redesign of functional binding proteins, enzyme-like catalysts, membrane-spanning ion-channels, and peptide-based materials [71,72].

Future challenges for this aspect of the field include: the design of more structures, that is, populating empty elements of the periodic table of coiled coils [61], and beyond into the dark space of coiled-coil structures; the incorporation of dynamics, which will be key to delivering functional designs, but are poorly understood; and the further introduction of binding and catalytic functions [71,72].

**β-Propellers, solenoids and other potentially parameterizable protein structures**

Moving into protein-fold space more widely will present more challenges. The coiled coil has advantages of symmetry, inherent stability, and clear parameter sets for defining and creating the possible backbone scaffolds. For other protein structures — both natural and dark — symmetry is reduced, stability is less guaranteed, and there are few or no parameter sets. Nevertheless, good progress is being made.

Tame and colleagues explore the design of new β-propeller folds, dubbed Pizza proteins [73*]. They start with a single blade from non-symmetric six-bladed propeller, from which they generate a fully symmetric six-bladed structure by blade duplication (the pizza slices), concatenation and optimisation in RosettaDock. The high-resolution X-ray crystal structure of the resulting single-chain protein, Pizza6, is closely similar to the designed model, Figure 4a. Shorter polypeptides, Pizza2 and Pizza3, trimerase and dimerise, respectively, to give six-bladed structures again. This demonstrates the robustness of both the designed blade and the approach. It also tallies with how multiply blade structures may have arisen during evolution through gene duplication. Interestingly, larger sequences encoding >6 blades, form higher-order assemblies, which are mostly reconciled as discrete aggregates of the six-bladed design target; although Pizza7 behaves in more-complex ways.

Different curves and twists on another repeat protein are described by the groups of André and Baker [74**,75**]. They have both targeted the leucine-rich repeats LRRs, which form an array of horseshoe-like proteins that bind macromolecules via their inner surfaces formed by parallel β-strands connected on the outer faces by α-helices, Figure 4b. Rather than focus on similarities between repeats, and develop consensus sequences for a repeat, which has been successful in generating both structured and functional solenoid designs [76–78], both groups employ structure-based design to control the curvature (projected angle between successive repeats in the structure) and twist (the angle out of the plane between each segment). In this way, the groups aim to produce structures with all manner of shapes, combining set curves and twists.
between repeats, to deliver high-affinity and high-specificity de novo binding proteins for any macromolecular target.

André’s group use a repeat of the ribonuclease inhibitor (RI) fold as a starting point, Figure 4b, maintaining many of the hydrophobic-core residues that specify the local, repeat units [74**]. The other residues are mutated in silico to select combinations that should give a specified curvature between repeats, and zero twist. (In these respects, this approach combines elements of redesign and de novo design.) Concatenation of 10 self-complementary units, with additional capping units at the termini, into a single polypeptide gives stably folded and discrete proteins, although high-resolution structures will be required to verify the designs completely. Encouragingly, leaving out the capping units produces dimers, which are of the right dimensions in solution and by electron microscopy to fit the design hypothesis that the protein forms a flat semi-circular structure, two copies of which can interact head to tail to complete a circular, donut-like structure.

The approach of Baker’s group is subtly different [75**]. They target a series of de novo self-complementary repeat modules, which when homo-oligomerized give defined curvatures. In addition, they develop junction, or wedge units to allow different repeat modules to be linked together to alter curvature within a single protein structure. Thus, in principle, these building blocks can be mixed and matched to create an array of polypeptides and define different shapes on demand. These could be regular, as with the design from the André group, or irregular. In this way, different binding surfaces could be built up for specific targets. The versatility of the approach is illustrated with 12 new proteins complete
with X-ray protein crystal structures [75**, Figure 4c–f. This approach is reminiscent of the drive in peptide self-assembly to generate toolkits of peptide-based building blocks that can be characterised once and then used in different contexts [36*,70].

What will we learn about natural proteins through this emerging area?

The topic of the evolution of protein folds is hotly debated [81]; as is the notion of what a fold is precisely, and how to classify them [82]. However, in evolutionary terms natural protein structure space is often considered discrete, whereas in terms of possible protein structures the space is by definition continuous. Grishin does not see these at odds, but simply as part of a duality in our understanding and theories of protein structure [81]. One emerging view is that this arises because evolved stable protein structures represent islands of stability within a sea of instability [83]; although, the sea does appear to be furred in places [84]. The question is: can protein designers build more islands and links de novo?

In Arrival of the Fittest [85], Wagner describes how evolution has bridged large spaces through connected genotype networks. This is as true of metabolic pathways, as it is for nucleic acid and protein-sequence spaces. These networks allow innovation — that is, the exploration of new genotypes and generation of new phenotypes — but, at the same time, the evolving systems can survive. Another way to put this is that for a given natural sequence or function there are many similar solutions nearby in sequence space, but it may be hard to escape from such regions.

In short, natural systems are robust. This presents a problem for protein redesign and design: if natural proteins are over-determined, are they necessarily good starting points for new designs, structural or functional? In other words, if point mutations or small numbers of changes keep you where you are, how can we move into completely new territory? If this is correct, and if we are approaching the limit of innovation with natural protein sequences and structures as scaffolds, perhaps it is time to look to the dark side? So, is there hope? Wagner also makes the point that there are multiple solutions to a given phenotype, and that sequence space is vast. Thus, solutions to the phenotype problem are not special or privileged per se; they are inevitable. If this extends to fully de novo protein folds, we should not have any problems in discovering them.

What will we do with fully de novo proteins once made?

There are clear uses for some of the structures described in this review. For example, a reliable set of α-helical barrels could be used as the basis for introducing binding or even catalytic properties into protein lumens of defined size and chemistry [55,71]; and, if polymerised, to produced peptide and protein-based nanotubes [72]. For the solenoid-type folds macromolecular binding presents the mostly likely function to be targeted [76,78]. Whereas access to β-propeller folds with different numbers of blades may open routes to many different functions [86]. More generally, access to a wide variety of robust, de novo protein folds — for which we have a clear understanding of their sequence-to-structure relationships, and that are free from sequence constraints or idiosyncrasies from millions of years of evolution — will provide future protein engineers with an extremely versatile toolkit of scaffolds to graft functions onto.

That said, with the exceptions of TOP7 and CC-Hept [44,64**], all of the de novo structures described so far are mimics of, or variations on natural protein folds — what is needed now is a push towards fully de novo folds. Aside from exploring new functions, this search for and study of such entirely new proteins is legitimate in itself for a number of reasons. Primarily, as with the entire ethos and origins of protein design, we pursue such goals as the ultimate test of our understanding of protein structure, folding, assembly and function; that is, after Feynman, ‘What I cannot make, I do not understand’. Extending this objective, the pursuit of entirely new protein structures, pushes the boundaries of what may be possible with seemingly straightforward polypeptide chains; the restrictions that are imposed on chain folding by Ramachandran; and, if we stick with proteinogenic amino acids, what can be encoded with just 20 amino acids. Furthermore, the exploration of such de novo proteins and assemblies fits with the emerging field of synthetic biology, for which there are high hopes in terms of new innovation and applications in biotechnology, medicine, materials science and beyond.

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Engineering and design

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