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THE PROTEIN TRINITY: STRUCTURE/FUNCTION RELATIONSHIPS THAT INCLUDE INTRINSIC DISORDER

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INTRODUCTION. The current, dominant view relating protein structure to function can be expressed as amino acid sequence \rightarrow 3-D structure \rightarrow function. Currently, more than 200 counter examples in which function depends on nonfolded or incompletely folded regions of protein have been described. Furthermore, reviews on intrinsically disordered proteins are beginning to appear[1,2,3]. In one of these reviews, it is suggested that the existence of proteins with intrinsic protein disorder calls for a re-assessment of the protein structure-function paradigm[3].

METHODOLOGY. From literature and database searches, we collected a set of proteins that are structurally characterized to have regions of disorder or to be wholly disordered under physiological conditions. Once our database was assembled, we used various computational techniques to construct predictors of natural disordered regions (PONDRs).

Protein	LDR/Protein Size	Function
c-Jun	35/331	DNA binding
FlgM	97/97	Protein binding, Channel transport
4E-BP1	118/118	Protein binding, Phosphorylation
Neurofilament H	679/1087	Entropic bristle, Phosphorylation, Gycoslyation
Titin	2174/~33,000	Entropic spring

TABLE 1. Representative samples from our database of over 90 nonhomologous proteins that contain long disordered regions (LDRs) of \geq 30 contiguous amino acids. "LDR/Protein Size" represents the length of the disordered region and the protein respectively. "Function" refers to the function of the disordered region.

Kingdom	# of Proteomes	Disorder
Prokaryia	22	7–33%
Archaea	7	9-37%
Eukaryia	5	36–63%

TABLE 2. Each proteome from the three kingdoms was scanned for disorder using a PONDR predictor. The score for each proteome represents the percentage of the proteins in each proteome

predicted to contain LDRs of \geq 40 contiguous amino acids. The disorder scores for each kingdom are reported as the range of disorder observed throughout all of the proteomes in each kingdom.

CONCLUSIONS. Using the PONDR predictors, we were able to confirm the hypothesis that, just as amino acid sequence determines 3-D structure, it also determines the lack of 3-D structure. Results from these predictors further suggest that the most common type of protein in eukaryotic cells contains both ordered and intrinsically disordered regions. This is likely due to the large amount of signaling proteins in eukaryotic cells. Given that proteins with intrinsically disordered regions are so common, the model relating protein structure to function needs to be reassessed. Here, to accommodate intrinsic disorder, we propose The Protein Trinity (Fig. 1). Following Ptitsyn[4], this proposal envisions three distinct protein states, described approximately as the ordered state, the collapsed state and the extended state. According to The Protein Trinity, any one of these three forms (not just the ordered state) can exist in native proteins.

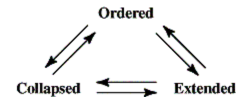


FIGURE 1. The Protein Trinity.

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