Neural network models
of protein domain evolution

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Abstract: Protein domains are complex adaptive systems, and here a novel procedure is presented that models the evolution of new functional sites within stable domain folds using neural networks. Neural networks, which were originally developed in cognitive science for the modeling of brain functions, can provide a fruitful methodology for the study of complex systems in general. Ethical implications of developing complex systems models of biomolecules are discussed, with particular reference to molecular medicine.

Keywords: models in biochemistry, protein domain evolution, neural networks, ethics of modeling.

Introduction
Everywhere in nature, matter organizes itself into complex patterns and structures. This self-organizing tendency of matter culminates in biological organisms and their constituent macromolecules, the most complex chemical entities known to us. Proteins are by far the most abundant and diverse class of biomolecules and mediate the vast majority of biochemical processes. With the recent explosion of protein sequence data from all three kingdoms of life, the archea, prokarya and eukarya, we have come to even more fully appreciate the modular nature of proteins, and the complex ways in which their functional and structural units, protein domains, are conserved and recombined during evolution (Fig. 1). Domains are thermodynamically stable and fold independently within the context of the whole protein. Novelty in protein function often arises as a result of the gain or loss of domains, or by reshuffling existing domains along the linear amino acid sequence. Thus, protein domains can arguably be seen as stable units of evolution.

New domain functions evolve within the constraints of maintaining thermodynamic stability and autonomous folding capability. This gives rise to a complex interplay of molecular organization and evolutionary dynamics,
which is still a largely unexplored area of research. My aim in the present paper is to approach this problem from a perspective informed by recent developments in complexity theory. This work employs distributed representation by neural networks in building relational models of protein domain evolution. I will also address the explicit ethical dimension inherent in choosing and developing models in biomedical science. Ethical implications of developing complex systems models of biomolecules are discussed on this premise.

![Cartoons of some widely found protein domains.](image)

**Figure 1**: Cartoons of some widely found protein domains. Alpha helices are shown as **barrels**, beta-sheets as **arrows**, and loop regions as **lines**. Protein Data Bank (PDB) identifiers of the crystallographic structures are given following the domain names.
1. Protein domains are complex adaptive systems

Can protein domains legitimately be classed as complex adaptive systems? At present, a consensus on the characteristics of complex systems is still elusive, both qualitatively and quantitatively, however, the following characteristics have found general agreement (Cilliers 1998, p. 3) and are present in protein domains:

1. **Complex systems consist of a large number of elements.** At the atom level, protein domains typically consist of thousands of elements. At a higher level of description, the amino acid level, they are comprised of up to several hundred elements. Whilst description and modeling at the atom level is computationally intractable at present, domain systems can be modeled at the amino acid level.

   Please note that, for reasons that will become clear when neural network modeling of domain evolution is discussed below, the positions along the protein sequence, rather than the amino acids themselves, are defined as the elements of the systems. These elements can be in one of 20 different states (be filled by one of the 20 amino acids). The state of an element can change, i.e., a positions can mutate to a different amino acid.

2. **The elements of a complex system interact in a dynamic fashion and these interactions change over time.** Dynamic interactions between amino acids (positions in certain states) mediate the folding process and a stable pattern of interactions subsequently determines the three-dimensional fold of the domain. Dynamic interactions are also fundamental to domain functions that are mediated by conformational changes. During evolution, the pattern of interactions between fold positions changes as a consequence of amino acid substitutions (gain or loss of hydrogen bonds, salt bridges, or van der Waals interactions).

3. **The interactions between elements are richly connected – any one element influences, and is influenced by, a large number of others.** In a domain fold, amino acid positions along the linear protein sequence are engaged in multiple local (involving positions that are close in the linear sequence) and non-local (involving positions that are distant in the linear sequence) physical interactions. With the exception of neutral positions, each fold position makes an individual fitness contribution and simultaneously affects the fitness of many other positions within the domain. Fitness is here defined as the capacity of the domain to maintain its structural integrity and to carry out specific function(s).

4. **The interactions between elements are non-linear. Small causes can have large results, and vice versa.** Complexity results from the patterns of richly connected interactions between the elements. Complex systems exhibit so-called emergent properties, properties that are only seen in systems of an equivalent
degree of complexity. In other words, the behavior of complex systems cannot be derived on the basis of knowledge of their parts. One of the key processes responsible for emergence is self-organization (Cilliers 1998, p. 89; Holland 1998, pp. 115, 225). This co-ordinated behavior results from the non-linear interactions of its components which leads to collective effects. Self-organization also leads to spontaneous transitions into new collective states, at times as adaptive responses to changes in their environment.

The non-linearity of interactions between amino acid positions is a major reason why certain amino acid substitutions at only one or a few positions may unravel a domain fold. And, conversely, is a reason why amino acid sequences can at times diverge from homologous sequences beyond any statistically significant similarity, while the shared domain fold is still conserved intact. We are unable to explain or predict these phenomena (at least for now), and so they also illustrate how non-linearity severely limits predictability. Another related issue is the persistent elusiveness of a solution to the ‘folding problem’, despite three decades of intense efforts.

(5) The interactions between elements are relatively short-range. Physical constraints and information are mostly transmitted between immediate neighbors. However, this does not mean that there can not be long-range influences. In a richly connected network, the path between two elements can usually be covered in a small number of steps. Influences can be enhanced, suppressed, or modulated in some way along the path. Amino acids in domain cores are packed in an energetically favorable arrangement, and strong local constraints on amino acid variation are present. The network of amino acids that are in contact with each other collectively constrains mutational change. Although this mechanism is mediated by local interactions, it can propagate throughout the domain to distant sites via “chains of local interactions” (Lapedes et al. 1997). Non-linear constraint modulation along such interaction chains occurs due to the rich connectivity between elements (multiple physical interactions and mutual constraints).

(6) There are recurrent interaction pathways. The effects of a state change at one element can feed back on itself, either directly or via a number of intervening states. The feedback can be either enhancing or inhibiting. Depending on its nature, a mutation (state change) at one domain position may enhance or inhibit the probability of a particular amino acid substitution (after selection) at coevolving positions. These subsequent mutations may in turn enhance or inhibit further change at the first position.

(7) Complex systems have a history. They evolve through time, and their present state is constrained by their past. Present-day protein domains have evolved from ancestral domains. Domain evolution can only occur within the constraints of maintaining thermodynamic stability and autonomous folding capability.
It can be difficult to precisely delineate the boundaries of a complex system. Boundary definitions are often derived for descriptive purposes and are influenced by the position of the observer. Molecular biology in general follows a top-down approach. Bodies are broken down into tissues, tissues into cells, and cells into molecules, biochemical compounds, and atoms. Reductionism then seeks to explain the functioning of the organism on the basis of the chemistry and physics of its constituent parts. Complexity theory asserts the importance of balancing an analytical top-down approach, indispensable for the identification of the building-blocks of a system, with a bottom-up approach, in order to study how living systems emerge from the laws of physics and chemistry. Whilst all biological processes are consistent with the physical and chemical laws of our universe, and in this sense can ultimately be ‘reduced’ to chemistry and physics, there is a growing awareness among scientists that biological phenomena require an approach that equally addresses the problem of emergence. Emergent phenomena result from the complex, rule-governed, interactions of a large number of biomolecules, in a highly context-dependent manner. Consciousness, to mention a familiar example, arises out of the unimaginably densely connected interactions of billions of neurons, and is not a property of any one brain region, let alone of the neurons themselves. Consciousness is an emergent property of the brain as a whole.

From a different perspective, one which pays close attention to process and interaction across multiple levels of biological complexity, the living world appears as a multidimensional whole of complex systems within complex systems. The demarcation lines between different levels of organizational complexity, and the delineation of any one of these systems, rest on boundaries defined according to criteria that will always, to some extent, be contingent on the perspective of the observer. This notwithstanding, the discussion so far has shown that protein domains can legitimately be seen as complex systems in their own right, far down in a nested hierarchy of proteins, protein complexes, structural and functional networks in cells, whole cells and organisms.

2. Domains evolve as complex adaptive systems:
Hormone-binding domains in nuclear receptors

Ligand-binding sites in homologous protein domains can diverge greatly during evolution. This poses a particularly interesting problem in those cases where the ligand-binding site is situated in, or close to, the domain core, or where ligand-docking induces dramatic conformational changes. These fea-
tures are present in many receptors and enzymes; the hormone-binding domain of the nuclear receptors for steroids and retinoids, for example, exhibits both characteristics. This raises the interesting question how binding sites for diverse ligands evolve in core regions of structurally dynamic domains. Are evolutionary changes locally restricted to the ligand-binding site, or are they distributed throughout the domain?

Steroid, thyroid and retinoid hormones comprise the broadest class of gene-regulatory ligands known. Their receptors belong to the diverse superfamily of nuclear receptors (NRs) that are present in all metazoans from cniderians onward and have had a central part in the evolution of biological complexity since the Cambrian explosion (Escriva et al. 1997, Laudet et al. 1992). As ligand-inducible transcription factors, NRs play essential roles in the regulatory pathways that transmit signals, originating from the extra- and intra-cellular environment, to large genetic networks through a complex sequence of molecular interactions. These genetic networks regulate many aspects of development and function; specifically, higher morphology, the immune system, the nervous system, as well as reproductive and metabolic systems.

The ligand-binding domain of nuclear receptors possesses a unique fold that is partly disordered in the absence of ligand, termed the “antiparallel α helical sandwich” (for refs., see Nagl et al. 1999). The helices are grouped into three layers around an internal ligand-binding core. Crystallographic studies of ligand-bound NRs suggest a structural role for ligand that is fundamental to the allosteric control mechanisms found in the ligand-binding domain. The ligand is completely buried within the domain interior and contributes to the hydrophobic core of the active conformation of the NR. Therefore, ligand binding directs the alignment of the secondary structural elements critical for receptor function, and strongly constrains the conformational freedom of the ligand-binding domain.

During the evolution of the NR superfamily, the ligand-binding pocket has evolved to allow binding of ligands possessing strikingly diverse chemical structures. Escriva et al. (1997) proposed that the ancestor of the superfamily was an orphan receptor without ligand-binding capability. Their study of NR evolution suggests that liganded receptors have arisen relatively recently and have gained the ability to bind ligands independently. Since the ligand-contacting residues line the binding pocket in the domain core, they perform a dual role; a functional role in ligand recognition and a structural role as core residues. With respect to ligand recognition, they can be seen to constitute an ‘interior interaction surface’. In principle, this would allow great scope for the evolution of the ligand-binding pocket. However, since the hydrophobic ligand is an integral part of the domain core in the active conformation, the ligand and the ligand-binding residues combined need to be able to maintain
structural stability and domain dynamics (conformational changes). How is this potential conflict between structural constraints and functional diversity resolved within the domain fold? In an earlier study, it was shown that the ligand-contacting residues in the hormone-binding pocket are evolutionarily linked to an extensive, hierarchically organized, network of coevolving positions (Fig. 2) (Nagl et al. 1999). The nature of the mutations in correlated positions suggests that they compensate for the destabilization resulting from the binding of diverse ligands and preserve the structural integrity and the conformational dynamics of the ligand-binding domain. In conclusion, a distributed evolutionary mechanism, involving the domain fold as a whole, is present in the ligand-binding domains of nuclear hormone receptors. It is suggested that this mechanism maintains a thermodynamically favorable interplay between molecular organization and evolutionary dynamics.

**Figure 2**: Retinoic acid-contacting positions and first-order covarying positions in the ligand-binding domain of the retinoic acid receptor. Ligand contacts are shown in black, covarying positions are shown in grey (α-carbons, spacefill mode). The ligand is shown in black (stick mode).
3. Neural network models of protein domain evolution

3.1 An information-theoretic approach to protein domain evolution

Gene duplication and recombination are thought to be the primary mechanisms for the generation of protein diversity. In this process, one gene copy maintains the original function while the other is free to evolve new functions. The concept of a functional space, as an abstract representation of all possible functions that can evolve within the structural constraints of a domain fold, is useful for the investigation of domain evolution. Within this conceptual framework, the emergence of new functions can be understood as the result of adaptive walks in sequence space. During this adaptive evolution, duplicated genes accumulate successive mutations that progressively enhance the new function.

In this work, where protein domains are studied as complex adaptive systems, the positions along the linear amino acid sequence of the domain are conceptualized as the elements, or 'agents', of the system that can each assume one of 20 different states (i.e., the 20 amino acids) (Fig. 3). Four classes of fold positions can be distinguished in domains that are descended from a common ancestral domain: (i) positions with conserved amino acid identities; (ii) positions with conserved physicochemical properties; (iii) positions with variable physicochemical properties (often belonging to the distributed network of coevolving positions (see Sect. 2); and (iv), unconstrained positions accumulating neutral mutations. Positions in the coevolutionary distributed network to be modeled by neural networks belong to class (ii) or (iii).

![Figure 3: The positions along the linear amino acid sequence of the domain constitute the 'agents' of the system, and can each assume one of 20 different states (amino acids in single letter code).](image)

The evolutionary history of a domain, contained in a sequence alignment, is a record of successful mutagenesis experiments carried out by nature. A multiple sequence alignment indicates the extent to which specific residues may be changed without destroying domain structure. At the same time the alignment can identify those residues that need to be changed in order to create a
new function within a similar structural framework. Coevolving positions can be identified from a sequence alignment of a domain family using mutual information, a measure of correlation for discrete symbols. A formal measure of variability at position $i$ is the Shannon entropy, $H(i)$. $H(i)$ is defined in terms of the probabilities $P(s_i)$, of the different symbols, $s$, that can appear at a sequence position (i.e., for amino acid sequences $s = 20$, for the 20 possible states of amino acid occurrence) (Korber et al. 1993). $H(i)$ is defined as

$$H(i) = -\sum_s P(s_i) \log P(s_i)$$  \hspace{1cm} (1)

Mutual information is defined in terms of entropies involving the joint probability distribution, $P(s_i, s'_j)$, of occurrence of symbol $s$ at position $i$, and $s'$ at position $j$. The associated entropies for each position $i$ and $j$ are

$$H(i) = -\sum_s P(s_i) \log P(s_i)$$  \hspace{1cm} (2)

$$H(j) = -\sum_{s'} P(s'_j) \log P(s'_j)$$  \hspace{1cm} (3)

And the joint entropy is defined as

$$H(i, j) = -\sum_{s,s'} P(s_i, s'_j) \log P(s_i, s'_j)$$  \hspace{1cm} (4)

The mutual information, $M(i, j)$, is defined as

$$M(i, j) = H(i) + H(j) - H(i, j)$$  \hspace{1cm} (5)

If the positions are independent, their mutual information is 0. If, on the other hand, the positions are correlated, their mutual information is positive and achieves its maximum value if there is complete covariation.

Given a set of sequences that are assumed to be independent and identically distributed samples from a probability distribution, one can independently estimate each pairwise probability distribution for every pair of positions by frequency counting. However, sequences belonging to a domain family are not independent samples, but are related through shared ancestry described by a phylogenetic tree. If two mutations occur independently in an ancestral sequence and these are subsequently inherited by many of the descendants further down the tree, the two positions involved will receive a high mutual information score. To estimate the mutual information content between position pairs that is created by tree inheritance alone, and not by covariation, a simulation experiment can be performed (Nagl et al. 1999, Lapedes et al. 1997). This procedure simulates the evolution of sequences by random mutations along a phylogenetic tree obtained from the domain sequence alignment. Using the outgroup as a seed, random sequences are evolved following the phylogenetic tree obtained from the real data set. During simulated random mutation of sequences, the states of the sequences are
duplicated at a bifurcation point in the tree, and the two copies are then independently evolved. Every amino acid can mutate with equal probability to any other amino acid. The procedure is repeated numerous times, and significance threshold values are determined from the frequency distributions of the mutual information scores in the control and real data sets. Any mutual information score greater than the lower boundary value, has a low probability of being caused by inheritance through the tree.

3.2 Neural network modeling of domain evolution

The coevolutionary relationships identified by mutual information analysis can have very high interconnectivity, where each position in the coevolutionary network constrains, and is constrained by, many other positions. This was the case in the nuclear receptor ligand-binding domain (Nagl et al. 1999). Each amino acid is uniquely defined by its physicochemical properties, such as shape, volume, polarity, hydrophobicity and charge among many additional, less well understood properties. Depending on the location of coevolving positions within the network, different physicochemical properties may be crucial in determining the pattern of coevolution. As has been learnt from homologous domain alignments, in many cases volume conservation is of paramount importance (Gerstein et al. 1994), while other properties are less constrained. In other cases, the hydrophobicity value or charge may be crucial; or a combination of several properties. Presumably, the greater the number of properties involved, or the more restricted the allowed range of a single property, the stronger will be the mutual constraints on allowed states for each position. All these factors combined result in constraints of high dimensionality. Modeling coevolutionary networks with artificial neural networks (ANNs) can represent the complexity, and the parallel-distributed nature, of this evolutionary process.

ANNs are computer algorithms that attempt to model the way the brain works and draw on the analogies of adaptive biological learning. One particularly valuable and intriguing characteristic of information processing in biological brains seems to be also present in ANNs – the ability to make decisions based on very complex, noisy, irrelevant and/or partial information. While the comparison with the human brain has led to some exaggerated claims concerning ANNs, this analogy is a very useful way to describe the construction and function of neural nets. An ANN is composed of a large number of highly interconnected processing elements that are analogous to neurons and are tied together with weighted connections that are analogous to synapses. Interconnected neurons, whether biological or artificial, have certain neuro-‘logical’ properties and can be seen as logic gates: They receive input signals from a large number of other neurons, process these signals ac-
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According to specified transformation functions, and produce an output signal as a result of this processing. Brains and artificial neural networks represent information in a distributed fashion; information is encoded by the patterns of synaptic connection strengths (weights) between neurons. The distributed networks of neurons perform many transformation steps in parallel, a style of computation known as parallel distributed processing (PDP). When fully connected neural networks are used, a combination of a large set of connection weights and nonlinear transfer functions allows models of any complexity to be fitted between the response and the input parameters. Neural networks are therefore highly efficient nonlinear data modeling devices, and can be seen as universal models for information processing in complex system. Arguably, the evolution of functional sites within the coevolutionary network of a domain family can be conceptualized as a type of PDP (Fig. 4). It should be well noted that this statement is not meant to imply a direct correspondence in architecture between the coevolutionary network and an ANN, but refers to an analogous information-processing mode. Furthermore, as all parallel-distributed computational steps are executed simultaneously, ANN models of domain evolution do not represent the historical sequence of step-wise mutation at coevolving sites over evolutionary time. This temporal aspect of coevolutionary networks can be analyzed and modeled by reconstruction of ancestral states by parsimony.

For the purpose of building an ANN model of a coevolutionary network, we return to our previous representation of a protein as a chain of agents in a linear sequence, each of which can take on one of 20 states (amino acids) (Fig. 3). The agents are understood as mechanisms for mediating interactions (Holland 1998, p. 6), and state transitions in agents (mutations) lead to a modification in the patterns of interactions, sometimes resulting in a change in structure/function. The state transitions are constrained by rules (Holland 1998, p. 116), and all possible state sequences are the outcomes of a succession of transitions specified by these rules. In this way, the rules generate evolutionary novelty. Structure/function can now be re-conceptualized as an emergent property, the result of context-dependent interactions, that changes over time.

It is possible to encode the state transition rules in the values of the connection weights of an ANN model of the coevolutionary network. Specifically, the evolution of new functional sites within the coevolutionary network can be modeled by a classical fully-connected feedforward neural network (Fig. 4) (for a detailed mathematical treatment of feedforward network properties and behavior, see, for example, Skapura 1995, Mehrotra et al. 1996, Livingstone et al. 1997).
Figure 4: ANN model of the evolution of new functional sites in a domain family. The network architecture is that of a classical feedforward network whose size can vary dependent on the coevolutionary network to be modeled (closed arrows). Sequence positions (agents) function as fully connected processing elements (squares). Each agent is represented as a binary or real number vector (open arrows; see below). A heteroassociative mapping is performed that maps the input vector matrix (agent states in the functional site) to the output vector matrix that ranges over a different vector space (states of coevolving agents). After training, the ANN encodes the state transition rules of the coevolutionary network.

An important decision to be made concerns how to encode the states of the agents. To name just two alternatives, they can be encoded as binary vectors (bitstrings), or as vectors of real numbers (any value between -1 and 1), depending on which aspect of the states we wish to model. If we want to encode amino acid identities (A, W, S, D, etc.), bitstring encoding and a discrete ANN model suggest themselves as the most appropriate choice. If we want to encode information about certain physicochemical properties of the amino acids (hydrophobicity, hydrophilicity, charge, polarity, volume, etc.), this can be achieved by using real number vectors, where each property is expressed by a normalized value between -1 and 1, and a continuous ANN model.

The inbuilt directionality of this type of neural net corresponds to selection pressure on the domain for evolving new functions. During training, the
network is presented with instances of functional sites (input) and associated amino acid identities at coevolving positions (output) taken from domain family sequence alignments, and trained to associate outputs with input patterns. When the network is subsequently used for modeling, it identifies the input pattern and tries to produce the associated output pattern. The power of neural networks comes to life when a pattern that has no output associated with it, is given as input. In this case, the network predicts the output based on the rules learnt in the training phase. This property is responsible for the power of a neural network in evolutionary modeling. When, for example, given an artificially designed functional site as input, it will predict the likely compensatory states of the coevolving agents based on the learnt state transition rules. The ANN modeling procedure is expected to be valuable for the design of novel ligand-binding capabilities for a given domain fold. ANN modeling, based on the coevolutionary relationships between ligand-binding sites and coevolving positions, may enable one to overcome otherwise prohibitive limits to binding-site modifications. Predictions of fold-stabilizing mutations located at coevolving positions throughout the domain may be used to maintain the stability of the modified fold.

4. Ethical implications of model choice
Biomolecular engineering is fast acquiring the technical know-how for the design and large-scale manufacture of novel proteins. This current progress in engineering goes hand in hand with revolutionary advances in molecular medicine, that are generating an unparalleled increase in knowledge about human diseases. In the future, it will be possible to design proteins for novel therapeutic properties, and these ‘designer molecules’ will make up a major part of the new molecular materia medica. Accompanying these developments is a far-reaching conceptual shift that is leading to a radical redefinition of the human body as a hugely complicated molecular machine. This new vision of human biology, with its concomitant engineering approach to the treatment of disease, has profound ethical implications, as it increasingly determines how we choose to intervene in the functioning of the body. Within such a framework, knowing, representing and intervening can clearly not be separated. We represent in order to intervene (Hacking 1983). The demarcation line between ‘pure’ and ‘applied’ science becomes ever more illusionary. Therefore, any choice, concerning the modeling of biomolecular structure and function, ought to be made in the awareness that models are never only descriptive tools for knowledge representation, but are also prescriptive. Keller (1992, p. 5) observed:
Since representations are necessarily structured by language (hence, by culture), no representation can ever ‘correspond’ to reality. At the same time, some representations are clearly better (more effective) than others. In the absence of a copy of truth, we need to search for the meaning of ‘better’ in a comparison of the uses to which different representations can be put, that is, in the practices they facilitate. From such a perspective, scientific knowledge is value-laden (and inescapably so) just because it is shaped by our choices – first, of what to seek representations of, and second, of what to seek representations for. Far from being value-free, good science is science that effectively facilitates the material realization of particular goals, that does in fact enable us to change the world in particular ways.

Model choice, while unquestioningly having to satisfy cognitive values such as accuracy, consistency, simplicity, breadth of scope and fruitfulness (Longino 1996), also has an explicit ethical dimension that scientists have a responsibility to confront (Nagl 1998). A critical awareness of which kind of knowledge, and which kind of goals, are likely to be facilitated by the chosen mode of representation ought to inform any modeling project. I will argue that, when dealing with complex systems – be they molecules, human bodies or ecosystems – we have a duty to study and represent them as such. I will show that this duty derives from the classical bioethical principles of beneficence and nonmaleficence (Beauchamp & Childress 1989), and then will extent my argument by discussing aspects of Longino’s ‘theoretical virtues’ (Longino 1996, p. 44).

The Hippocratic oath expresses a duty of nonmaleficence, or not inflicting harm, together with a duty of beneficence, or doing good (Beauchamp & Childress 1989, p. 120). These duties are absolutely fundamental to biomedical ethics and the practice of medicine. In contemporary medicine, biomedical research scientists are doctors’ close partners, and thus it can be argued that the ethical prescriptions of the Hippocratic oath ought to extend to their branch of the life sciences. Scientists’ duties may be loosely phrased as follows: “I will pursue my scientific work to help the sick according to my ability and (best) judgement, but I will never use it to injure or harm them.”

In the light of the immeasurable potential benefits of artificially designed proteins in molecular medicine, it is greatly desirable that we develop a repertoire of design methods that would enable us to create proteins for therapeutic uses. Thus, one can assert that a duty to develop such techniques exists, and that this duty derives from the principle of beneficence. However, many problems still hamper the attainment of these goals. On the one hand, it has been recognized for some time that protein design can draw vital insights from evolutionary principles. On the other hand, the complex interplay of molecular organization and evolutionary dynamics is still poorly understood, and this lack of understanding presently limits potentially extremely fruitful evolutionary approaches to protein design. An awareness of these problems,
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together with the insight that proteins are complex adaptive systems – in other words, that they evolve as complex systems – leads to a duty to study their complex systems properties. Such a duty is grounded in the reasonable expectation that such a research program will enable the gain of new knowledge, and the development of new design techniques, that are inaccessible from within other conceptual frameworks. This duty therefore also follows from the principle of beneficence. The work on complex systems models of protein domain evolution presented in this paper was carried out in the hope that it will help elucidate how new functions evolve in stable folding architectures, and be a contribution towards overcoming the current limitations in protein design.

Whenever there are consequences to human welfare, the duty to treat complex systems as complex systems is also grounded in the principle of nonmaleficence, or not inflicting harm. This is quite immediately obvious in the case of large complex systems, such as ecosystems or whole biological organisms. An environmental risk assessment, or a drug trial, that employs models that are inadequate for detecting effects due to complex systems properties, may pose great risks to people. A duty to avoid the use of such models, and to develop alternatives that can model complex systems behavior, can be easily appreciated. It may, however, at first be less obvious how such a duty due to nonmaleficence could be postulated for models of much smaller complex systems, models of proteins for example.

Here, we need to briefly digress to consider that models are in a certain sense metaphorical constructions (Nagl 1998; Holland 1998, p. 207). As such, they carry with them not only explicit messages but also implicit content. A model is a device for seeing the world in a particular way. A well-developed scientific model accumulates a complicated assortment of techniques, interpretations, standards of proof, and so on; and may well have a cognitive impact far transcending the original context in which it was conceived. Much of this remains unwritten, but is understood by everyone who has been socialized within the research tradition associated with the model.

Importantly, models shape our habits of thought. It seems therefore unwise to think that, while we may feel an ethical obligation to develop models that embody the complexities of the human body, we may ‘get away’ with ignoring the complex systems properties of biomolecules. Our fundamental orientation toward life is always at issue, no matter what part of it we happen to focus on at the time. Habits of thought that prompt us to take heed of the complexity inherent in all biological entities, will direct our thinking away from seeing and representing such entities in simplistic terms – away from mechanistic conceptions of molecules or visions of machine-like bodies. They will hopefully also stop us from intervening in the human body from this fragmented, and potentially extremely harmful, perspective. It is within
this wider context that nonmaleficence can be seen as a guiding principle, supporting the duty to study the complex systems properties of biological entities. On the positive side, these novel habits of thought may direct us toward an understanding of the world as a multidimensional whole of complex systems within complex systems. Such a change in thinking may subsequently lead to new biomolecular therapies that seek to cooperate with, rather than control, living systems. The central role of biomolecules in molecular medicine, a tremendously powerful and influential new research field, may make complex systems models of these molecules instrumental in bringing about such a global change in scientific attitude.

This leads on to some final points I wish to make. My reflections on ethical issues of model choice find an echo in Longino’s work on theoretical virtues (Longino 1996, p. 44). These virtues complement the cognitive values of accuracy, consistency, simplicity, breadth of scope and fruitfulness, that are commonly applied to assess the merits of scientific models. Longino’s virtues of ‘novelty’ and ‘mutuality of interaction’ are especially pertinent to my concerns. Longino defines ‘novelty’ as models or theories that differ in significant ways from presently accepted ones by (i) attempting to elucidate phenomena that have not been previously studied, (ii) postulating different processes, (iii) adopting different principles of explanation, and (iv), incorporating alternative metaphors (p. 45). As Longino (1996) states, “treating novelty as a virtue reflects a deep skepticism that mainstream theoretical frameworks could be adequate to the problems confronting us” (p. 46). It is certainly from a great disquiet regarding the present state of our biomedical models that I argue for an urgent need for complex systems models, which can be seen to fulfil all four of Longino’s criteria of novelty. Finally, the virtue of ‘mutuality of interaction’ values theories and models that treat relationships between entities and processes as mutual, avoid causal explanations based on single factors, and take complex interaction as a fundamental principle of explanation (Longono 1996, p. 47). Clearly, complex systems models are embodiments of this virtue par excellence. In conclusion, complex systems approaches, informed by the theoretical virtues of ‘novelty’ and ‘mutuality of interaction’, are highly relevant to current biomedical research, if one is to fulfil the duties of the Hippocratic oath.

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