

Proteome Res. Author manuscript; available in PMC 2008 September 19.

Published in final edited form as:

J Proteome Res. 2007 May; 6(5): 1882–1898. doi:10.1021/pr060392u.

Functional Anthology of Intrinsic Disorder. I. Biological Processes and Functions of Proteins with Long Disordered Regions

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Abstract

Identifying relationships between function, amino acid sequence and protein structure represents a major challenge. In this study we propose a bioinformatics approach that identifies functional keywords in the Swiss-Prot database that correlate with intrinsic disorder. A statistical evaluation is employed to rank the significance of these correlations. Protein sequence data redundancy and the relationship between protein length and protein structure were taken into consideration to ensure the quality of the statistical inferences. Over 200,000 proteins from Swiss-Prot database were analyzed using this approach. The predictions of intrinsic disorder were carried out using PONDR VL3E predictor of long disordered regions that achieves an accuracy of above 86%. Overall, out of the 710 Swiss-Prot functional keywords that were each associated with at least 20 proteins, 238 were found to be strongly positively correlated with predicted long intrinsically disordered regions, whereas 302 were strongly negatively correlated with such regions. The remaining 170 keywords were ambiguous without strong positive or negative correlation with the disorder predictions. These functions cover a large variety of biological activities and imply that disordered regions are characterized by a wide functional repertoire. Our results agree well with literature findings, as we were able to find at least one illustrative example of functional disorder or order shown experimentally for the vast majority of keywords showing the strongest positive or negative correlation with intrinsic disorder. This work opens a series of three papers, which enriches the current view of protein structure-function relationships, especially with regards to functionalities of intrinsically disordered proteins and provides researchers with a novel tool that could be used to improve the understanding of the relationships between protein structure and function. The first paper of the series describes our statistical approach, outlines the major findings and provides illustrative examples of biological processes and functions positively and negatively correlated with intrinsic disorder.

Keywords

Intrinsic disorder; protein structure; protein function; intrinsically disordered proteins; bioinformatics; disorder prediction

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Introduction

Among other objectives, computational biology aims to enable an understanding of the relationships between the primary sequence, the higher order structure and the function of proteins. Each protein function is generally thought to originate from a specific 3-dimensional (3-D) structure. Formulation of this view began more than 100 years ago with the lock-and-key model proposed by Fischer. More than 70 years ago Wu, and slightly later, Mirsky and Pauling equated denaturation with loss of specific structure. The dependence of function on 3-D structure was accepted by the time of the protein folding studies of Anfinsen and colleagues. The flood of protein 3-D structures determined by X-ray diffraction and by nuclear magnetic resonance (NMR) spectroscopy has overwhelmed alternative concepts.

In contrast to the dominant view given above, proteins for which intrinsic disorder is required for function have been reported in the literature for many years. By "intrinsic disorder" we mean that the protein (or protein region) exists as a structural ensemble, either at the secondary or at the tertiary level. Thus, both extended regions with perhaps some elements of secondary structure and collapsed (molten globule-like) domains with poorly packed side chains are included in our view of intrinsic disorder. 6 More detailed analysis of extended disordered proteins/regions revealed that they can be further divided in two groups, random coil-like and pre-molten globule-like conformations. ⁷ Recently, more than 150 proteins have been identified as containing functional disordered regions, or being completely disordered, yet performing vital cellular roles. ^{8, 9} Twenty-eight separate functions were assigned to these disordered regions, including molecular recognition via binding to other proteins, or to nucleic acids.⁸, ¹⁰ A complementary view is that functional disorder fits into at least five broad classes based on the mode of disordered protein/region action. ¹⁰ Obviously, for these proteins, the predominant structure-function paradigm is insufficient, which suggests that a more comprehensive view is needed. ¹¹ In fact, a new paradigm was recently offered to elaborate the sequence-to-structure-to-function scheme in a way that includes the novel functions of disordered proteins. 6, 7, 12 The complex data supporting this revised view were summarized in "The Protein Trinity" hypothesis, which suggested that native proteins can exist in one of three states, the solid-like ordered state, the liquid-like collapsed-disordered state or the gaslike extended-disordered state. 12 Function is then viewed to arise from any one of the three states or from transitions between them. Later this paradigm was extended to "The Protein Quartet" model to include one more extended-disordered conformation, the pre-molten globule state. For structured proteins: i.e., proteins that form crystals without partners or have ordered globular forms without partners in NMR experiments, we will use the terms "structured", "intrinsically ordered" or just ordered.

Recent studies revealed that many proteins lack rigid 3-D structure under physiological conditions *in vitro*, existing instead as highly dynamic ensembles of interconverting structures. Indeed, the literature on these proteins, known as intrinsically disordered, natively unfolded, or intrinsically unstructured, has virtually exploded during the last decade. ^{7, 13} This literature explosion is consistent with bioinformatics studies predicting that about 25 to 30% of eukaryotic proteins are mostly disordered, ¹⁴ that more than half of eukaryotic proteins have long regions of disorder, ^{14, 15} and that more than 70% of signaling proteins and the vast majority of cancer-associated proteins have long disordered regions. ¹⁶ As it has been already mentioned, despite the fact that intrinsically disordered proteins fail to form fixed 3-D structures by themselves under physiological conditions, they carry out numerous important biological functions. ^{6–11, 13, 16–22} Intrinsically disordered regions are typically involved in regulation, signaling and control pathways in which interactions with multiple partners, and high-specificity/low-affinity interactions are often involved. ^{21–23} Furthermore, sites of posttranslational modifications (acetylation, hydroxylation, ubiquitination, methylation, phosphorylation, etc.) and proteolytic attack are frequently associated with regions of intrinsic

disorder. ^{6, 16} Given the high frequency of intrinsically disordered proteins and their crucially important functions, a curated Database of Disordered Protein (DisProt) has been recently initiated. ²⁴ This database provides structure and function information about proteins that lack a fixed 3D-structure under putatively native conditions, either in their entireties or in part. ²⁴ In spite of all of this, the phenomenon of intrinsic disorder in proteins is still severely underappreciated; not a single biochemistry textbook discusses these proteins. ²⁵

There is a large gap between the number of proteins with experimentally confirmed disordered regions and actual number of such proteins in nature. Although studies of functional properties of known disordered proteins are helpful in revealing the functional diversity of protein disorder, they are bound to provide only a limited view. In this study, we propose a statistical approach for comprehensive study of functional roles of protein disorder. This approach relies on use of the VL3E²⁶ predictor that is currently the most accurate predictor of long disordered regions with estimated accuracy of above 86%. ²⁶ The high accuracy of VL3E ensures that most disordered regions could be successfully detected with only a small fraction of ordered regions being incorrectly labeled as disordered. The VL3E predictor was applied to over 200,000 Swiss-Prot²⁷ proteins, many of which were annotated with one or more functional keywords. Then, the disorder-and order-correlated functions were detected as those that are overrepresented by proteins predicted to have long disordered regions (> 40 amino acid residues) in comparison with a random selection of proteins with the same length distribution. The proposed approach ensures that adverse effects of sequence redundancy and sequence length are eliminated. Disorder predictors were previously used to analyze functions of disordered proteins. For example, it was shown that a large fraction of cancer-related proteins are likely to be disordered. ¹⁶ In another study ²⁸ it was demonstrated that many processes in yeast are related to protein disorder. The current study provides a comprehensive analysis of disorder-related functions by using a much larger set of proteins (i.e., the entire Swiss-Prot database).

Given a list of functions positively and negatively correlated with disorder, we performed an extensive literature survey to find experimental evidence supporting the findings. We were able to find at least one illustrative experimentally validated example of functional disorder/ order for a large majority of functional keywords. This work opens a series of three papers dedicated to finding and description of protein functions and activities that are positively and negatively correlated with long disordered regions. Being the first in the series, this paper deals with the description of the statistical approach used here and delineates the major results of the application of this tool for the analysis of over 200,000 proteins from Swiss-Prot database. This paper also provides illustrative literature examples related to the Swiss-Prot keywords associated with the biological processes and functions positively and negatively correlated with intrinsic disorder. The second paper of the series portrays keywords related to the cellular components, domains, technical terms, developmental processes and coding sequence diversity associated with long disordered regions, ²⁹ whereas keywords correlated with ligands, postranslational modifications and diseases associated with long disordered regions are the topic for the last paper of the series. 30 The overall result is that this series of papers represents a functional anthology of intrinsic disorder that includes both the results of our bioinformatics analysis and illustrative literature examples for the majority of functional keywords possessing strongest positive or negative correlation with the intrinsic disorder prediction.

Materials and methods

Dataset

The dataset for analysis was constructed using the Swiss-Prot database (release 48, 2005) containing 201,560 proteins. ²⁷ In this study we used the 196,326 proteins with length longer than 40 amino acid residues. Each protein in Swiss-Prot is annotated with keywords that

describe its functional or structural properties. Out of the 874 keywords used by Swiss-Prot, 710 were associated with at least 20 proteins. Swiss-Prot is statistically redundant, as it contains a large number of homologous proteins with highly similar sequences. I Ignoring the redundancy would significantly bias statistical inference. To reduce redundancy, TribeMCL was applied to cluster the protein sequences from Swiss-Prot into families. TribeMCL uses the Markov clustering algorithm for the assignment of proteins into families based on the similarity matrix generated from the all-against-all BLASTp³³ comparison of sequences. It is able to produce high quality families despite presence of multi-domain proteins, peptide fragments, and promiscuous domains. The obtained BLAST profiles were imported into TribeMCL software package (http://micans.org/mcl/) and clustering was performed with all parameters set at default. As a result of application of this redundancy reduction procedure, the sequences were grouped into 27,217 families.

Predicting long disordered regions in proteins

Previous studies suggested that in comparison with ordered sequences, disordered sequences tend to have lower aromatic content, higher net charge, 17 , $^{34-36}$ higher values of the flexibility indices, greater hydropathy values, 34 , 36 and lower sequence complexity. 37 Following these observations, the VL3E predictor 26 was developed using 162 long (>30 residues) disordered regions from non-redundant set of 152 DisProt proteins 24 , 38 and 290 completely ordered proteins. The predictor consists of an ensemble of neural network classifiers and it achieves ~87% cross-validation accuracy on balanced data with equal number of ordered and disordered residues. We used the VL3E predictor to predict Swiss-Prot proteins with long disordered regions. Each of the 196 ,326 Swiss-Prot proteins was labeled as putatively disordered if it contained a predicted intrinsically disordered region with $^{>40}$ consecutive amino acids and as putatively ordered otherwise. For notational convenience, we introduce disorder operator d such that $^{d}(s_i) = 1$ if sequence s_i is putatively disordered, and $^{d}(s_i) = 0$ if it is putatively ordered.

Relationship between long disorder prediction and protein length

The likelihood of labeling a protein as putatively disordered increases with its length. To account for this length dependency, we estimated the probability, P_L , that VL3E predicts a disordered region longer than 40 consecutive amino acids in a SwissProt protein sequence of length L. Probability P_L was determined by partitioning all SwissProt proteins into groups based on their length. To reduce the effects of sequence redundancy, each sequence was weighted as the inverse of its family size; if sequence s_i was assigned to TribeMCL cluster $c(s_i)$, we calculated n_i as the total number of SwissProt sequences assigned to this cluster and set its weight to $w(s_i) = 1/n_i$. In this manner, each cluster is given the same influence in estimation of P_L , regardless of its size. To estimate P_L , all SwissProt sequences with length between L-l and L+l were grouped in set $S_L = \{s_i, L-l \le |s_i| \le L+l\}$. The probability P_L was estimated as

$$P_{L} = \frac{\sum_{s_{i} \in S_{L}} w(s_{i})d(s_{i})}{\sum_{s_{i} \in S_{L}} w(s_{i})}.$$

Window size l allowed us to control the smoothness of P_L function. In this study we used window size equal to 20% of the sequence length, $l = 0.1 \cdot L$. We show the resulting curve in Figure 1 together with the same results when l = 0.

Extracting disorder-and order-related Swiss-Prot keywords

For each of the 710 SwissProt keywords occurring in more than 20 SwissProt proteins, we set to determine if it is enriched in putatively disordered or ordered proteins. For a keyword KW_i , j = 1...710, we first grouped all SwissProt proteins annotated with the keyword to S_i . To

take into consideration sequence redundancy, each sequence $s_i \in S_j$ was weighted based on the SwissProt TribeMCL clusters. If sequence s_i was assigned to cluster $c(s_i)$, we calculated n_{ij} as the total number of sequences from S_j that belonged to that cluster and set its weight to $w_j(i) = 1/n_{ij}$. Then, the fraction of putatively disordered proteins from S_j was calculated as

$$F_{j} = \frac{\sum_{s_{i} \in S_{j}} w_{j}(s_{i})d(s_{i})}{\sum_{s_{i} \in S_{j}} w_{j}(s_{i})}.$$

The question is how well this fraction fits the null model that is based on the length distribution P_L . Let us define random variable Y_i as

$$Y_{j} = \frac{\sum_{s_{i} \in S_{j}} w_{j}(s_{i}) X_{|s_{i}|}}{\sum_{s_{i} \in S_{L}} w_{j}(s_{i})},$$

where X_L is a Bernoulli random variable with $P(X_L = 1) = 1 - P(X_L = 0) = P_L$. In other words, Y_j represents a distribution of fraction of putative disorder among randomly chosen SwissProt sequences with the same length distribution as those annotated with KW_j .

If F_j is in the left tail of the Y_j distribution (i.e. the p-value $P(Y_j > F_j)$ is near 1), the keyword is enriched in ordered sequences, while if it is in the right tail (i.e. the p-value $P(Y_j > F_j)$ is near 0) it is enriched in disordered sequences. We denote all keywords with p-value < 0.05 as disorder-related and those with p-value > 0.95 as order-related.

The distribution Y_j is hard to derive analytically, so we randomly generated 1,000 realizations and calculated the empirical p-value as the fraction of times these realizations were larger than F_j . We also calculated the mean μ_j and standard deviation σ_j of the 1,000 realizations. We observed that, when $|KW_j|$ is large, distribution of Y_j resembles a Gaussian distribution with mean μ_j and standard deviation σ_j . Using the Gaussian approximation, we calculated the Z-score of KW_j as $(F_j - \mu_j)/\sigma_j$ and its p-value as $1/2(1 - erf(Z_j/2))$, where erf() is the error function. The Gaussian approximation is useful since using the fraction of 1,000 replicates is not accurate in estimating p-values below 0.01 or above 0.99. We report the Z-scores together with the empirical p-values in the results.

Results

Estimating correlation between long disordered regions and Swiss-Prot keywords

We applied the procedure described above to each of the 710 Swiss-Prot keywords occurring each in more than 20 Swiss-Prot proteins. These 710 keywords can be grouped into 11 functional categories, which are listed in Table 1. We denote keywords with p-value > 0.95 as disorder-related and the ones with p-value < 0.05 as order-related. Keywords with p-value between 0.95 and 0.05 are ambiguous. These functions might depend on structured of disordered regions but simply exhibit signals that are too weak. Alternatively these functions might depend on short regions of disorder or might require both ordered and disordered regions.

The number of keywords strongly correlated with disorder and order is significantly larger than expected by the random model. This is evident by observing that, for a p-value threshold of 0.05, a random predictor would result in about 5% (~36) of order and 5% of disorder-related keywords. These results suggest that presence or absence of disordered regions is an important factor in majority of biological functions and processes. Overall, this analysis shows that 238 Swiss-Prot functional keywords are disorder-related, whereas 302 are order-related. Interestingly, only two of the categories, "Biological Process" and "Ligand", are enriched in

order-related keywords, while the remaining 9 are enriched in the disorder-related keywords. This result supports an earlier conjecture that disordered regions have a larger functional repertoire than the ordered regions. 20

To further understand these function-disorder relationships, we carried out manual literature mining and studied a large number of individual experimental examples. To organize the presentation of these results, the keywords from various functional categories, which are most significantly associated with protein order and disorder arranged into specific groups (Table 2–Table 6). In each table, the disorder-function relationships are ranged by their Z-scores (see Materials and Methods). The Z-scores for all 710 functions are given in Supplementary Materials (see Table S1). One of the major goals here was to determine for each example whether the indicated function was carried out by regions of disorder or regions of structure. After all, the keyword-disorder correlations established by the method of Figure 2 do not determine whether the indicated association implies direct involvement of disorder with function or not.

Biological processes associated with intrinsically disordered proteins

The set of top 20 Swiss-Prot annotated cellular processes associated with predicted disorder are listed in Table 2. Presented below are several illustrative examples of biological processes from this list for which the associations with long disordered regions have been experimentally determined. The data below are organized in the following way: each discussed keyword is placed at the beginning of the corresponding paragraph and *Italized*. If the following description involves other keywords discussed in this and the subsequent papers ²⁹, ³⁰ these keywords are presented using the *Italic* font.

Differentiation—In developmental biology, cellular differentiation describes the process by which different cell types are derived from a single fertilized egg cell. Differentiation is a continuously regulated process, with specific interactions between the cell and its environment playing a major role in maintaining stable expression of differentiation-specific genes.³⁹ Obviously, numerous intracellular and extracellular proteins are involved in the differentiation control and regulation. For example, extracellular matrix (ECM), which is an important component of the cellular environment, was shown to play a role in regulating differentiation and the differentiated phenotype of cells. ⁴⁰, ⁴¹ An ECM is present within mammalian embryos from the two-cell stage and is a component of the environment of all cell types, although the composition of the ECM and the spatial relationships between cells and ECM differ between tissues. The ECM offers structural support for cells, and can also act as a physical barrier or selective filter to soluble molecules. ⁴⁰ The ECM is composed of glycoproteins, glycosaminoglycans and proteoglycans that are secreted and assembled locally into an organized network to which cells adhere. 42 Cells interact with the ECM via numerous cellbinding sites located within individual ECM glycoproteins and ECM receptors. For example, in case of fibronectin the primary determinant of cell-binding activity for many cell types resides in the sequence GRGDSP, which occurs in one of the type III repeats that form the central domain of the molecule. 43 Intriguingly, this cell-binding sequence exclusively consists of strongly disorder-promoting amino acid residues,³⁷ thus it very likely is intrinsically disordered. Importantly, it has been experimentally shown that the fibronectin binding domains from several different species of Gram-positive bacteria 44 as well as the N-terminal domain of BBK32, a fibronectin-binding lipoprotein from Borrelia burgdorferi, the causative agent of Lyme disease, are all intrinsically disordered.⁴⁵

Transcription and transcription regulation—Transcription, being one of the key processes in the living cell through which a DNA sequence is enzymatically copied to produce a complementary RNA, is the transfer of genetic information from DNA into RNA. In the case

of protein-encoding DNA, transcription initiates the process that ultimately leads to the translation of the genetic code into a functional protein. Transcription is strongly regulated by a number of proteins, especially transcription factors that include activators, repressors and enhancer-binding factors. Transcription factors function through the recognition of specific DNA sequences and the recruitment and assembly of the transcription machinery. Thus, both protein-DNA and protein-protein recognition are central processes in function of transcription factors. Several examples of intrinsically disordered proteins in transcriptional regulation have been reported. ^{18, 19} For example, the C-terminal activation domain of the bZIP protooncoprotein c-Fos unstructured and highly mobile, yet this protein effectively suppresses transcription in vitro. 46 The C-terminal domain of the transcriptional corepressor CtBP, which serves as a scaffold in the formation of a multiprotein complex hosting the essential components of both gene targeting and coordinated histone modifications, is also intrinsically disordered, as determined by using several complementary approaches (bioinformatics, NMR, CD, and small-angle X-ray scattering).⁴⁷ Recent analysis of high-resolution structures of transcription factors in the Protein Data Bank revealed that these proteins are, on average, largely disordered molecules with over 60% of amino acids residing in 'coiled' configurations. ⁴⁸ The abundance of intrinsic disorder in transcriptional regulation was further demonstrated using a set of bioinformatics tools, including the Predictor Of Natural Disorder Regions (PONDR). This analysis showed that up to 94% of transcription factors have extended regions of intrinsic disorder. Furthermore, the analysis of the disorder distribution within the transcription factor datasets revealed that the degree of disorder is significantly higher in eukaryotic transcription factors than in prokaryotic transcription factors. ⁴⁹ The complementary analysis of human transcriptional regulation factors revealed that although their average sequence is more than twice as long as that of prokaryotic proteins, the fraction of human sequences aligned to domains of known structure in PDB is less than half of that found for bacterial transcription factors, ⁵⁰ suggesting that the increased length of eukaryotic transcription factors results to a significant degree from the addition of disordered regions.

Spermatogenesis—Spermatogenesis is the formation and development of mature spermatozoa from stem cells by *meiosis* and *spermiogenesis*. As *spermatogenesis* progresses, there is a widespread reorganization of the haploid genome followed by the extensive DNA condensation suggesting that the dynamic composition of chromatin is crucial for the activities of enzymes that act upon it. Histone variants such as H3.3, H2AX, and macroH2A play important roles at the various stages of spermiogenesis. Furthermore, posttranslational modifications of different histones, including specifically modulated acetylation of histone H4 (acH4), ubiquitination of histones H2A and H2B (uH2A, uH2B), and phosphorylation of histone H3 (H3p), are also involved in the regulation *spermatogenesis*. ⁵¹ Furthermore, during the final stages of spermatogenesis, the DNA of sperm in most organisms is compacted due to the replacement of somatic-type histones by DNA-condensing sperm nuclear basic proteins (SNBPs), sperm histones (H type), protamine-like (PL type), and protamines (P type). 52 Analysis of amino acid composition of PL-I sperm nuclear protein from Spisula solidissima revealed that it contains high amounts of lysine and arginine (24.8 and 23.1%, respectively). ⁵³ Also, the PL-I has been shown to possess a tripartite structure, consisting of N- and Cterminal flexible "tails" flanking a globular, trypsin-resistant core of 75 amino acids. 54–56

DNA condensation—DNA condensation *in vivo* relies on electrostatics-driven interaction of DNA with small cations and/or a number of abundant proteins including histones. In eukaryotes, the basic unit of chromatin (a condensed form of DNA) is commonly defined as a nucleosome, which is made up of DNA wrapped in two left-handed superhelical turns around a proteinaceous core. ⁵⁷ The nucleosome core contains eight histone proteins, two dimers of H2A–H2B that serve as molecular caps for the central (H3–H4)₂ tetramer. ⁵⁸ Thus, nucleosome represents the first level of chromatin condensation and is often termed 'beads on a string'.

Other crucial components of chromatine are the linker (H1 family) histones, which bind to the DNA that enters and exits the nucleosome and which facilitate the shift in equilibrium of chromatin towards more condensed, higher order forms.⁵⁷ It was established long ago that purified core histones being dissolved in water with no added salt, behave as polypeptides in an "extended loose form". 58–63 Recently, using a combination of bioinformatics tools with several biophysical techniques it has been shown that in low salt all bovine core histones are typical natively unfolded proteins; i.e., they possess exceptionally high level of intrinsic disorder. ⁶⁴ Importantly, in the presence of high enough salt concentrations, core histones adopt a folded conformation. 58–64 In the crystal structure, histones are highly helical proteins, with α -helices accounting for 65–70% of the total structure. Only 3% of residues can be assigned to short parallel β -sheets, the remainder, approximately 30%, is not ordered. 65, 66 It has been also emphasized that the N-terminal "tail" domains (NTDs)⁶⁷ of the core histones and the Cterminal tail domain (CTD) of linker histones are intrinsically disordered, yet they are able to bind to many different macromolecular partners in chromatin. ⁶⁸ Particularly, histone tails are known to be involved in the conformational changes of the nucleosome core particle (NCP) as well as in the structural phase transitions occurring at the supramolecular level. It is generally accepted that these tails interact with DNA at low salt and are extended outside of the particle at salt concentrations above ~0.2 M monovalent salt.⁶⁹ Analysis of the extension process of isolated NCP tails as a function of ionic strength has been reported. The addition of salt simultaneously screens Coulombic repulsive interactions between NCP and Coulombic attractive interactions between tails and DNA inside the NCP. 70

Cell cycle, cell division, mitosis, meiosis—The cell cycle consists of an ordered series of events between the two cell divisions and involve the growth, replication, and division of a eukaryotic cell. Depending on the type of cell, the cell division might result in two different outcomes: in the division of somatic cells (mitosis), daughter cells are identical to the parent cell and contain a complete copy of the parental chromosomes; in meiosis (the division of sex cells), the daughter cells contain a half of the genes of the parent. Progression through the cell cycle is controlled in part by the activity of cyclin-dependent kinases, which are considered to be the major timekeepers of cell division. ⁷¹ Cdks are regulated by binding to their *cyclin* protein partners thus forming active heterodimeric complexes. Eight Cdk family members (Cdk1–Cdk8) and nine cyclins (A–I) have been identified so far. Interestingly, each Cdk pairs with a separate cyclin class, most of which have at least two members. ^{72, 73} For example, Cdk1 together with cyclin B1 directs the G2/M transition. Exit from G1, in contrast, is primarily under the control of cyclin D/Cdk4/6. Finally, two other cyclins (A and E) that pair with Cdk2 are required for the G1/S transition and progression through the S phase. ⁷², ⁷³ The activity of Cdks throughout the cell cycle is known to be precisely regulated by a combination of several mechanisms, including the control of cycle-dependent variations in the levels of activating partners, cyclins; coordination of Cdk phosphorylation and dephosphorylation; and variations in the levels of the Cdk inhibitor proteins, CKIs, which are responsible for the deactivation of the Cdk–cyclin complexes. ^{71, 74} Five major mammalian CKIs are known: p21^{Waf1/Cip1/Sdi1} and p27Kip1 inactivate Cdk2 and Cdk4 cyclin complexes by binding to them, p16INK4 and p15^{INK4B} are specific for Cdk4 and Cdk6, whereas p57^{Kip2} is specific for Cdk2.⁷¹, ⁷⁴ The p21^{Waf1/Cip1/Sdi1},⁷⁵ p27^{Kip1},^{76–78} and p57^{Kip2} CKIs⁷⁹ are all intrinsically disordered proteins that undergo sequential folding upon binding to their functional partners.

mRNA processing and splicing—An average gene in higher eukaryotes is very large due to the interruption of the coding sequence with large noncoding introns. Introns are known to be co-transcriptionally removed with great accuracy by pre-messenger RNA (mRNA) splicing. A large number of proteins are involved in generating specificity in pre-mRNA processing. Among the different pre-mRNA processing possibilities, alternative splicing is the most prevalent mechanism to generate proteomic diversity. The role of intrinsic disorder

in alternative splicing is discussed in the second paper of this series. ²⁹ Astounding examples of extensively alternatively spliced genes includes the Down syndrome cell adhesion molecule gene (Dscam) from *Drosophila*, the Neurexin and CD44 genes in humans, which can produce as many as about 38,000, 3000 and 1000 different splice forms, respectively. ^{80–82}. *Splicing* involves the stepwise assembly of five (U1, U2, U4, U5 and U6) small *ribonucleoprotein* particles (snRNPs) and a large number of proteins onto the pre-mRNA to form a large complex called the *spliceosome*. ⁸³ The role of intrinsic disorder in the *spliceosome* function is discussed below in the Section entitled *Cellular components associated with intrinsic disorder*.

Apoptosis—Apoptosis is the programmed death of a cell. Regulation and control of apoptosis is crucial for the normal functioning of the organism. On the other hand, cancer cells avoid apoptosis and continue to multiply in an unregulated manner. The tumor suppressor protein p53 represents an outstanding example of this concept. The p53 molecule regulates expression of genes involved in numerous cellular processes, including cell cycle progression, apoptosis induction, DNA repair, as well as others involved in responding to cellular stress. 84 When p53 function is lost, either directly through mutation or indirectly through several other mechanisms, the cell often undergoes cancerous transformation. 85, 86 Cancers showing mutations in p53 are found in colon, lung, esophagus, breast, liver, brain, reticuloendothelial tissues and hemopoietic tissues. 85 When activated, p53 accumulates in the nucleus and binds to specific DNA sequences. 86, 87 It has been shown to induce or inhibit over 150 genes, including p21, GADD45, MDM2, IGFBP3, and BAX. 87 The p53 protein can be divided into three functional domains: an amino-terminal transactivation region, a central DNA binding domain, and a carboxy-terminal tetramerization and regulatory region. At the physiological temperature of 37°C and in the absence of modifications or stabilizing partners, wild-type p53 is more than 50% unfolded. 88 According to NMR analysis, the isolated transactivation domain lacks rigid structure, 89, 90 although it does possess an amphipathic helix that forms secondary structure part of the time, which can be stabilized by binding to Mdm2⁹¹ or in the membrane environment. 92 Besides Mdm2, the transactivation domain interacts with numerous other proteins including TFIID, TFIIH, RPA, CBP/p300 and CSN5/Jab1, 84 thus playing a crucial role in the regulation of p53 function. For example, p53 can be inhibited by interaction with E3 ubiquitin ligase Mdm2, 93 which is bound to a short stretch of p53, specifically to residues 13–29. 91 As this region of p53 is within the transactivation domain, p53 cannot activate or inhibit other genes when Mdm2 is bound. Thus, p53-Mdm2 interaction exemplifies a crucial mechanism of vital regulation via the binding-induced folding of one of the interacting partners. The C-terminal regulatory domain is also unstructured. ^{94, 95} The structures of the core domain bound to DNA ⁹⁶ and of several oncogenic mutants have been solved. ⁹⁷ The crystal and NMR structures of the tetramerization domain are also known. ⁸⁹, ⁹⁸, ⁹⁹ The high disorder content of p53 may help to explain its inherent instability and extreme oncogenic potential. 100, 101

The BH3-only proteins belong to the proapoptotic family of proteins that function as key initiators of the programmed cell death. The BH3-only members of the Bcl-2 family proteins (those with a single Bcl-2 homology (BH) domain), including Bim, Bid, Bmf and Bad, are key initiators of *apotosis*, and they interact specifically with numerous binding partners. 102 In a healthy cell BH3-only molecules are either repressed, or are present in an inactive state. 103 The molecular mechanism of apoptosis initiation involves a stage of BH3-only proteins activation by a death stimulus, followed by their interaction and inactivation of prosurvival Bcl-2 proteins (such as CED-9 in *Caenorhabdhitis elegans*, and Bcl-x_L in humans). 102 Until recently, the structural knowledge about BH3-only proteins has been limited to peptide fragments bound to their targets. For example, in the complex of Bcl-x_L and a 33 residues long peptide corresponding to the BH3 domain of Bim, the peptide forms an α -helix upon binding to the hydrophobic groove of Bcl-x_L. 104 Analysis of crystal structures of BH3 domain peptides bound to the prosurvival proteins CED-9 and Bcl-x_L revealed that the nature of this interaction is highly conserved despite only a low level of shared sequence identity, with the conserved

leucine and aspartic acid residues of the LXXXGDE motif, which defines BH3-only proteins, making critical contacts with conserved residues in the hydrophobic binding groove of CED-9 and Bcl-x_L. $^{104-107}$ However, in another structure a 9-residue peptide of Bim forms a β -strand upon binding to the component of dynein motor complex, DLC1. 108 Recently, using methods such as CD, NMR, analytical centrifugation, size exclusion chromatography and limited proteolysis, it has been established that the BH3-only proteins Bim, Bad and Bmf are unstructured in the absence of binding partners. 109 Intriguingly, the majority of the Bim residues remains disordered when this protein binds and inactivates prosurvival proteins, with the only the short α -helical molecular recognition element 110 becoming structured. 109 Furthermore, detailed sequence analyses suggest that most BH3-only proteins are unstructured. 109 The disorder of this proapoptotic protein family is likely to be important for several biological functions such as promiscuous binding, extensive splicing, and regulation via phosphorylation.

Ubl conjugation pathway—Posttranslational modification via the covalent attachment of ubiquitin and different ubiquitin-like proteins (Ubls, including SUMO, ISG15, Nedd8, and Atg8) is a crucial regulatory cellular mechanism, which plays a number of important roles in controlling cell division, signal transduction, embryonic development, endocytic trafficking and the *immune response*. 111 For example, conjugation of *ubiquitin-like proteins* (the *Ubl* conjugation pathway) to components of the transcriptiol machinery is an important regulatory mechanism allowing switching between different activity states. While ubiquitination of transcription factors is associated with transcriptional activation, their SUMOylation is most often connected with transcriptional repression. 112 Recent bioinformatic analysis of a limited number of known ubiquitination substrates showed that protein ubiquitination sites are preferentially located within surface exposed, flexible loop regions. 113 In addition. the sequence analysis of ubiquitination sites and regions adjacent to them showed that their properties such as low sequence complexity, high negative net charge and low hydrophobicity, are similar to those of intrinsically disordered regions (Iakoucheva and Radivojac, personal communication). An example of SUMOylation occurring within intrinsically unstructured region is the conjugation of transcription factor Ets-1 with SUMO-1.¹¹⁴ Using NMR spectroscopy it has been shown that the sumoylation motif of Ets-1 containing Lys15 is located within the unstructured N-terminal segment of Ets-1 preceding its PNT domain. 114 The authors hypothesize that flexibility of the linking polypeptide sequence may be a general feature contributing to the recognition of SUMO-modified proteins by their downstream effectors. 114

Wnt signaling pathway—Wnt is a critical pathway for embryogenesis, carcinogenesis, and cancer stem cells. 115, 116 Detailed information on this pathway can be found on the Wnt Homepage (http://www.stanford.edu/~rnusse/wntwindow.html). The Wnt pathway shows evolutionary conservation across a wide range of species, ranging from the freshwater polyp *Hydra* to vertebrates. 117 Mammals have 19 Wnt genes that can be grouped into 12 subfamilies. 118 Surprisingly, at least 11 of these subfamilies are present in Cnidaria (specifically, the sea anemone *Nematostella vectensis*) suggesting that they are not the result of any recent evolutionary diversification. 119 This indicates that the acquisition of the Wnt subfamilies was an early development in the evolution of metazoa and likely occurred about 650 million years ago. 119, 120

The best-understood Wnt pathway is often called the Wnt/β -catenin pathway, in which the Wnt signal leads to activation of the nuclear functions of β -catenin. These functions activate expression of a number of genes leading to cell survival, proliferation, or differentiation. A second vertebrate Wnt pathway, the Wnt/Ca^{2+} pathway, promotes intracellular Ca^{2+} release and regulates cell movements in development and in some cancers. A number of Wnt protein isoforms are generated by *alternative splicing*. 123-125

Wnt proteins comprise a large family of highly conserved secreted *growth factors* that activate target-gene expression in both a short- and long-range manner and regulate cell-to-cell interactions during embryogenesis. Wnt signaling is involved in virtually every aspect of embryonic development and also controls homeostatic self-renewal in a number of adult tissues. 115, 117

Glycogen synthase kinase 3β (GSK3 β) is a Ser/Thr protein kinase, which is one of the major players in the Wnt signaling pathway as GSK3 β hyperphosphorylates β -catenin, thus promoting its ubiquitination and targeted destruction. ¹²⁶ The crystal structure of human GSK3 β (420 residues) has been solved at 2.8 Å. ¹²⁷ Clear electron density was only evident for the 351 residues from Lys35 to Ser386, with the segments of the polypeptide preceding Lys35 and following Ser386 being disordered in the crystal. ¹²⁷ The structure of the ordered part of GSK3 β agrees with the consensus observed for "activation-segment" protein kinases, consisting of an N-terminal β -sheet domain, coupled to a C-terminal α -helical domain. The visible N-terminal domain (35–134) consists of a seven-stranded β -sheet, which folds to a closed orthogonal β -barrel. The core of the C-terminal α -helical domain (152–342) has a similar topology to the equivalent region in such mitogen activated protein kinases, such as MAPK, as ERK2 and p38. ¹²⁷ It is important to emphasize that the major difference between the C-terminal α -helical domain of GSK3 β and MAPK is the absence of the second helix in the hairpin segment from 276–293 in the GSK3 β domain. Furthermore, in GSK3 β this region represents a highly mobile and poorly defined 285–299 loop. ¹²⁷

Neurogenesis—Numerous proteins are involved in *neurogenesis*, the formation and development of nervous tissue. Among these proteins are the *transcription factors* Pax3, ¹²⁸ Pax6, ¹²⁹ Glis2, ¹³⁰ and Erm, ¹³¹ which play an important regulatory role in this process. These transcription factors, like transcription factors in general, are highly disordered. For example, Pax3 has a highly flexible linker (53 amino acids) separating two DNA binding domains: a paired domain (128 amino acids) and a paired type homeodomain (60 amino acids). ¹³² Similar to Pax3, transcription factor PAX6 has two DNA-binding domains, a paired domain and a homeodomain (HD), joined by a glycine-rich linker and followed by a proline-serine-threonine-rich (PST) transactivation region at the C terminus. ¹³³ Structural analysis revealed that the central 250 amino acid residues of the transcription factor Erm has very little (if any) ordered structure. ¹³⁴

Chromosome partition—Chromosome partition in two daughter cells is a complex process that involves a number of proteins. For example, proteins such as topoisomerase IV and XerCD recombinase, as well as MukB and FtsZ are related to chromosome partition in *Escherichia coli*. ¹³⁵ MukB exists as two thin rods (long antiparallel coiled coils) with globular domains at the ends emerging from the very flexible (read disordered) linker domain (123 amino acids). ¹³⁶ The flexibility of the hinge is crucial for the MukB function, as the arms can open up to 180°, separating the terminal domains by 100 nm, or close to near 0°, bringing the terminal globular domains together. ¹³⁶

Immune response—The immune system is capable of generating specific antibodies against an almost infinite diversity of physiological or synthetic antigens. However, the repertoire of antibodies produced in any organism is fixed, suggesting that the immune system is an example of nearly unlimited functional multiplicity based on limited sequence diversity. ¹³⁷ The high flexibility of antigen-binding sites in the immunoglobulin, which provides the antibody with a unique capability to access a great variety of configurations with similar stabilities, was long ago proposed to be the basis of this binding diversity. ¹³⁸ In more detail the interplay between the intrinsic disorder, antigenic structure and immunogenicity has been recently overviewed to emphasize the crucial role of intrinsic disorder in the development of immune response. ²² For example, the conformational flexibility of antibodies drives their

polyreactivity, thus expanding the antigen-binding capacity of the antibody repertoire. On the other hand, short intrinsically disordered regions are likely important for the antigenicity of continuous determinants. Furthermore, the conformational flexibility and spatial adaptation play important roles in the antigen-antibody recognition and interaction. Finally, short intrinsically disordered regions are good antigens, whereas several long disordered regions and intrinsically disordered proteins initiate weak immune responses or are even completely non-immunogenic. ²² Based on these observations it has been hypothesized that the role of intrinsic disorder in immunogenicity and antigenicity of a protein depends on the length of the disordered segment: short disordered regions (usually five to eight residues) are important for the development of the immune response to continuous epitopes, whereas long disordered regions (longer than 30 amino acids) are less likely to be immunogenic. ²²

The role of intrinsic disorder in autoimmune diseases has also been emphasized recently. 139 The observation that the majority of the nuclear systemic autoantigens are extremely disordered proteins allowed the authors to introduce a new model of autoimmunity, disorder-based epitope spreading. 139

Another example that illustrates the importance of disorder for *immune response* is the unstructured nature of the interferon tails. 140

Finally, cytoplasmic domains of several immune receptors members of the family of multichain immune recognition receptors (MIRRs) (e.g., T-cell receptors (TCRs), B-cell receptors (BCRs), and the high-affinity IgE receptor) have signaling subunits carrying so-called immunoreceptor tyrosine-based activation motif (ITAM). ^{141–143} ITAM-containing cytoplasmic domains of signaling subunits of several MIRRs are intrinsically disordered. ¹⁴⁴, An intriguing feature of these signaling subunits is their tendency for the specific homooligomerization, which is not accompanied by their folding. ¹⁴⁵, ¹⁴⁶

Ribosome biogenesis and rRNA processing—Ribosomes are responsible for the production of the entire complement of proteins required for cellular maintenance, growth, and survival. Eukaryotic ribosomes contain four RNA molecules: 25S, 18S, 5.8S, and 5S. The 5S rRNA is transcribed by RNA polymerase III, while the three other rRNA molecules are transcribed as a long 35S polycistronic precursor by RNA polymerase I. 147 Ribosome biogenesis and rRNA processing are universal cellular processes, which encompass complicated series of events involving hundreds of transiently interacting components. It has been shown, for example, that in Saccharomyces cerevisiae the biogenesis of pre-18S ribosomal RNA is controlled by a large ribonucleoprotein (RNP) complex, which contains the U3 small nucleolar RNA (snoRNA) and 28 proteins. ¹⁴⁸ The analysis yielded five small subunit ribosomal proteins (Rps4, Rps6, Rps7, Rps14 and Rps28) among other proteins. Intriguingly, in eukaryotic cells, ribosomal protein S6 (Rps6) is the major phosphorylated protein on the small ribosomal subunit, ¹⁴⁹ suggesting that this protein might contain functionally important intrinsically disordered regions (see below, section dedicated to the posttranslational modifications). Furthermore, bioinformatics analysis revealed that 14 of the U three proteins (Utps) bear different repeats comprising crucial regions of their protein-protein interaction domains (WD repeats, coiled-coil domains, HEAT repeats and a crooked-neck-like (crnlike) tetratrico peptide repeat (TPR)). ¹⁴⁸ The crn-like TPR is found in several proteins involved in other RNA processing events including pre-mRNA splicing (Prp42, Prp6 and Clf1) and polyadenylation (RNA14). ¹⁵⁰ NMR analysis of the solution structure of the cytosolic TPR domain of Tom20 in the complex with the presequence peptide revealed that the C-terminal region of this protein (residues 105–145) is disordered. ¹⁵¹

Chondrogenesis—This is the earliest phase of skeletal development, the process by which the cartilage is formed. Cartilage is an elastic connective tissue found in such parts of the body

as the joints, outer ear, and larynx. Furthermore, cartilage represents the major constituent of the embryonic and young vertebrate skeleton, which is converted largely to bone with maturation. Chondrogenesis involves multiple steps, including mesenchymal cell recruitment and migration, condensation of progenitors, chondrocyte differentiation, and maturation, resulting in the formation of cartilage and bone during endochondral ossification. ¹⁵² This complex process is precisely controlled by interactions with the surrounding matrix, growth and differentiation factors, as well as other environmental factors responsible for the initiation or suppression of the cellular signaling pathways and for the regulation of transcription of specific genes. ¹⁵³ For example, the development of vertebrate limb is controlled by the fibroblast growth factor, bone morphogenetic protein (BMP, a secreted signaling molecule, multifunctional growth factor belonging to the transforming growth factor β superfamily), Wnt and hedgehog pathways. ¹⁵⁴ Recently, crucial roles of different mediators (including GADD45β, transcription factors of the Dlx, βHLH, leucine zipper, and AP-1 families, and the Wnt/β-catenin pathway) that interact at different stages during chondrogenesis have been revealed. ¹⁵³ Also, members of the mammalian RUNX protein family, which includes three transcription factors RUNX1, RUNX2, and RUNX3, are expressed during chondrogenesis. These transcription factors also play active roles in mesenchymal condensation, chondrocyte proliferation, and chondrocyte maturation, and regulate transcription of target genes. ¹⁵⁵ Thus, regulation and control of chondrogenesis involve multiple players, many of which possess functional disordered regions. For example, the abundance and functional roles of intrinsic disorder in transcription factors were already discussed (see section entitled Transcription and transcription regulation), whereas the role of disorder in the leucine zippers will be discussed in the second paper of this series.²⁹

Growth regulation—Numerous proteins and pathways are implemented in *growth regulation*. For example, *cyclin* G was shown to be highly expressed in regenerating hepatocytes and motoneurons and in rapidly growing cancer cells and to have growth-promoting functions. ¹⁵⁶, ¹⁵⁷ *Cyclin* G interacts with cyclin-dependent kinase 5 (cdk5) and GAK, a cyclin G-associated kinase, ¹⁵⁸ as well as with with the B' subclass of PP2A phosphatase. ¹⁵⁹ In addition, cyclin G directly interacts with Mdm2 and can stimulate the ability of PP2A to dephosphorylate Mdm2. ¹⁵⁹ Furthermore, cyclin G was one of the earliest p53 target genes to be identified. ¹⁶⁰ This suggests that cyclin G is a key regulator of the p53-Mdm2 network. The role of intrinsic disorder in p53 function was already discussed.

Functions associated with intrinsically disordered proteins

Table 3 presents a list of the top 20 SwissProt functional keywords associated with intrinsic disorder.

Ribonucleoproteins—Numerous facts have been accumulated to demonstrate intrinsic disorder is crucial for function of different *ribonucleoproteins*. In fact, ribonucleoprotein assembly is nearly always accompanied by changes in the conformation of the interacting RNA or protein, or both. $^{161-163}$ For example, the inter-domain linkers of sex-lethal protein (SXL) possess significant disorder, which provides the RNA recognition motifs (RRMs) with a possibility to be flexibly tethered in solution. 164 Another example is ribonuclease P (RNase P), a ribonucleoprotein complex containing one RNA subunit and at least one protein subunit. RNase P is involved in pre-tRNA processing. 165 In *E.coli*, RNase P consists of a small (119 amino acid residue) C5 protein bound to the much larger (377 nucleotide) P RNA subunit. 166 The C5 protein of *E.coli* is essentially disordered in buffer alone, but gains significant amount of ordered secondary structure in an anion-dependent manner. 167 A similar behavior was also described for the *Bacillus subtilis* RNase P. 168

Ribosomal proteins—The assembly of the ribosome, which involves the sequential binding of numerous proteins *via* multiple pathways leading to large-scale changes in the conformation of the associated RNA and proteins, represents an extreme case involving dramatic structural changes induced by protein-RNA interaction. ¹⁶⁹—172 In fact, many ribosomal proteins have been shown to be significantly disordered prior binding to rRNA and to acquire ordered structure during ribosome formation. ⁷, ¹⁶, ¹⁷

Developmental proteins— α -Fetoprotein (AFP), a member of the family of albumin-like proteins, is a serum glycoprotein belonging to the intriguing class of onco-developmental polypeptides. AFP is homologous to human serum albumin (HSA). ¹⁷³ Similarly to HAS, AFP is able to bind a number of small molecules, including metal ions, estrogens and different fatty acids (reviewed in ¹⁷⁴). Importantly, the removal of all ligands from AFP is accompanied by a complete loss of rigid 3-D structure (reviewed in ¹⁷⁴).

Hormones and growth factors—Growth hormone (GH), prolactin (PRL) and placental lactogen (PL), being the pituitary hormones, are members of an extensive cytokine superfamily of hormones and receptors that share many of the same general structure-function relationships in expressing their biological activities. 175 These hormones were shown to have two receptor-binding sites that have different topographies and electrostatic character. This feature is crucial for the regulation of these systems by producing binding surfaces with dramatically different binding affinities to the receptor extracellular domains. The receptor evidently possesses an exceptional conformational plasticity to be able to bind the topographically dissimilar sites on the hormone. 175 Human parathyroid-hormone-related protein (hPTHrP) is a hormone that is over-expressed by a large number of tumors and is produced by a variety of normal cells. The N-terminal fragment (1–34) of hPTHrP is responsible for the major biological functions of this hormone. Furthermore, this fragment is mostly unstructured possessing only a small content of α -helical secondary structure. 176 Secretin, a gut hormone consisting of 27 amino acid residues, was shown to be completely unfolded in aqueous solution but gain a fully ordered structure in the presence of 40% trifluoroethanol. 177

Activators, repressors, cytokines, protease inhibitors, antimicrobial peptides and amphibian defense peptides—Several other function-related keywords, that are associated with intrinsically disordered proteins or regions are illustrated below. Protein AphA is a homodimeric member of a family of transcriptional activators. Transcriptional activators, including nuclear receptors, activate target genes through two broad classes of coactivators - those that remodel/modify chromatin and those that directly interfere with the general transcription machinery to facilitate formation and/or function of the preinitiation complexes. The AphA monomer is highly unstable by itself (i.e., likely it is highly disordered) and the dimer is formed in such a way that the two AphA chains wrap around each other, ¹⁷⁸ suggesting that the dimmer arose via a disorder-to-order transition. The first 61 amino acid residues of the DNA-free lac repressor (i.e., a fragment which includes headpiece and the hinge region) are disordered, and thus are unobserved in the crystal structure. ¹⁷⁹ Lymphotactin (Ltn) is unique member of family of pro-inflammatory activation-inducible cytokines. Ltn possesses a unique C-terminal extension, which is required for biological activity 180, 181 and which is disordered and highly mobile. ¹⁸² Analysis of the Bowman-Birk *protease inhibitor* by Raman optical activity revealed that it possesses a "static" type of disorder similar to that in disordered states of poly(L-lysine) and poly(L-glutamic acid). ¹⁸³ The pediocin-like class IIa bacteriocins, which are *antimicrobial peptides* from lactic acid bacteria, ¹⁸⁴, ¹⁸⁵ were shown to be significantly unstructured in water. ¹⁸⁶ Similarly, hylaseptin P1, an amphibian defense peptide, is in a random coil conformation in aqueous solutions. 187

Neuropeptides—Pituitary adenylate cyclase activating polypeptide (PACAP), 188 which occurs naturally in two forms consisting of a 38 amino acid peptide amide (PACAP38) and its 27 amino acid N-terminus (PACAP27), belongs to the secretin/glucagons/*vasoactive* intestinal peptide (VIP) family. 189 Structural analysis of PACAP38 and PACAP27 revealed that these two neuropeptides are mostly disordered and retain only small transitory amounts of stable structure in aqueous solution. 190 Other opioid peptides are the enkephalins. The term enkephalin mainly refers to two peptides, [Met]-enkephalin and [Leu]-enkephalin, that both are products of the proenkephalin gene. [Met]-enkephalin is Tyr-Gly-Gly-Phe-Met; [Leu]-enkephalin has Leu in place of Met. Recently performed structural characterization of methionine and leucine enkephalins by hydrogen/deuterium exchange and electrospray ionization tandem mass spectrometry revealed that the monomer forms of both peptides adopt an unfolded conformation in aqueous solvent, whereas they prefer β-turn secondary structure under the membranemimetic environment. 191

GTPase activation and GTPase-activating proteins (GAPs)—The GTP-GDP conversion by guanine nucleotide binding proteins (GNBPs) represents an important timer in intracellular signaling and transport processes. GNBPs are highly abundant in different genomes. For example, there are at least 140 small GTPases encoded in human (including the Ras, Rho, Arf, Rab and Ran GTPases), with various subclasses of this protein superfamily being implicated in almost all aspects of cell biology, including proliferation, nucleocytoplasmic transport, differentiation, vesicle trafficking, cytoskeletal organization and gene expression. ¹⁹² These small GTPases are considered to be molecular switches, the cycling of which between active and inactive forms is regulated by cellular factors. ¹⁹² There are two major classes of GNBP regulators, the guanine nucleotide exchange factors (GEFs), which promote the formation of active GTP-bound GTPases and the GTPase activating proteins (GAPs), which promote GTPase inactivation by stimulating GTP-hydrolysis activity. ¹⁹³ In fact, the natural rate of GNBP-mediated GTP hydrolysis is slow but the reaction is accelerated by up to five orders of magnitude by the interaction of GNBPs with GAPs. 194 At least 160 human genes have been recently predicted to encode proteins that resemble GAPs for various members of the Ras GPTase superfamily. 195 Furthermore, ~ 0.5% of all predicted human genes likely encode GAPs suggesting that these proteins have widespread and important roles in GTPase regulation. Finally, such famous domains as ankyrin, BAR, BTK, CH, CNH, PDZ, PTB, RUN, SAM, SH2, SH3, WW and many others are all GAPs. 196

Chromatin regulator—Several nuclear proteins serve as *chromatin regulators*, being involved in modulation of chromosome structure, chromatin and nucleosome remodeling and therefore playing a role in the controlling of gene transcription. Members of the HMGA family of non-histone chromatin proteins (formerly known as HMGI/Y proteins) serve as an illustrative example of such *chromatin regulators*. ¹⁹⁷ HMGA proteins are the founding members of a new class of regulatory elements called 'architectural transcription factors' that participate in a wide variety of cellular processes including regulation of inducible gene transcription, integration of retroviruses into chromosomes, the induction of neoplastic transformation and promotion of metastatic progression of cancer cells. ¹⁹⁸ HMGA proteins are highly flexible and are characterized by the total lack of ordered structure.

Pyrogen—Substances that can cause a rise in body temperature are known as *pyrogens*. Fever is the multiphasic response of elevation and decline of the body core temperature regulated by central thermoregulatory mechanisms localized in the preoptic area of the hypothalamus. Some *cytokines* (which are highly inducible, secreted proteins mediating intercellular communication in the nervous and immune system), including interleukin 1 (IL-1), interleukin 6 (IL-6) and the tumor necrosis factor alpha (TNF α), act as endogenous pyrogens. ²⁰² The role of intrinsic disorder in cytokine function was already discussed.

Opioid peptides and endorphin—Opioid peptides are short natural peptides that mimic the effect of opiates in the brain and therefore are potent pain suppressants. Some opioid peptides (e.g., endorphin, dynorphin and endorphin) are produced endogenously, some are produced by microbes (deltorphins and dermorphine), whereas others are absorbed from partially digested food (casomorphins, exorphins, and rubiscolins). Opioid peptides mediate their physiological and pharmacological effects through three major opioid receptor types (μ , δ , κ), 203 which are the members of the G protein-coupled receptor (GPCR) family. 204 The 1 H-NMR spectra of human β -endorphin (the largest natural opioid peptide of 31 amino acid residues) indicate that the peptide exists in random-coil conformation in aqueous solution but becomes helical in mixed solvent. 205

Inhibitors—The activity of many important proteins is regulated by specific proteins-inhibitors. It has been already mentioned that the functionality of *Cdk inhibitor proteins* relies on intrinsic disorder. *Serine protease inhibitor* elafin is a 57 amino acid residue peptide inhibiting human leukocyte elastase, porcine pancreatic elastase and proteinase-3. Elafin was shown to be almost completely unfolded in aqueous solutions. ²⁰⁷

Protein phosphatase inhibitors—Calcineurin (CaN) is a calcium- and calmodulin-dependent protein serine/threonine phosphate, which is critical for several important cellular processes, including T-cell activation. 208 CaN is a heterodimer composed of subunits A and B (CaNA and CaN, respectively). 208 CaN phosphatase activity is regulated by 22 binding to CaNB and by 22 -induced binding of calmodulin (CaM) to CaNA. 209 Activity of CaN is modulated by a number of factors, including an autoinhibitory domain (residues 45 -479), which binds in the active site cleft in the absence of 22 -/CaM and inhibits the enzyme, acting in concert with the CaM binding domain to confer CaM regulation. 209 Analysis of CaN crystal structure revealed that CaNA residues $^{1-13}$, 374 -468, and 487 -521 are not visible in the electron density map; i.e., disordered. 209 Importantly, long disordered region 374 -468 includes the CaM-binding helix. 209 Therefore, this transient helix in CaN becomes bound and surrounded by CaM, turning on the CaN's serine/threonine phosphatase activity. Locating the CaN helix within the disordered region is essential for enabling CaM to surround its target upon binding. 6

Cyclin—The progression of cells through the *cell cycle* is regulated by several specific proteins, known as *cyclins* and cyclin-dependent kinases. The concentrations of cyclins vary in a cyclical fashion during the cell cycle. They are produced or degraded as needed in order to drive the cell through the different stages of the cell cycle. The crucial role of intrinsic disorder in function and regulation of Cdk–cyclin complexes was already discussed.

Biological processes and functions associated with ordered proteins

Table 4 and Table 5 list the top 20 biological processes and functions that are significantly associated with predicted order. An examination of order-correlated keywords suggests the presence of 5 major functional categories: (1) catalysis (this category includes all functions listed in Table 5; i.e., oxidoreductase, transferase, hydrolase, lyase, glycosidase, kinase, isomerase, ligase, decarboxylase, glycosyltransferase, protease, acyltransferase, monooxygenase, aminotransferase, metalloprotease, methyltransferase, aminoacyl-tRNA synthetase, aminopeptidase, and dioxygenase); (2) transport (electron transport, sugar transport, and transport), (3) biosynthesis (amino-acid biosynthesis, GMP biosynthesis, gluconeogenesis, amino-acid biosynthesis, pyrimidine biosynthesis, peptidoglycan synthesis, lipopolysaccharide biosynthesis, aromatic amino acid biosynthesis, branched-chain amino acid biosynthesis, purine biosynthesis, lipid a biosynthesis, and lipid synthesis) (4) metabolism (carbohydrate metabolism, tricarboxylic acid cycle, aromatic hydrocarbons catabolism, and one-carbon metabolism) (5) trans-membrane proteins (porins). Note that catalysis overlaps

strongly with biosynthesis and metabolism in that all of the proteins associated with these keywords are enzymes. The proteins associated with transport are often membrane proteins and are necessarily structured so that their backbone hydrogen bonds are formed in the low dielectric environment of the membrane. Other transport-associated proteins (which are not the membrane proteins) often need to bind very small molecules (or a single atom such as metals, or even electrons), which requires a precise coordination, and therefore, a well-defined structure. Proteins from the fifth category, *porins*, represent are transmembrane proteins that are large enough to facilitate passive diffusion. They are prevalent in the outer membrane of the mitochondria and Gram-negative bacteria. Porins are almost entirely composed of beta sheets and control the diffusion of small metabolites like sugars, ions, and amino acids. ²¹⁰ They have been shown to have a highly stable structure using various characterization methods. ²¹¹ For example, porin from *Paracoccus denitrificans* is extremely stable toward heat, pH, and chemical denaturants. ²¹² Thus, current result agrees with our previous observations that proteins involved in catalysis, transport, biosynthesis and metabolism are less disordered than regulatory proteins.

Interestingly, the smaller disorder content as well as the greater amount of structural information (*e.g.* a greater PDB coverage) has previously been reported for proteins from the first four functional categories as compared to highly disordered signaling and cancerassociated proteins. ¹⁶ Thus, the current result agrees with our previous observations that proteins involved in catalysis, transport, biosynthesis and metabolism are less disordered than regulatory proteins.

Finally, one noticeable exception should be mentioned here. Although glycosidases are among the top 20 proteins with predicted functional order (Table 5), many of them in fact possess large disordered regions, even though their catalytic function requires a well defined structure. This is especially true for cellulases (Biological process: cellulose degradation, strong correlation with predicted order, see Table S1) for which protein disorder has been experimentally determined. ²¹³, ²¹⁴ These cellulases are composed of a catalytic domain, linked to a cellulose binding domain through a long disordered linker (109 amino acid residues in Cel5G, an endoglucanase from *Pseudoalteromonas haloplanktis*), which could be considered as an entropic spring. In fact, the SAXS analysis of dimensions, shape, and conformation of Cel5G full length in solution and especially of the linker between the catalytic module and the cellulose-binding module revealed that the linker is unstructured, and unusually long and flexible. ²¹³ This modular organization and the presence of a disordered linker are crucial to optimize the biphasic process of crystalline cellulose degradation.

Another example of an enzyme that possesses functional disordered regions is retinaldehyde dehydrogenase II (RalDH2). This enzyme converts retinal to the transcriptional regulator retinoic acid in the developing embryo. It has been shown that a 20-amino acid span in the substrate access channel is disordered, but folds during the course of catalysis and provides a means for an enzyme that requires a large substrate access channel to restrict access to the catalytic machinery by smaller compounds that might potentially enter the active site and be metabolized. Therefore, RalDH2 represents a unique example of a protein that exhibits a catalytic activity in which a large disordered region folds upon catalysis.

Comparing the identified disorder functions with literature findings

Recently, literature analysis identified 28 functions associated with 98 confirmed disordered regions containing 30 or longer contiguous disorder residues. ^{8,9} These functions were grouped into four broad categories: molecular recognition, molecular assembly, protein modification, and entropic chains. Entropic chains carry out functions that depend directly on the disordered state, and so such functions are simply outside the capabilities of fully folded structures. ^{8,9} The use of partially folded subunits for molecular assembly appears to have significant

advantages compared to the use of ordered subunits. 21 , 22 Molecular recognition appears to be a common function for both ordered and disordered proteins: molecular recognition by disordered proteins may be primarily used for signaling whereas recognition by ordered proteins may be primarily used for catalysis, 8 , 9 or for the assembly of functional complexes. Finally, sites of some types of posttranslational modification frequently occur within the regions with very strong preference for disorder. 8 – 11 , 18 , 19 , 22 , 216

Out of 28 functions associated with the confirmed disordered regions, 13 were also found in the Swiss-Prot keyword list. Eleven of these functions are strongly correlated with predicted disorder with p-value above 0.95. Table 6 lists these 11 functions together with their z-scores. Furthermore, previously reported ¹⁶ strong functional correlations to intrinsic disorder were also found by using the method proposed in this study. These results strongly support the validity of the proposed statistical methodology for finding disorder-correlated functions.

Conclusions

We proposed a statistical approach that estimates the correlations between protein structure and protein function. As an application, we studied the relationship between intrinsic protein disorder and function in the Swiss-Prot database. Overall, 238 Swiss-Prot functional keywords were discovered to be strongly associated with predicted intrinsic disorder, and 302 function keywords were shown to be strongly correlated with predicted order. We validated a significant fraction of these findings by comparing them to literature data. The numerous correlations between the known experimental data and the results of the analysis demonstrate the general validity of our approach. However, a more thorough comparison with literature data will be needed to determine the frequencies with which exceptions occur as compared to the observed trends. In our original, manual curation of disorder-function correlations, we determined that each function was unambiguously associated with a specific region of disorder.^{8, 9} While the current methodology does not determine whether each function is directly associated with a disordered segment, the literature data verify that these functions are indeed, for the most part, associated with regions of disorder. Of special interest is that disease related proteins were shown to have the high correlation with disordered regions of proteins (see the last paper of this series³⁰). Overall, this approach provides an innovative and relevant method to examine protein structure-function relationships.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgment

The authors express their deepest gratitude to Celeste Brown and Predrag Radivojac for numerous valuable discussions. This work was supported by grants from the National Institutes of Health LM007688-0A1 (A. K. D. and Z. O.) and GM071714-01A2 (A.K.D and V.N.U.), and by the Indiana Genomics Initiative (INGEN) (A. K. D.). INGEN is supported in part by Lilly Endowment Inc. The Programs of the Russian Academy of Sciences for the "Molecular and cellular biology" and "Fundamental science for medicine" provided partial support to V. N. U., and L. M. I. was supported by the NSF grant MCB 0444818.

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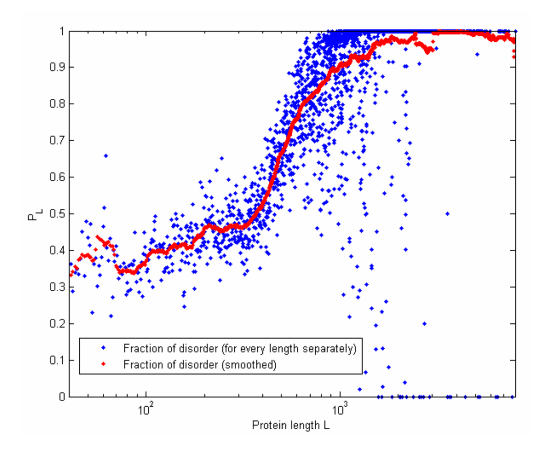
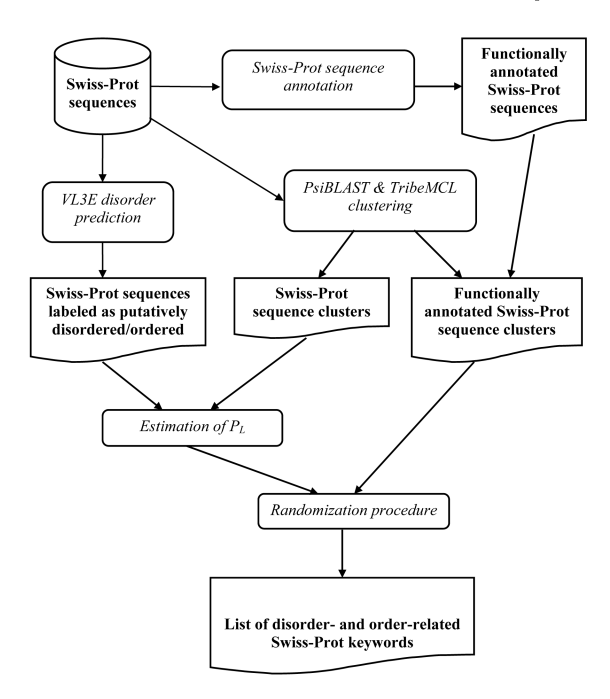


Figure 1. Fraction of putative disorder as a function of sequence length. The smoothed curve uses averaging window of size equal to 20% of the sequence length.



Schematic representation of the algorithm for extracting disorder- and order-related keywords.

Table 1

Summary of association between the prediction of long disordered regions and keywords for each of the 11 functional categories. For each category, the table lists the total number of keywords associated with it (out of the 710 Swiss-Prot functional keywords), as well as number of keywords associated with predicted order and disorder.

Functional category	# Keywords	# Keywords (p-value < 0.05)	# Keywords (p-value > 0.95)
Biological process	301	174	73
Cellular component	77	23	33
Coding sequence diversity	9	0	6
Developmental stage	4	0	3
Disease	17	0	11
Domain	34	9	21
Ligand	72	41	17
Molecular function	143	37	51
PTM	37	11	18
Technical term	12	7	2
Tissue	4	0	3
TOTAL	710	302	238

Table 2 Top 20 of processes that have strongest correlation with predicted disorder

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Keywords	Number of proteins	Number of families	Average sequence length	Z-score	P-value
Differentiation	1406	422	439.25	18.81	-
Transcription	11223	1653	442.64	14.62	
Transcription regulation	9758	1554	413.31	14.33	
Spermatogenesis	332	189	280.49	13.9	
DNA condensation	317	130	300.06	13.34	-1
Cell cycle	4278	612	494.17	12.17	
mRNA processing	1575	249	515.55	10.92	
mRNA splicing	716	180	459.06	10.13	-1
Mitosis	718	215	620.43	9.42	
Apoptosis	810	211	465.48	9.35	
Protein transport	3081	579	421.73	8.77	1
Meiosis	284	170	639.16	8.7	1
Cell division	3466	385	451.63	8.51	
Ubl conjugation pathway	1254	244	525.99	8.13	1
Wnt signaling pathway	417	41	476.84	6.58	
Neurogenesis	322	74	667.4	6.56	
Chromosome partition	556	<i>L</i> 9	495.39	6.39	1
Ribosome biogenesis	319	71	391.79	5.9	
Chondrogenesis	64	9	332.85	5.58	
Growth regulation	155	45	354.9	5.14	1

Table 3 Top 20 of functions that have strongest correlation with predicted disorder

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Keywords	Number of proteins	Number of families	Average sequence length	Z-score	P-value
Ribonucleoprotein	12236	412	150.55	22.13	-
Ribosomal protein	11692	330	140.58	20.63	-
Developmental protein	3260	721	477.93	19.28	
Hormone	1187	161	141.13	15.58	
Growth factor	785	84	255.7	11.16	-
Cytokine	668	110	213.28	10.21	
Neuropeptide	268	209	95.08	9.65	
Activator	3086	573	428.47	9.04	-
GAP protein	47	2	232.96	7.42	
Antigen	1113	455	437.48	6.99	
Repressor	2309	449	374.46	6.92	-
Chromatin regulator	334	100	801.24	6.7	
Pyrogen	37	2	262.59	6.44	
Vasoactive	125	39	160.39	5.56	-
Amphibian defense peptide	123	148	50.64	5.44	
GTPase activation	311	70	831.03	5.36	
Endorphin	42	4	226.68	5.35	-
Opioid peptide	24	4	216.96	5.14	
Protein phosphatase inhibitor	47	∞	366.51	5.07	-
Cyclin	182	25	430.58	4.88	1

NIH-PA Author Manuscript **Table 4** Top 20 of processes that have strongest correlation with predicted order NIH-PA Author Manuscript NIH-PA Author Manuscript

Keywords	Number of proteins	Number of families	Average sequence length	Z-score	P-value
CMB his mostly and	300	·	11 227	62.61	
GMF prosynthesis	677	c	4/3.11	-17.02	0
Amino-acid biosynthesis	2004	212	361.5	-17.11	0
Transport	19888	2199	378.13	-14.87	0
Electron transport	4633	346	272	-13.72	0
Lipid A biosynthesis	533	13	291.25	-13.22	0
Aromatic hydrocarbons catabolism	320	105	300.36	-12.37	0
Glycolysis	2255	50	390.64	-12.14	0
Purine biosynthesis	1208	28	445.46	-11.89	0
Pyrimidine biosynthesis	1310	27	383.27	-11.7	0
Carbohydrate metabolism	1797	180	404.2	-11.68	0
Branched-chain amino acid biosynthesis	963	26	404.12	-11.11	0
Lipopolysaccharide biosynthesis	481	102	335.93	-11.09	0
Sugar transport	903	109	387.37	-11	0
Antibiotic resistance	1203	177	354.24	-10.66	0
Lipid synthesis	2184	122	328.02	-10.17	0
Tricarboxylic acid cycle	1013	54	460.88	-10.04	0
Arginine biosynthesis	1353	17	414.06	-9.53	0
Ion transport	5275	459	464.46	-9.37	0
Rhamnose metabolism	85	4	372.84	-9.12	0
Peptidoglycan synthesis	1839	38	372.73	-9.03	0

Table 5 Top 20 of functions that have strongest correlation with predicted order

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Keywords	Number of proteins	Number of families	Average sequence length	Z-score	P-value
Oxidoreductase	14995	992	376.63	-29.54	0
Transferase	26525	1606	445.17	-24.25	0
Lyase	7262	347	377.92	-22.64	0
Hydrolase	20464	1995	430.68	-21.75	0
Isomerase	4487	220	383.98	-14.18	0
Glycosidase	1826	244	444.73	-13.98	0
Glycosyltransferase	2950	261	437.53	-12.51	0
Acyltransferase	2239	179	402.83	-10.85	0
Methyltransferase	3524	224	349.6	-10.53	0
Kinase	7017	322	448.29	-10.22	0
Ligase	8010	230	529.41	-10.06	0
Decarboxylase	1293	63	345.26	99.6–	0
Monooxygenase	1668	73	444.87	-9.26	0
Metalloprotease	1100	109	553.73	-7.89	0
Aminopeptidase	452	39	509.17	-7.55	0
Dioxygenase	360	99	433.2	-7.32	0
Aminoacyl-tRNA synthetase	3402	37	571.83	-7.15	0
Protease	4423	380	549.7	-7.1	0
Aminotransferase	955	28	420.27	-6.02	0

Table 6

All (11) Swiss-Prot keywords associated with at least one of the 98 confirmed long disordered protein regions^{8,9}. For each function, number of the associated regions (out of 98) and z-ratio are listed.

Function	Number of regions associate with function in literatures (Dunker, et al 2002)	Z-ratio in SwissProt database
Protein-DNA interaction	19	18.2
Phosphorylation	16	27.1
Structural mortar	>10	3.0
Ubiquitination	7	8.7
Protein-rRNA interaction	5	11.5
Fatty acylation	4	6.0
Protein-genomic RNA binding	3	17.7
Glycosylation	3	6.8
Methylation	1	2.9
ADP-ribosylation	1	2.0
Protein-tRNA interaction	1	1.8