



Deep learning for reconstructing protein structures from cryo-EM density maps: Recent advances and future directions

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Abstract

Cryo-Electron Microscopy (cryo-EM) has emerged as a key technology to determine the structure of proteins, particularly large protein complexes and assemblies in recent years. A key challenge in cryo-EM data analysis is to automatically reconstruct accurate protein structures from cryo-EM density maps. In this review, we briefly overview various deep learning methods for building protein structures from cryo-EM density maps, analyze their impact, and discuss the challenges of preparing high-quality data sets for training deep learning models. Looking into the future, more advanced deep learning models of effectively integrating cryo-EM data with other sources of complementary data such as protein sequences and AlphaFold-predicted structures need to be developed to further advance the field.

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Introduction

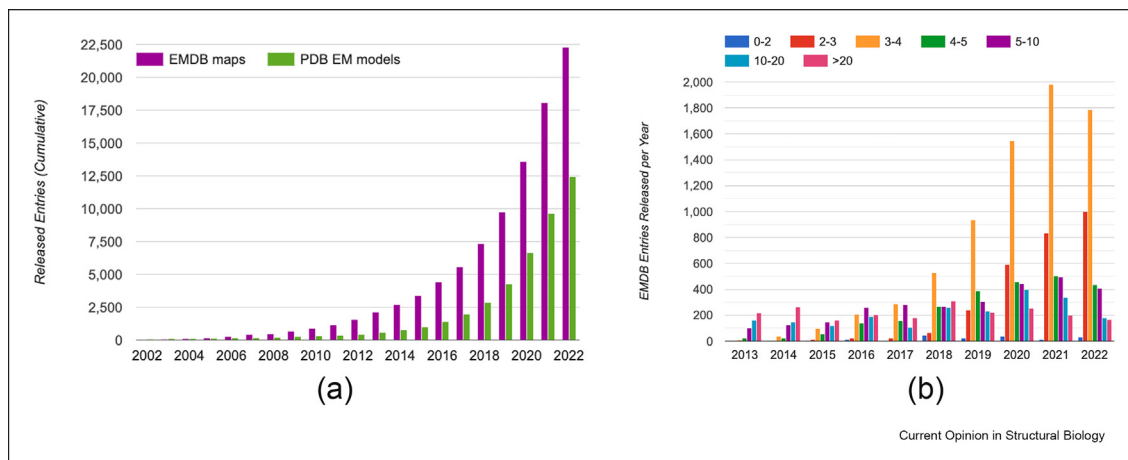
Cryo-EM is revolutionizing structural biology due to its unique capability of determining the structures of large protein complexes and assemblies. The atomic-resolution structure determination for proteins enabled by cryogenic electron microscopy (cryo-EM) [3], allows

us to understand the complex bio-logical processes carried out by proteins as well as to identify potential therapeutic protein targets for drug discovery. However, reconstructing *de novo* protein structures from high-resolution ($\sim 3\text{--}4\text{ \AA}$) cryo-EM density maps, which accounts for a large portion of cryo-EM density maps deposited currently in the EMDB [2], is time-consuming and challenging when homologous template structures for target proteins are not available. For instance, as shown in Figure 1, in the current year 2022, only about 12,500 out of 22,300 density maps of high-resolutions deposited to EMDB have a complete atomic structure available in Protein Data Bank (PDB) [40].

Accurately reconstructing protein structures from cryo-EM maps is a challenging process because the data is often noisy and incomplete and target protein structures can be large and complex. Traditional methods based on energy optimization such as EM-Fold [23], Gorgon [24], Rosetta [25], Pathwalking [26], MAIN-MAST [27,28], VESPER [51], and Phenix [29] have made valuable progress in reconstructing protein structures from cryo-EM density maps. These methods rely on extensive physics-based or statistical potential-based optimization algorithms that require high computational resources. These methods often need manual intervention and trials to extract features from the cryo-EM density maps to obtain accurate reconstruction of protein structure.

A different strategy to automatically determine protein structures from cryo-EM density maps is to use the data-driven machine learning approach [44], a kind of artificial intelligence (AI) technology, to directly learn a mapping from cryo-EM density maps to protein structures from the large amount of known cryo-EM data and their corresponding protein structures (i.e., labels). Early AI methods in the field are based on shallow machine learning techniques such as k-nearest neighbor, support-vector machines, or k-means clustering techniques. These methods such as RENNSH [30], SSELearner [31], and Pathwalking [26] are able to identify only secondary structures or simplified backbone structures and often are unable to achieve the optimal solution.

Figure 1



The growth of cryo-EM density maps and cryo-EM-derived protein structures and the distribution of the resolution of the density maps. The statistics was obtained from EMDDataResource [2], an unified data resource for 3-Dimensional Electron Microscopy (3DEM) on 2022-09-14.

To overcome the challenges of the traditional optimization methods and early machine learning methods, deep learning methods [45] have been developed to automatically reconstruct three-dimensional (3D) protein structures from cryo-EM density maps with significant success in recent years (see Figure 2 for a summary of a general cryo-EM protein structure determination pipeline powered by deep learning). In this article, we review the recent development of deep learning technology in the field, analyze their impacts, investigate the challenging issues in preparing data to train deep learning models, and discuss some new trends to further advance the field.

Deep learning reconstruction of protein structures from cryo-EM density maps

Deep learning, also called deep neural network, is currently the most powerful machine learning method of predicting the properties of an object from the input data describing the object. It has achieved great success in many fields including a recent major breakthrough in predicting protein structure from sequence by AlphaFold [1]. Compared to other machine learning methods, deep learning has a unique capability of extracting informative features for pattern recognition from raw data automatically, making it suitable for reconstructing protein structures from raw density maps in which only a large amount of numbers rather than informative features are available.

It is worth noting that deep learning has been applied to almost all the areas of cryo-EM data analysis [35,32,19–22,38] from sample preparation, particle picking, density map denoising, and to the final step of 3-D structure determination. Due to the space limit,

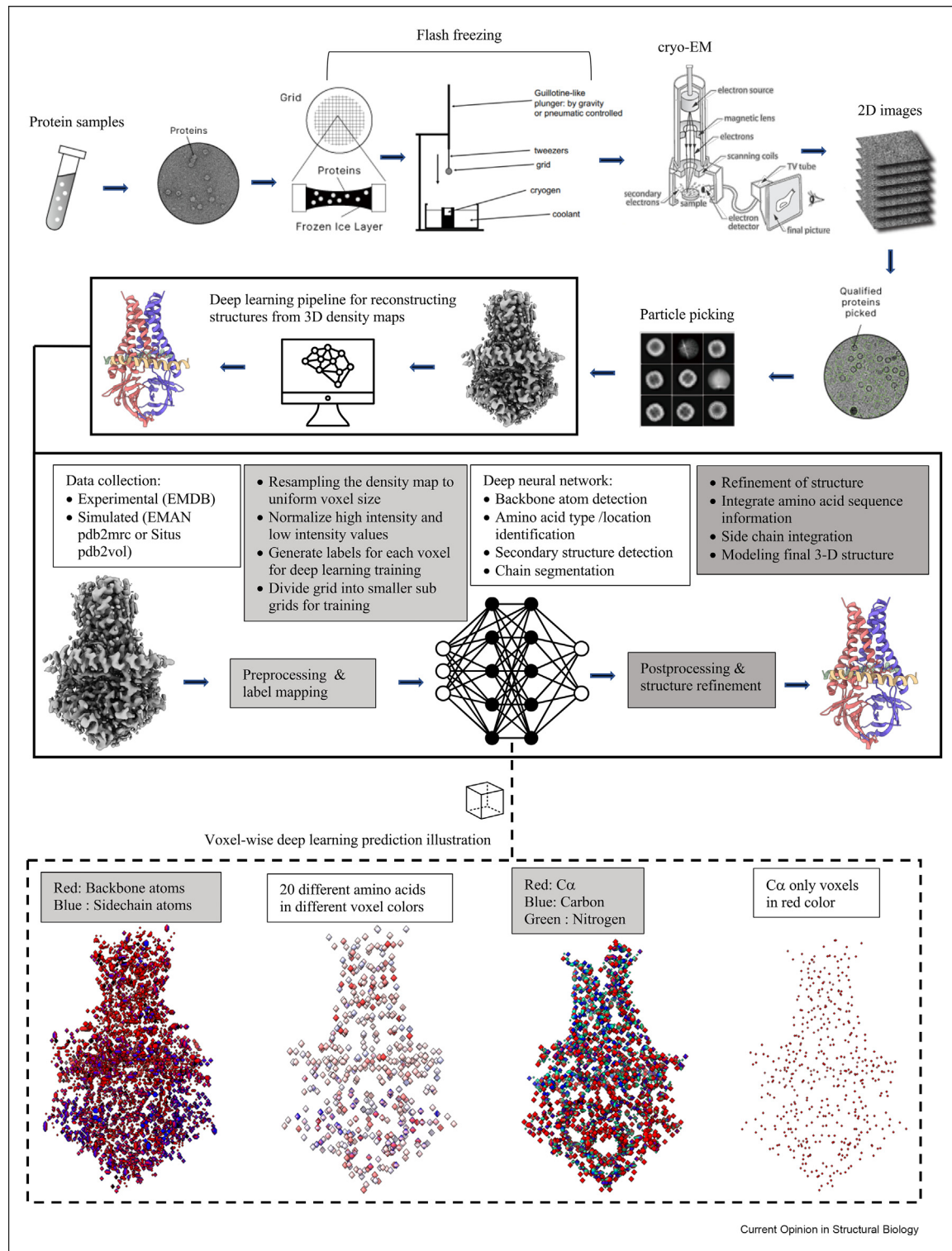
this review is focused on the last step of cryo-EM data analysis - reconstructing protein structures from density maps. The deep learning architectures designed for this task and how to prepare data to train them are discussed in the two subsections below.

Deep learning architectures for reconstructing protein structures from cryo-EM density maps

Deep learning methods for inferring protein structures from cryo-EM density maps can be classified into different categories based on the neural network architectures, for example, convolutional neural network (CNN) [33], U-Net [34,43], graph convolutional network (GCN) [41], and long- and short-term memory network (LSTM) [42] they use and the output (e.g., 3D structure and secondary structure) they generate from density map input. Early deep learning methods aimed to identify secondary structures from low- and medium-resolution density maps [11]. As more and more high-resolution density maps became available [3], recent deep learning methods targeted at directly reconstruct 3D backbone structures (i.e., locations of carbon and nitrogen atoms on the protein backbone) and even full-atom 3D structures (i.e., locations of all/most heavy atoms and amino acid identity/type) from density maps [10,7,14–16]. An example of deep learning reconstruction of protein structure from cryo-EM density map is showed in Figure 3.

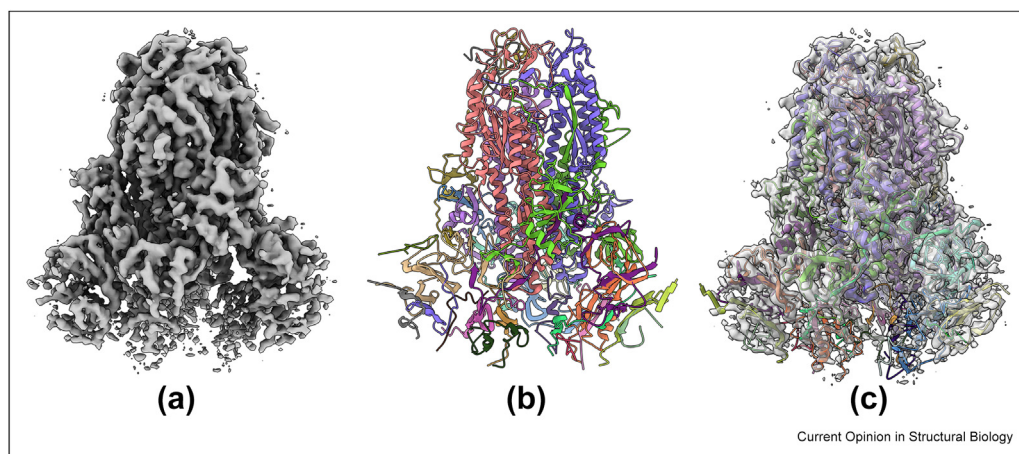
One of the most widely used deep learning architectures of obtaining protein structural information from density maps is convolution neural network (CNN). CNNs use a mathematical operation known as convolution to extract features from spatially organized data such as a 2D-image or 3D density map to predict the properties of the data

Figure 2



A summary of a cryo-EM density map generation and protein structure reconstruction pipeline powered by deep learning. The density map (EMD-22898) illustrated in the figure is for SARS-CoV-2 ORF3a [39]. PDB ID: 7KJR.

Figure 3



An example of reconstructing a structure from the cryo-EM density map of SARS-CoV spike glycoprotein by deep learning. **(a)** Density map of SARS-CoV spike glycoprotein [48] (EMD-6732) in resolution of 3.8 Å at recommended contour level of 0.06 (11.0 σ). **(b)** The structure reconstructed from EMD-6732 by a deep learning method - DeepTracer. The RMSD is 1.023 Å with respect to the ground truth structure (PDB ID: 5XLR). **(c)** The overlay of the density map and reconstructed structure at 0.5 transparency level by UCSF ChimeraX [49].

(e.g., classifying voxels in a density map into amino acid types). Several CNN methods (mostly 3D-CNN architecture) including Generator [7], Emap2sec [8], AAnchor [9], CNN Based [11], Cascaded-CNN [10], and CR-I-TASSER (mostly 3D CNN) [15] have been developed to determine secondary structures [8,11], backbone-/full-atom 3D structures [15,7,9] or both from cryo-EM density maps [10]. Cascaded-CNN is the first deep learning *de novo* method of directly reconstructing 3D structures of proteins from cryo-EM density maps, even though it focuses on building backbone structures. CR-I-TASSER combines the 3D-CNN prediction from cryo-EM maps and an advanced protein structure prediction method - I-TASSER [46] to build full-atom protein structures.

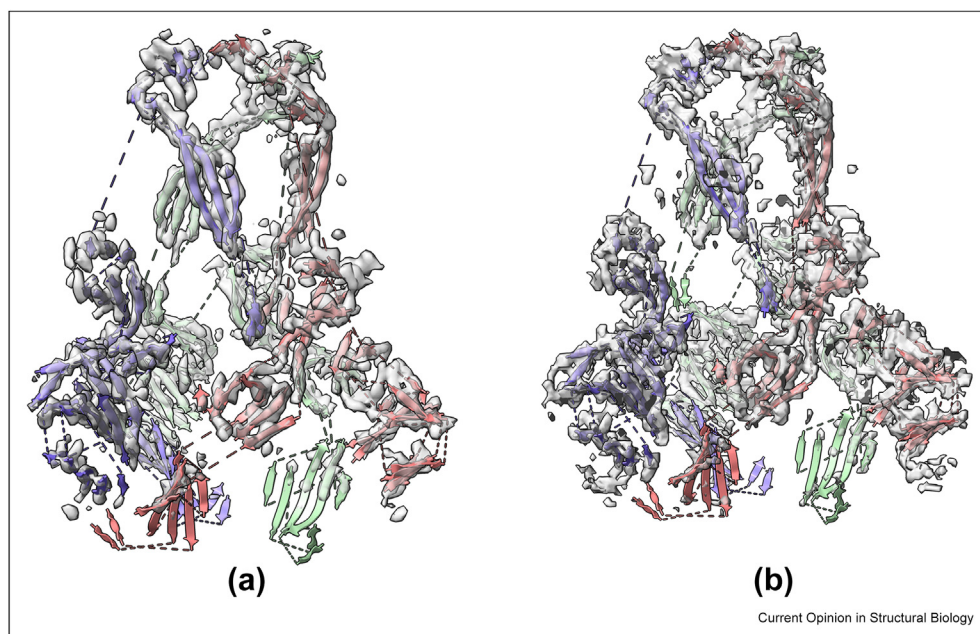
Another widely used convolutional neural network architecture in the field is U-Net [34], originally designed for biomedical image classification and segmentation tasks. U-Net consists of a series of convolution-based down-sampling layers to condense the input images into smaller dimensions and a series of convolution-based up-sampling counterpart layers to reconstruct the data of the same dimension as in the down-sampling process to classify/segment pixels in the input images. Compared to the standard CNN architectures, U-Nets can be more effective in extracting multi-level abstract representations of the data through the down-sampling and up-sampling processes. The 2D U-Net architecture has been generalized to 3D U-Net architectures in Haruspex [12] and EMNUSS [17] to detect secondary structures from cryo-EM density maps (e.g., Figures 4 and 5), and in DeepTracer [13] and EMBuild [16] to reconstruct 3D protein structures from cryo-EM density

maps. DeepTracer has been successfully applied to reconstruct the structures of some SARS-CoV proteins from cryo-EM density maps (e.g., Figure 3).

In addition to CNN and U-Net, other deep learning architectures such as graph convolutional networks (GCN) and long- and short-term memory network (LSTM) have also been used with CNN to reconstruct protein structures from cryo-EM density maps [7]. A summary of different deep learning-based methods, their function (e.g., input and output) and availability is presented in Table 1.

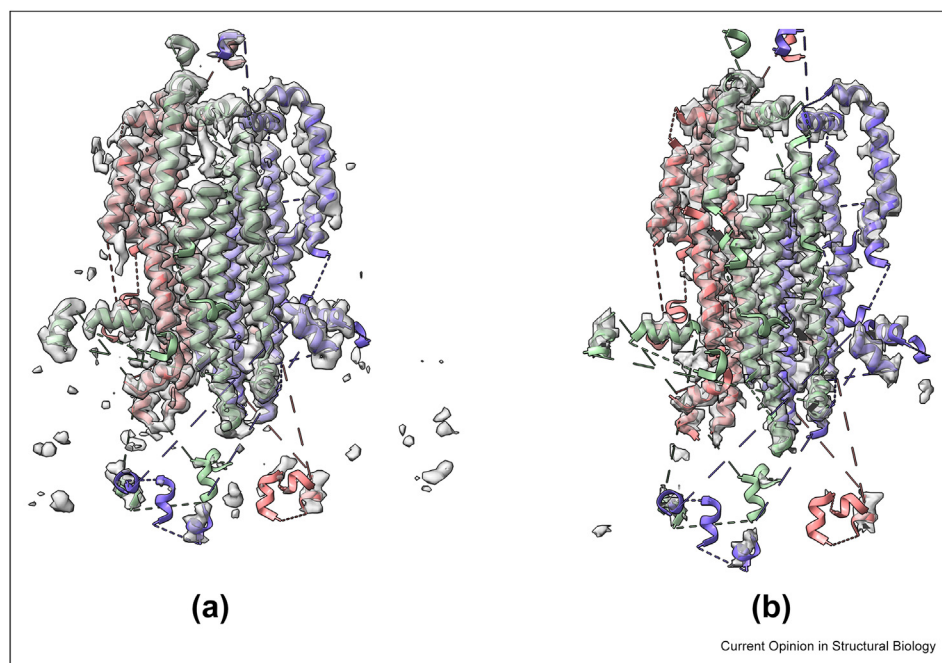
Inspired by the recent breakthrough in developing deep learning methods of predicting protein structures from sequences such as AlphaFold [1] and RoseTTAFold [5], a new trend is to integrate deep learning methods of reconstruct protein structures from cryo-EM density maps with the advanced computational (e.g., deep learning) methods of predicting protein structures from sequences to obtain more accurate structural models. For instance, DeepTracer ID [14] first uses DeepTracer to build an initial structure from cryo-EM density maps and then search the structure against a database of AlphaFold-predicted structures to identify similar structural hits to enhance the reconstructed structure. EMBuild [16] combines the structures reconstructed from cryo-EM maps, AlphaFold-predicted structural models and other protein structural refinement methods to construct accurate structures for protein complexes. ModelAngelo [18] refines the geometry of protein chains by combining information extracted from cryo-EM data, prior knowledge of protein geometries, and amino acid sequence data. DeepProLigand [4]

Figure 4



An example of secondary structure annotation in cryo-EM density map of SARS- CoV spike glycoprotein [48] (EMD-6732) by deep learning. PDB ID: 5XLR. **(a)** Haruspex [12] predicted strands in transparent gray overlapped with deposited PDB structure strands. **(b)** EMNUSS [17] predicted strands in transparent gray overlapped with deposited PDB structure strands.

Figure 5



An example of secondary structure annotation in cryo-EM density map of SARS- CoV spike glycoprotein [48] (EMD-6732) by deep learning. PDB ID: 5XLR. **(a)** Haruspex [12] predicted helices in transparent gray overlapped with deposited PDB structure helices. **(b)** EMNUSS [17] predicted helices in transparent gray overlapped with deposited PDB structure helices.

Table 1

Summary of deep learning based methods for protein structure reconstruction from cryo-EM density maps.

Methods	Architecture	Function	Open source
Structure Generator [7]	3-D CNN, GCN, Bidirectional LSTM	First use 3-D CNN to identify amino acids and their rotameric identities in an EM map and then GCN and LSTM to build protein structures	✓
Emap2sec [8]	3-D CNN	Take voxel cubes as input to identify secondary structures of protein	✓
AAnchor [9]	3-D CNN	Take in voxel cubes to identify amino acid types and locations	✓
A CNN Based Method [11]	3-D CNN	Take in voxel cubes to detect secondary structures of protein from background	×
CascadedCNN [10]	Cascaded 3-D CNN	Take in voxel cubes to identify Cα atoms of protein backbone and secondary structures to generate 3D protein structures	✓
Haruspex [12]	3-D U-Net	Take in voxel cubes to predict the probabilities of 4 different classes; α -helix, β -sheet, nucleotide, or unassigned to assign secondary structures	✓
DeepTracer [13]	3-D U-Net	Take in voxel cubes to identify the location of backbone atoms, secondary structures and amino acid types simultaneously to build 3D structure	×
DeepTracer ID [14]	DeepTracer (3-D U-Net) and pre-calculated AlphaFold2 protein library	Use DeepTracer to generate an initial 3D protein structure to search AlphaFold2DB to identify similar structural hits for refinement	×
CR-I-TASSER [15]	3-D CNN, I-TASSER	Predict C α using 3-D CNN for selecting structural templates for I-TASSER to generate 3D protein structure	✓
EMBuild [16]	3-D U-Net++, AlphaFold	Integrate AlphaFold structure prediction, FFT-based global fitting, domain-based semi-flexible refinement, and graph-based iterative assembling with main-chain probability maps predicted by U-Net++ to build 3D protein structure	✓
EMNUSS [17]	3-D U-Net++	Take in voxel cubes to identify secondary structures of protein	✓
ModelAngelo [18]	Graph Neural Network	Refines geometry of protein chains and classifies amino acid for each nodes	×

integrates the protein structural models reconstructed from cryo-EM density maps by DeepTracer with the known template structures containing ligands to model protein-ligand interaction, which was ranked first in the **ligand prediction in 2021 EMDDataResource Ligand Model Challenge**.

Data preparation for training deep learning methods to reconstructing protein structures from cryo-EM density maps

Cryo-EM density map data collection

Collecting a sufficient amount of high-quality data to train and test deep learning models is critical for any deep learning task. The common way to acquire the *experimental cryo-EM density maps* is through the Electron Microscopy Data Bank [2]. An alternative approach employed by some methods such as **Cascaded-CNN** [10] and SSELearner [31] is to simulate the density map from the PDB protein structure. Cascaded-CNN applies *pdb2mrc* from EMAN2 package [50], and VESPER uses *pdb2vol* from Situs package [52] to generate the *simulated maps*. **However, simulated maps**

lack complex noise, missing density values, and experimental artifacts which can arise from particle alignment errors, interaction of electron beam with the atoms, or movement of atoms during image capture. Therefore, the deep learning models trained on simulated maps may not work as expected on very noisy experimental data. To address the problem, CR-I-TASSER, EMNUSS and Emap2sec employs a hybrid training approach that uses both simulated maps and experimental maps in the training and validation process.

Training data preprocessing

Prior to using the cryo-EM density map to train deep learning models, it is generally necessary to normalize and standardize the data to make them suitable for deep learning as shown by **Cascaded-CNN and DeepTracer, which perform data grid resampling, density value normalization, and grid division**. These preprocessing steps ensure the uniformity among density maps and help deep learning models to extract features and recognize patterns more easily. During the grid division, the 3D cryo-EM is split into the cubes of a specific size

(e.g., $64 \times 64 \times 64 \text{ \AA}^3$ by Cascaded-CNN and DeepTracer, $50 \times 50 \times 50 \text{ \AA}^3$ by CR-I-TASSER, $40 \times 40 \times 40 \text{ \AA}^3$ by Haruspex, and $11 \times 11 \times 11 \text{ \AA}^3$ by Emap2sec and AAnchor). Each of these cubes is then processed by the deep learning method to classify the voxels into the targeted classes such as amino acid types (identities) and secondary structures.

Future directions

Deep learning has made a significant impact on protein structure reconstruction from cryo-EM density maps. However, the field is still in the early stage of development. The latest deep learning technology such as graph neural networks [53] and attention mechanisms [47] have not been extensively used in the field. While CNNs and U-Nets based on convolution are currently the most used methods for structure reconstruction, they have some short-coming for 3D structural modeling. CNNs are translation-equivariant, but not fully rotation invariant that is desirable for 3D structure analysis. Moreover, the convolution mechanism propagates message in the constrained local receptive field, which is not as effective as the attention mechanism [47] that can leverage all the input information by automatically weighting the input features according to their relevance as demonstrated by the remarkable success of AlphaFold2 in protein structure prediction. More sophisticated deep learning models like attention-based Transformer models [36], 3D-equivariant graph neural networks [37], and AlphaFold2-like deep learning models need to be developed to better use cryo-EM data to improve reconstruction accuracy.

Another important direction is to use deep learning to integrate cryo-EM data with multiple other sources of complementary data such as protein structural models predicted from sequences, structural templates in the Protein Data Bank (PDB), and protein sequences to more accurately reconstruct protein structures from noisy density maps that often miss the density values of some atoms. The current integration process is limited to shallow data combination. For instance, DeepTracer ID uses AlphaFold models to refine the structural models predicted from structural models reconstructed from deep learning. More comprehensive, end-to-end deep learning models to combine multiple sources of data to generate accurate final protein structures can be developed to automatically and accurately reconstruct protein structures from the data.

Moreover, it is important to integrate cryo-EM based deep learning methods of reconstructing protein structures with the advanced methods developed in the field of protein structure prediction. The structural models directly reconstructed from cryo-EM data by deep learning generally have correct overall topology, but the reconstructed models may not satisfy physico-chemical restraints such as bond length and bond angles and not

have all the molecular details (e.g., the precise location of all side chain atoms) [10,4]. Linking the atoms of amino acids identified from the density maps into full peptide chains consistent with protein sequences and physical-chemical restraints is still challenging. However, the modeling techniques such as protein structure refinement and molecular dynamics to fix these problems have been established for protein structure prediction [1]. Some methods such as CR-I-TASSER have started to integrate the two kinds of technologies. More synergistic integration of the two are needed to generate high-quality realistic protein structures from cryo-EM data.

The development of high-quality deep learning models to reconstruct protein structures from cryo-EM density maps critically depends on the availability of sufficient high-quality training data. Although experimental cryo-EM data and its related ground truth structure are freely accessible through EMDB [2] and RCSB PDB [40], these datasets still need to be pre-processed and labeled before they can be used for deep learning training. Curating a large amount of high-quality training and test data is challenging and time consuming, but often receives little attention. Currently, there are few well-curated experimental cryo-EM data sets publicly available for training and evaluating deep learning models in the field. Therefore, more effort needs to be devoted to creating such data sets and make them to publicly available for the community to use.

Conclusion

A number of useful deep learning models have been developed to reconstruct protein structures from cryo-EM density maps, demonstrating deep learning is a promising technology to further push the frontier of applying cryo-EM technology to determine protein structures. As the deep learning field is evolving very fast, many more state-of-the-art deep learning architectures (e.g., AlphaFold2-like models and transformers) have yet to be applied to further advance the emerging field. More sophisticated deep learning methods need to be developed to seamlessly integrate cryo-EM data with other complementary data such as predicted protein structures, protein sequences, and template structures to further improve cryo-EM-based structure determination. A synergistic integration of cryo-EM based protein structure determination techniques and latest protein structure prediction techniques is also important for generating highly accurate native-like protein structures

To speed up the development, more effort is need to create a large amount of high-quality cryo-EM training and test data for the community to use.

Conflict of interest statement

Nothing declared.

Data availability

Experimental cryo-EM data and its related ground truth structure are freely accessible through EMDB and RCSB PDB.

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* of special interest

** of outstanding interest

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