Meta-structure Correlation in Protein Space Unveils Different Selection Rules for Folded and Intrinsically Disordered Proteins

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Abstract

The number of existing protein sequences spans a very small fraction of sequence space. Natural proteins have overcome a strong negative selective pressure to avoid the formation of insoluble aggregates. Stably folded globular proteins and intrinsically disordered proteins (IDP) use alternative solutions to the aggregation problem. While in globular proteins folding minimizes the access to aggregation prone regions IDPs on average display large exposed contact areas. Here, we introduce the concept of average meta-structure correlation map to analyze sequence space. Using this novel conceptual view we show that representative ensembles of folded and ID proteins show distinct characteristics and responds differently to sequence randomization. By studying the way evolutionary constraints act on IDPs to disable a negative function (aggregation) we might gain insight into the mechanisms by which function-enabling information is encoded in IDPs.
Intrinsically Disordered Proteins (IDP) are now recognized as a major class of natural proteins following a long invisibility period arising mainly from their failure to adopt a single, well-defined conformation capable of forming three-dimensional crystal lattices. IDPs are especially abundant in eukaryotes and predominantly associated with regulatory processes. These two facts are unlikely to be unrelated and the exploitation of IDP may be one of the molecular features that enabled evolution beyond simple unicellular prokaryotes. The important functional roles of IDPs are supported, among others, by evidences about their tight regulations at all levels.1,2

Recognition of the widespread occurrence of IDPs was made possible by the access to the exhaustive list of primary sequences of complete organisms through full genome sequencing and the development of bioinformatic tools that could correctly predict the probability of disorder from the primary sequence.5-14 Hallmarks of protein disorder are the low frequency of aromatic and other hydrophobic residues combined with a high frequency of charged, polar and structure breaking residues.

The growing evidence that a unique folding may not be required to perform a function in the case of IDP suggests that the structure-activity paradigm has to be expanded into a more general one involving more sophisticated concepts about ensemble averaging and sampling of accessible conformational space. While the rules by which information is encoded in flexible proteins are still elusive, successful evolution of protein sequences (Survival of the fittest) in general points to the relevance of a common property: avoiding the formation of amyloids. In other words, amyloid formation (or its prevention) can be considered a common functional requirement to which all proteins have adapted. Although there exist different ways to achieve it, looking for common principles that apply to whole classes of proteins may provide some insight into the information-encoding capabilities of IDPs.
In this article we explore two aspects of IDPs that can be derived from the analysis of primary sequences. First we compare the occurrence of amyloidogenic regions, as predicted by the Waltz algorithm,\textsuperscript{15} in folded and intrinsically disordered proteins. Second, we introduce the average meta-structure correlation maps (AMCM) and compare natural and randomized sequences to unveil conserved features that are different in IDPs and folded proteins beyond their different amino acid composition. We suggest that the observed differences may reflect different strategies to avoid amyloid formation by folded and disordered proteins.

**Methods**

The propensity for forming insoluble amyloids was assessed by the Waltz algorithm.\textsuperscript{15} The Waltz program allows for the analysis of amyloidogenesis exclusively from the primary sequence and gives the location of sequences predicted to have high propensities to form amyloids.

The meta-structure concept was recently introduced by one of us to extract implicit structural information encoded in the sequence and derived from topological pairwise propensities extracted from the analysis of proteins of known structure.\textsuperscript{16} It was shown that although meta-structures can be derived directly from sequences, they are much more conserved in evolutionarily related proteins.\textsuperscript{16} In the meta-structure concept, a 3D protein structure is perceived as a network of residue interactions, in which nodes refer to residues and edges indicate the existence of (through space) neighbourhood relationships. The mutual topological relationship between two residues (A,B) is quantified by the shortest path length across the network and characteristically depends on the amino acid types (A,B) and their primary sequence distance, \( l_{AB} \). The frequency of different topological relationships was evaluated in a subset of structures taken from the PDB database and stored as pairwise statistical distribution functions.
The meta-structure analysis employs this statistically derived topological information to extract higher order information implicitly contained in the primary sequence, the secondary structure parameter and the local compaction: local 2nd structure information is quantified by a parameter that takes positive values for helical structures and negative values for predicted beta sheets. Compactness refers to the tendency of local regions of the sequence to be buried from solvent access. We refer to the sequence of pairs of local secondary structure and compaction values as the protein meta-structure. Although only structured protein parts were used in the derivation of the pairwise distribution function, the methodology is not limited to structured proteins but rather provides quantitative information about the most probable network topology of a given protein, folded or unfolded.

For a given protein, the average value of each of the meta-structure parameters can be calculated to provide a global pair of parameters. Notice that since the residue-specific meta-structure parameters reflect the sequence context, the global average values are sensitive to the actual protein sequence, and do not reflect simply the protein composition. The pairs of average meta-structure parameters for a given protein can be represented as a point in a two dimensional plot, that we refer to as average meta-structure correlation map (AMCM).

The potential of the meta-structure approach for high-throughput IDP identification was already demonstrated. Average residue compactness values of proteins were introduced as measures for protein foldedness. While stably folded proteins display average compactness values of about 300, significantly smaller values (<200) are found for structurally flexible proteins (intrinsically disordered/unstructured). Applications of this approach to proteins from different kingdoms (archea, prokaryote and eukaryotes) corroborated the widely accepted notion that lower organisms have only few unstructured proteins, IUPs/NUPs, (archea and prokaryotes: 1.7%-3.5%), whereas for eukaryotes a significant fraction of the proteome falls into this category.
This criteria (average compactness value < 200) was used to define a set of human IDP proteins (see below).

In this study, folded proteins are represented by a non-redundant representation of 27780 proteins form the Protein Data Bank (which we will refer to as PDB). A set of highly disordered proteins was selected by calculating the meta-structure parameters for all human proteins and selecting those with an average compactness value < 200. This set (referred to as IDP200) contains 1012 proteins. Of these, 50.4% have more than 90% of their residues disordered and 97% are predicted to be disordered in more than 50% of their sequence, according to the VL2 predictor accessed through the DisProt server. A third set was formed by 164 proteins (which we call DIS50) selected from the DisProt database and annotated to be more than 50% disordered. On a per residue basis, 40.8% of the DIS50 proteins are predicted by VL2 to have more than 90% of their residues in disordered regions and 85.9% of the proteins are more than 50% disordered. 23 out of the 164 sequences of DIS50 were also present in IDP200.

A comparison between predicted disorder and compactness at a residue level for DIS50 and IDP200 is shown in Figure 1. For the DIS50 residues, the frequency of compactness values for residues with a disorder score larger than 50% show a Gaussian shape centered at 200, while the distribution of the residues predicted to be ordered is centered at 300, the same average value found for PDB proteins. For the IDP200 set, which is biased to average compactness values below 200, the distribution of predicted disordered values is centered at around 150 while the small fraction of residues predicted to be ordered has compactness values centered around 250. Thus we conclude that meta-structure derived compactness values can be used as reliable parameters for identifying disordered segments in proteins.
A fourth set (AFR) was generated by collecting the regions of the proteins of the first three sets predicted to form amyloids by Waltz. Reference sets containing randomized sequences for each protein were prepared and are referred to as RPDB, RiDP200, RDIS50, and RA FR. An additional set of 592 sequences (β-protein interactors or BPI) contains natural proteins experimentally identified to be efficiently captured in vivo by amyloid forming peptides as described by Olzscha et al. and was used to validate the amyloidogenesis analysis.

Results and discussion

Firstly, we computed from the Waltz server output the percentage of sequences in the PDB, DIS50 and IDP200 sets that contain at least one amyloidogenic region (Figure 2). As a comparison we also included the BPI set of proteins captured by three different amyloid forming artificial proteins. A large proportion of the sequences of proteins in the PDB are predicted to have at least one amyloidogenic region confirming that folding is indeed preventing amyloid formation. Consistently, the fraction of amyloidogenic sequences computed by Waltz in the β-protein interactors set is larger than the average of the PDB, confirming the capacity of the Waltz algorithm to predict amyloidogenic regions.

The propensity to form amyloids by unfolded proteins in the IDP sets is much lower than that of the proteins present in the PDB in agreement with the notion that alternative strategies to avoid amyloid formation are required for unfolded proteins. Next we computed the total length of the amyloidogenic sequences and the fraction they represent of the total sequence for natural proteins and for a matching set made of randomized sequences. Figure 3 shows histograms of the frequencies of predictions of different proportions of amyloidogenic segments with respect to the total protein
length. The relative importance of amyloidogenic regions with respect to the total sequence is low for a large fraction of the natural sequences present in the PDB and IDP sets have very low tendencies to form amyloids. PDB sequences show a bimodal distribution with a sharp peak of non-aggregation prone sequences, followed by a broad distribution with a maximum at 4% and a slow decay with a substantial number of sequences comprising more than 20% of amyloidogenic regions. The IDP sets can be described by a much narrower distribution with most sequences having less than 10-12% of amyloidogenic regions. The bimodal distribution in the PDB set probably reflects an additional negative selection by the structural biology researchers being able to solve preferentially those proteins that have favorable solution properties.

The sets of randomized sequences show a general tendency to increased amyloidogenesis. This is more pronounced in the PDB. The tendency of RIDP200 to form amyloids is similar to the matching natural sequences. This observation is in agreement with the idea that naturally disordered proteins have an intrinsic lower tendency to aggregate than denatured globular proteins, which primarily comes from the residue composition of disordered proteins, including the avoidance of hydrophobic residues. However, general sequence constrains or preferences for IDPs have been observed,\textsuperscript{19} and are additionally shown in the meta-structure correlation results shown below.

Secondly, the proteins from the different data sets were subjected to a meta-structure analysis. The meta-structure information from each protein in the different sets was represented as points in an Average Meta-Structure Correlation Map (AMCM) in which the average value of the meta-structure derived residue compactness is plotted against the average secondary structure as predicted also from meta-structures. Figure 4 shows AMCM of naturally occurring proteins. The average compactness of the PDB set is, not surprisingly, larger than the DIS50 set. The IDP200
is, by construction, limited to compaction values lower than 200. For comparison we also calculated the AMCM plot for individual segments predicted to be prone to form amyloids (AFR set). The larger scatter is presumably due to the fact that here smaller peptide fragments are analyzed instead of entire protein averages. The compactness values of the aggregation-prone fragments of the AFR set is shifted to higher values than those of the sets of complete proteins from which they were derived.

A clear additional distinction is observed between the sets of globular and disordered complete proteins. The PDB set shows a negative correlation between secondary structure and average compactness indicating that the most compact structures are enriched in β sheets (Pearson correlation coefficient -0.36. Slope of the best linear fit: -0.19 for proteins with average compactness values between 200 and 400). In contrast both IDP enriched sets show a clear positive correlation (Pearson correlation coefficients (slopes): DIS50: 0.62 (0.56); IDP200: 0.46 (0.66), calculated for proteins displaying average compactness values above 100). Thus, more compact structures are associated to increasing helical contents, presumably reflecting the building principle of IDPs comprising locally defined structural elements. Only a few IDPs have negative secondary structure values. Amyloid forming peptides of the AFR set show a large scatter of secondary structure parameters and a low correlation between the two meta-structure parameters.

The observed correlations in full proteins suggests that intricate side-chain interactions leading to large compactness values are preferentially associated to β-strand formation in folded proteins. In contrast, IDP compaction is preferentially achieved by forming less aggregation-prone α-helices.

The location of protein sequences in the average meta-structure correlation maps shows a better correlation with their folding characteristics than the individual parameters. Figure 5 shows a smoothed representation obtained by clustering of
neighbor points in the AMCMs of the different protein sets. It can be seen that folded proteins, IDPs and aggregation prone peptides are located in distinct and only partially overlapping sub-spaces. We thus believe that the AMCM analysis provides a meaningful representation of "protein space". In addition to the analysis given here we anticipate further applications of the AMCM approach to global, large-scale analysis of the architecture and organization of the accessible protein space.

A comparison of the AMCM of natural and randomized sequences shows significant differences between globular and disordered protein sequences. In the data set of randomized IDP sequences the observed correlation between average compactness and 2nd structure changes from positive to negative (Pearson coefficients: DIS50 0.62; RDIS50: -0.25; IDP200: 0.46; RIDP200: -0.20). In contrast random protein sequences of folded proteins maintain a negative correlation similar to natural protein sequences (Pearson coefficients: PDB -0.36; RPDB: -0.50). These observations point to a different origin of the correlations in AMCMs of IDPs and globular proteins.

The physical background rules for IDPs involve both a bias in the amino acid composition, minimizing the occurrence of non-polar residues, and restrictions at the primary structure level that links increased compaction with the minimization of the formation of \( \beta \)-sheets. We suggest that the origin of the correlation between meta-structure parameters in IDPs is predominantly avoiding aggregation when local structures are formed, although other contributions to the low propensity to adopt \( \beta \)-structures by IDP cannot be ruled out, including the modulation of conformational entropy which is easier to achieve through helical conformations than with \( \beta \)-sheets.\textsuperscript{20}

IDPs and globular proteins have been selected during evolution using two distinct strategies to avoid the development of deleterious intermolecular associations.
Interestingly, while folded proteins partially overlap with amyloidogenic peptides, IDP sequences are distinctly different and clearly set apart from the reservoir of sequences prone to aggregation. This statement is in contrast to the widespread belief that disorder increases the tendency to aggregation and disease. However, while a number of disordered proteins are associated with amyloidosis leading to devastating diseases, it is not obvious that the fraction of naturally disordered proteins that can form amyloids is larger than the fraction of naturally occurring stable folded proteins that aggregate upon misfolding. Most globular proteins show a tendency to aggregate when solution conditions or point mutations disturb their natural folding while this is less obvious for naturally disordered proteins. Thus, while folding of globular proteins offers protection against aggregation, disordered proteins have naturally evolved alternative strategies to avoid amyloidogenesis.

An analogy could be drawn between the sequence universe and the surface of the Earth. Most of the Earth is covered with water. Likewise, most sequences would naturally form amyloids. Natural globular/folded proteins and IDPs can be associated to different "emerged continents". Both have in common that they are not "covered with water"/forming amyloids yet they are clearly different. Shores are regions that can be easily flooded by small changes in the environment and would represent "dangerous" sequences prone to form amyloids. They are present in both folded proteins and IDPs. Therefore, the observation of amyloids associated to some IDPs should not hide the fact that the vast majority of IDPs are not aggregating and this is the physical background on which function for IDPs can be built.

The structure-function relationship is one of the guiding paradigms of structural biology. IDPs challenge this principle as their function is associated to the availability of multiple conformations and the sampling of a wide conformational space within characteristic time scales thereby allowing for concerted recognition events by diverse
receptors or chemical reactions to proceed between closely placed complementary reacting centres.

Naturally occurring IDPs in part encode functionality in the form of restrictions of the “random-coil” conformational space actually sampled by a particular IDP sequence. In order to eventually decipher how IDPs information is encoded and stored in the primary sequence a more thorough analysis of network topologies will be valuable. We suggest that meta-structure analysis, in addition to provide a tool to separate IDPs from folded protein structures from the amino acid sequences, may be used in the future to differentiate between functionally related subsets of IDP *en route* to the challenging goal of uncovering how other functions are encoded in IDPs.

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Figure captions

Figure 1. Frequency distribution of compactness values of individual residues obtained from metastructure analysis of the proteins in the DIS50 (left) and IDP200 (right) databases. The compactness distribution of residues predicted to be disordered or ordered are represented in black or red, respectively.

Figure 2. Percentage of sequences with amyloidogenic regions in different datasets. Amyloidogenic regions were predicted using the Waltz algorithm. The β-protein interactors correspond to the sequences of the proteins captured by amyloid forming artificial peptides described in reference 17. Sequences were extracted using the Human Protein Reference Database identifier and include different isoforms (www.hprd.org). Error bars were generated by comparing random subsets of 10% of the databases.

Figure 3. Number of sequences containing different percentage of residues in amyloidogenic regions. The total number of residues present in the different amyloidogenic regions predicted by Waltz using the natural sequences and one randomized version of each protein were compared. Bin width is 2%. Only sequences for which the Waltz server provided a valid output for both the natural and randomized version were used and the total number of valid points is given in each graph.

Figure 4. The average meta-structure correlation map (AMCM) is a meaningful representation of protein space. The AMCM shows the correlation between sequence-derived compactness values and 2nd structure parameters. The two meta-structure parameters are calculated based on the primary sequence of a given protein and given as a protein average. Large compactness values are found for compact 3D structures with dense side-chain interaction networks, whereas small compactness values are
indicative of flexible polypeptide chains devoid of significant stabilizing interactions. 2\textsuperscript{nd} structure parameters are defined as follows: positive: \(\alpha\)-helix, negative: \(\beta\)-strand. Naturally occurring and random sequences are given in black and red, respectively (for details see text).

Figure 5: The structural hererogeniety of protein space is indicated by distinct cluster formation in the average meta-structure correlation map. Amyloidogenic fragments (AFR) are indicated in red, folded proteins (PDB) in blue and IDPs in green (IDP200) or violet (DIS50).
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Figure 1
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Figure 5