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An increasing number of density maps of macromolecular structures, including proteins and protein and DNA/RNA complexes, have been determined by cryo-electron microscopy (cryo-EM). Although maps at a near-atomic resolution are routinely reported, there are still substantial fractions of maps determined at intermediate (~ 4 Å) or lower resolutions, where extracting structure information is difficult. Considering limited approaches developed for maps with protein-nucleic acid complexes under intermediate resolution, we report a new computational method, Emap2sec+, which identifies DNA or RNA as well as the secondary structures of proteins in cryo-EM maps of 5 to 10 Å resolution (Nat. Commn, 2021). Emap2sec+ uses the 3D deep residual convolutional neural network (3D-ResNet) as its core of the architecture. Emap2sec+ assigns structural labels with associated probabilities at each voxel in a cryo-EM map, which can help structure modeling in an EM map. It adopts a two-phase stacked neural network architecture, where predictions in the first phase are further smoothed in the subsequent second phase by incorporating the context of neighboring voxels. Emap2sec+ showed stable and high assignment accuracy for nucleotides in low-resolution maps. Detection accuracy is remarkably stable for binary classification of protein and nucleotides even in the maps of a low resolution of 8-10 Å. The code is available at https://github.com/kiharalab/Emap2secPlus together with other cryo-EM software https://kiharalab.org/emsuites.

Figure 1