Determinants of protein function revealed by combinatorial entropy optimization

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Background. Genome projects are generating a rapidly increasing number of protein sequences, but our knowledge of functional details lags behind. Fortunately, functional constraints in evolution have created information-rich conservation patterns in protein families. If one can decode these patterns, one can derive detailed functional hypotheses. Here, we focus on decoding the patterns of specificity residues. Such residues are conserved in each protein subfamily, but differ between functionally diverse subfamilies.

Results. We present a new algorithm to solve the combinatorial complex problem of identifying specificity residues and, simultaneously, the corresponding optimal division into subfamilies. In our approach, called combinatorial entropy optimization (CEO), we optimize a conservation contrast function over different assignments of proteins to subfamilies. We validate the method by comparing sets of predicted specificity residues with sets of experimentally known functional residues, such as interaction residues observed in threedimensional macromolecular complexes, and get good agreement between prediction and observation.

Conclusion. The method, at http://proteinkeys.org, takes a multiple sequence alignment as input and returns subfamilies and a set of specificity residues. The computed subfamilies may be used, e.g., to assign a likely function to new protein sequences or to choose maximally informative targets for structural genomics projects. The computed specificity residues may be used to design highly specific mutation experiments that test function with minimal side effects; to build sharper and more informative evolutionary trees that more accurately reflect functional relatedness; to predict interactions with proteins; and, to estimate the functional consequences of genetic variation [1]-[7].

References

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