

## Scientists discover extensive array of protein folds unexplored in nature

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From the top, the protein name, topology diagram (circles represent  $\alpha$ -helices, triangles and arrows represent  $\beta$ -strands), computationally designed structures, and experimentally determined structures. The topology of NF8-01 forms a knot. Credit: Shintaro Minami, Naohiro Kobayashi, Toshihiko Sugiki, Toshio Nagashima, Toshimichi Fujiwara, Rie Tatsumi-Koga, George Chikenji & Nobuyasu Koga, "Exploration of novel  $\alpha\beta$ -protein folds through de novo design", Nature Structural & Molecular Biology (2023)

A new study has shed new light on the astonishing diversity of protein structures and their folds in nature. Researchers set out to reveal the extent to which nature has explored the vast landscape of possible



protein topologies. The results have unveiled an astounding array of unexplored protein folds, expanding our understanding and uncovering the depth of the protein universe.

This research has been published in the journal *Nature Structural & Molecular Biology*.

Proteins, the building blocks of life, fold into specific three-dimensional structures, enabling them to carry out their biological functions. The three-dimensional structures of proteins are dictated by their amino acid sequences. While experimental techniques have successfully unraveled the structures of numerous proteins over the years, the discovery of new protein folds, defined by the arrangement and connectivity of  $\alpha$ -helices and  $\beta$ -strands, has become increasingly infrequent. This raises the question: How extensive is the protein fold space not explored by nature? In attempts to address this long-standing question, theoretical studies have been conducted; however, experimental validation is lacking.

To address this question, the research team embarked on a study combining theoretical prediction for novel protein folds with experimental validation of their de novo designs.

The research team devised rules based on physical chemistry and protein structure data to theoretically predict possible protein folds. These rules were then employed to predict novel  $\alpha\beta$ -folds, which consist of a four to eight stranded  $\beta$ -sheet, not yet observed in the current Protein Data Bank (PDB). This led to the identification of a total of 12,356 novel folds. The team then attempted to computationally design proteins for the predicted novel folds from scratch to assess the foldability and fidelity of the novel folds.

"We attempted to computationally design proteins with all of the



predicted folds that have a four-stranded  $\beta$ -sheet, including one forming a knot-like structure," said Shintaro Minami, a researcher at Exploratory Research Center on Life and Living Systems (ExCELLS). "When designing proteins, we did not expect all of them, especially knotforming ones, to fold into the structures as anticipated."

The results of experimental testing were surprising. "For all of the folds, the computationally designed protein structures closely matched the experimental structures," said Naohiro Kobayashi, a senior research fellow at RIKEN.

These findings suggest the existence of at least approximately 10,000 unexplored foldable  $\alpha\beta$ -folds, a significant revelation considering only 400  $\alpha\beta$ -folds have been observed in nature. This suggests that many potential folds remain uncharted in the protein folding space.

These results have given rise to several hypotheses about the structure and evolution of proteins. One hypothesis is that proteins may have not been present in biology long enough for all possible folds to have been explored. Another hypothesis is that protein folds in nature are inherently biased due to all life on Earth having descended from a common ancestor.

"Proteins may have evolved by repeatedly reusing specific folds while expressing different functions. If extraterrestrial life does exist, it might be utilizing a different set of protein folds," said George Chikenji, an assistant professor at the Nagoya University.

Proteins are known for their diverse functions, which are generated from the diversity of protein <u>three-dimensional structures</u>. This study has revealed the existence of at least approximately 10,000 uncharted foldable  $\alpha\beta$ -folds in nature



"The design of proteins with these novel folds will lead to an even greater diversity of structures. This would pave the way for the de novo design of functional <u>protein</u> molecules, lead to breakthroughs in drug development, enzyme design, and other areas," said Nobuyasu Koga, a professor at the Exploratory Research Center on Life and Living Systems (ExCELLS), National Institutes of Natural Sciences (NINS).

**More information:** Shintaro Minami et al, Exploration of novel αβprotein folds through de novo design, *Nature Structural & Molecular Biology* (2023). DOI: 10.1038/s41594-023-01029-0

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