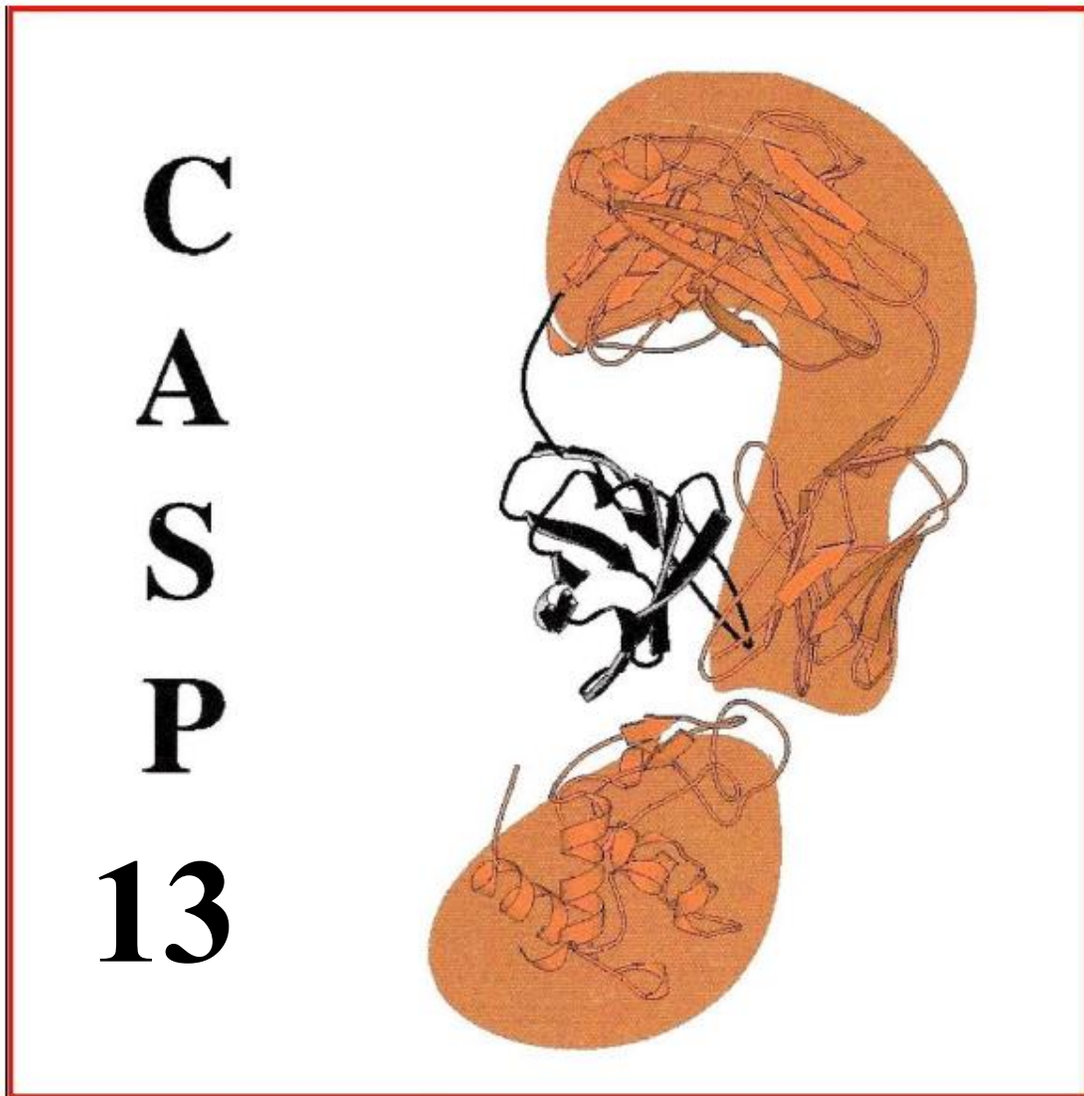


# CRITICAL ASSESSMENT OF TECHNIQUES FOR PROTEIN STRUCTURE PREDICTION



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## De novo structure prediction with deep-learning based scoring

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A7D CASP13 submissions were produced by three variants of an automatic free-modelling structure prediction system relying on scores computed with deep neural networks. Scoring relied on one of two neural networks: a predictor of inter-residue distances and a direct-scoring network. The basic method used a generative neural network for fragment generation for fragment assembly in memory-augmented simulated annealing. Successive rounds of simulated annealing used fragments from the memory. The third method used full-chain score minimization with gradient descent.

### Methods

The systems tested all use multiple sequence alignments (MSA) and profiles generated from HHblits<sup>2</sup> and PSI-BLAST<sup>3</sup>. No templates were used, nor were server predictions. No manual intervention was made except for domain segmentation of T0999 and final decoy ranking in a handful of cases. In protein complexes, each chain was processed independently.

### Scoring

Two neural networks were used for scoring. For the first, a very deep residual convolutional neural network was trained on a non-redundant database of proteins selected from the Protein Data Bank (PDB) to predict the distances between C-beta atoms of different residues, using MSA-based features. With these predictions and a reference distribution, a likelihood score was computed for candidate structures according to the realised distances.

A second deep residual convolutional neural network was trained to directly output a score as a function of structure geometry, MSA-based features and the contact predictions from the first network.

### Domain segmentation

Domain segmentation hypotheses for two or three domains were generated by automatic analysis of the full-chain contact matrix prediction derived from the inter-residue distance prediction. Each domain segmentation hypothesis (as well as full chain without segmentation) was folded independently up to eight times with the domains in each hypothesis being folded independently.

### Fragment assembly

Two approaches were used for structure modelling. The first was based on fragment assembly. For each domain, a DRAW<sup>4</sup> model of backbone torsion angles, trained on the same PDB subset was sampled to generate a set of overlapping 9-residue fragments. Fragments were inserted with simulated annealing using a score based on our distance predictions for the domain hypothesis plus Rosetta's<sup>1</sup> score2 (Variant 1) or the direct structure scoring without Rosetta (Variant 2).

Repeated rounds of simulated annealing were run, using evolutionary hyper-parameter optimization to tune run-length and start temperature, with successive rounds using fragments from the structures generated in previous rounds.

The best-scoring structures from simulated annealing were relaxed using Rosetta fast relax with our inter-residue distance prediction score and Rosetta's full-atom score.

### *Domain assembly*

After domain-level relaxation, for each domain segmentation, full-chain structures were assembled from domain structures with simulated annealing and further relaxed. The best-scoring full-chain structure for each run of each domain segmentation hypothesis for each method was chosen.

### *Direct structure optimization*

An alternative structure modelling approach was used for Variant 3 without any domain segmentation. Here we used gradient descent of a combination score (inter-residue distance prediction score + neural-network-based torsion angle prediction likelihood + score2) to optimize *full chain* structures, parameterised with torsion angles.

### *Decoy selection*

Five ranked structure predictions were submitted for each “all groups” target. Initial submissions used variants 1 & 2 in parallel, but submissions from T0975 on used variants 1 & 3 in parallel. The 5 candidate submissions were the best scoring from among the independent runs of the two different methods, with a bias towards selecting from variants 2 or 3, and manual ranking in a handful of cases.

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