Generating Tertiary Protein Structures via an Interpretative Variational Autoencoder

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Much scientific enquiry across disciplines is founded upon a mechanistic treatment of dynamic systems that ties form to function. A highly visible instance of this is in molecular biology, where an important goal is to determine functionally-relevant forms/structures that a protein molecule employs to interact with molecular partners in the living cell. This goal is typically pursued under the umbrella of stochastic optimization with algorithms that optimize a scoring function. Research repeatedly shows that current scoring function, though steadily improving, correlate weakly with molecular activity. Inspired by recent momentum in generative deep learning, this paper proposes and evaluates an alternative approach to generating functionally-relevant three-dimensional structures of a protein. Though typically deep generative models struggle with highly-structured data, the work presented here circumvents this challenge via graph-generative models. A comprehensive evaluation of several deep architectures shows the promise of generative models in directly revealing the latent space for sampling novel tertiary structures, as well as in highlighting axes/factors that carry structural meaning and open the black box often associated with deep models. The work presented here is a first step towards interpretative, deep generative models becoming viable and informative complementary approaches to protein structure prediction.

Additional Key Words and Phrases: Deep generative models, variational autoencoder, disentanglemment, protein structure prediction

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1 INTRODUCTION

Decades of scientific enquiry across disciplines have demonstrated just how fundamental form is to function [7]. A central question that arises in various scientific domains is how to effectively explore the space of all possible forms of a dynamic system to uncover those that satisfy the constraints needed for the system to be active/functional. In computational structural biology we find a most visible instantiation of this question in the problem of *de-novo* (or template-free) protein structure prediction (PSP). PSP seeks to determine one or more biologically-active/native forms/structures of a protein molecule from knowledge of its chemical composition (the sequence of amino acids bonded to one another to form a protein chain) [32].

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PSP has a natural and elegant formulation as an optimization problem [37] and is routinely tackled with stochastic optimization algorithms that are tasked with exploring a vast and high-dimensional structure space. These algorithms sample one structure at a time, biased by an expert-designed scoring/energy function that scores sampled structures based on the likelihood of these strutures being biologically-active/native [46]. While great progress has been made in stochastic optimization for PSP, there are fundamental challenges on how to achieve a resource-aware balance between exploration (exploring more of the structure space so as not to miss novel structures) and exploitation (improving structures so as to reach local optima of the scoring function) in vast and high-dimensional search spaces and dealing with inherently-inaccurate scoring functions that often drive towards score-optimal but non-native structures [28].

Stochastic optimization frameworks that conduct a biased (by the scoring function) exploration of a protein's structure space, do not learn not to generate structures that are unfit according to the scoring function. Efforts to inject a learning component by Shehu and others struggle with how to best connect the structure generation mechanism/engine in these frameworks with the evaluation engine [40]. A machine learning (ML) framework would seemingly provide a natural setting; however, while discriminative (ML) models can in principle be built to evaluate tertiary structures, and a growing body of research is showing their promise [10, 12, 13, 33], effective generative ML models for tertiary protein structures have so far been elusive.

The majority of work on leveraging deep neural networks (NNs) for problems related to PSP has been on the prediction of distance or contact maps. The latter are alternative representations of tertiary molecular structures that dispense with storing the Cartesian coordinates of atoms in a molecule and instead record only pairwise distances (or, as in contact maps, 0/1 bits to indicate a distance above or not above a threshold). Work on contact map (similarly, distance matrix) prediction focuses on the following setting: given the amino-acid sequence of a protein molecule of interest, predict one contact map that represents the most likely native structure of the target protein. The predicted contact map (or distance matrix) can be utilized to recover a compatible tertiary structure via optimization algorithms that treat the predicted contacts/distances as restraints [1, 50].

Early work on prediction of a contact or distance matrix from a given amino-acid sequence [29] employed bidirectional recursive NNs (BRNNs). RaptorX-Contact started a thread of productive research on residual convolutional NNs (CNNs) [52]. DNNCON2 leveraged a two-stage CNN [2]. DeepContact deepened the framework via a 9-layer residual CNN with 32 filters [35]. DeepCOV alleviated some of the demands for evolutionary information [23]. PconsC4 further limited input features and significantly shortens prediction time [41]. SPOT-Contact extended RaptorX-Contact by adding a 2D-RNN stage downstream of the CNN [19]. TripletRes put together four CNNs trained end-to-end [34].

It is worth emphasizing that the above NNs for contact or pairwise distance prediction predict one instance from a given amino-acid sequence. These models do not generate multiple contact maps or multiple distance matrices to give any view of the diversity of physically-realistic (and possibly functionally-relevant) structures in the structure space of a given protein. The objective of these models is to leverage the strong bias in the input features (which include amino-acid sequence, physico-chemical information of amino acids, co-evolutionary information, secondary structures, solvent accessibility, and other more sophisticated features) to get the model to output *one* prediction that is most likely to represent the native structure. While the performance of these models varies [49] and is beyond the scope of this paper, this body of work does not leverage generative deep learning and is not focused on generating a distribution of tertiary structures for a given amino-acid sequence (whether directly in Cartesian space or indirectly through the intermediate representation of contact maps or pairwise distance matrices).

Work in our laboratory has integrated some of these contact prediction models, such as RaptorX-Contact, in optimization-based structure generation engines to evaluate the quality of a sampled tertiary structure in lieu of or in

addition to a scoring/energy function [56]. AlphaFold can also be seen as a similar effort, where a deep NN is used to predict a pairwise distance matrix for a given amino-acid sequence and then predicted distances are encapsulated in a penalty-based scoring function to guide a gradient descent-based optimization algorithm assembling fragments into tertiary structures [45].

While it remains more challenging to generate a distribution of tertiary structures (whether directly or indirectly through the concept of contact maps or distance matrices), some preliminary efforts have been made [4, 5, 22, 43]. Work in [5] investigates the promise of such models for protein de-novo design. Other work that focuses on PSP remains preliminary and lack of rigorous evaluation on metrics of interest in PSP and, more generally, protein modeling does not expose what these models have actually learned. – need to provide more detail here to differentiate these efforts.

Inspired by recent momentum in generative deep learning, we propose here a novel approach based on generative, adversarial deep learning. Leveraging recent opportunities in graph generative learning, we put forth a disentanglement-enhanced contact variational autoencoder (VAE) to which we refer as DECO-VAE from now on. As any other deep neural network, DECO-VAE learns from data, which we obtain from an initial population generated by the state-of-the-art stochastic optimization engine Rosetta [31]. As a VAE, DECO-VAE learns from this data the latent space encapsulating the distribution and then directly generates more data (tertiary structures) from the latent space.

DECO-VAE addresses several hurdles that stand in the way of generative models for highly-structured data. First, each tertiary structure is represented as a graph via the concept of the contact graph, and DECO-VAE learns the latent space of such graphs, directly sampling novel contact graphs from this space. In this first version, readily-available 3D reconstruction webservers, such as CONFOLD [1], are used to obtain tertiary structures corresponding to DECO-VAE-generated graphs. Second, DECO-VAE addresses the issue of interpretability via the disentanglement mechanism. In one of several insightful analyses presented in this paper, we demonstrate how DECO-VAE is more powerful than a version with no disentanglement mechanism, CO-VAE, as well as other deep architectures. We show that DECO-VAE generates a distribution that does not simply reproduce the input distribution but instead contains novel and better-quality structures. A thorough comparison with several alternative generative models is also presented. In addition, the disentanglement mechanism integrated in DECO-VAE elucidates what exactly about the tertiary structures the "axes"/factors of the learned latent space control, thus opening the black box that is often associated with deep neural networks.

The paper proceeds as follows. First, we present a summary of related work. DECO-VAE, the input data, and other components of our approach (reconstruction of tertiary structures and various analyses and metrics) are then described in detail. The evaluation of the obtained distribution and the interpretability of DECO-VAE is presented next. The paper concludes with a summary of remaining challenges and future opportunities.

2 BACKGROUND AND RELATED WORK

The concept of a contact graph or a contact map has long been leveraged in molecular biology, particularly in protein modeling [53]. Briefly, the contact map is an alternatively, compressed representation of a tertiary molecular structure. We can think of it as an adjacenncy matrix, where the entry [i,j] encodes whether atoms at positions i,j are spatially proximate; the latter uses threshholds, which typically vary in [8,10]Åin computational biology literature. Typically, the main carbon atoms of each amino acid (the CA atoms) are considered. Some works focus instead on the carbon beta atoms (connecting the side chain atoms to the backbone atoms of an amino acid). In this paper, we take the convention of CA-based contact maps.

Interest in predicting contacts from a given amino-acid sequence has a long history in protein structure prediction, as it was thought to be perhaps less challenging that predicting other representations (such as Cartesian coordinates). Many machine learning approaches have been used, including support vector machines [10, 53], random forest [33], neural networks [15, 18, 42, 48, 51], and deep learning [12, 13], with varied success. These approaches struggle with how best to capture the inherent structure in the input training data and how to discover the pertinent latent space.

More recent methods leverage the concept of the latent space via variational autoencoders [6, 36, 39]. Specifically, Ma et al.[39] develops a deep learning based approach that uses convolutions and a variational autoencoder (a CVAE) to cluster tertiary structures obtained from folding simulations and discover intermediate states from the folding pathways. Anand et al. [6] applies generative adversarial networks (GANs) for *de-novo* protein design. The authors encode protein structures in terms of pairwise distances between CA atoms, eliminating the need for the generative model to learn translational and rotational symmetries. Livi et al. [36] is the first to treat the contact map as a graph and propose a generative model by learning the probability distribution of the existence of a link/contact given the distance between two nodes.

To the best of our knowledge, no deep graph-generative models have been applied to contact map generation. The DECO-VAE we present in this paper is the first to do so. The method leverages developments in graph-generative neural networks (GNNs). Most of the existing GNN-based graph generation for general graphs have been proposed in the last two years and are based on VAEs [44, 47] or GANs [8]. Most of these approaches generate nodes and edges sequentially to form a whole graph, leading to issues of sensitivity to generation order and high time costs for large graphs. Specifically, GraphVAE [47] represents each graph by its adjacent matrix and feature vector and utilizes the VAE model to learn the distribution of the graphs conditioned on a latent representation at the graph level. Graphite [16] and VGAE [27] encode the nodes of each graph into node-level embeddings and predict the links between each pair of nodes to generate a graph. In contrast, GraphRNN [55] builds an autoregressive generative model on these sequences with a long short-term memory (LSTM) model and shows good scalability.

These current graph generation models have two drawbacks: (i) the encoder and decoder architecture are not powerful to handle real-world graphs with complex structure relationships; (ii) they are black boxes, lacking interpretability. In the biology domain, we need to understand the features we learn from the observing data and how we generate the contact maps.

In this paper, we additionally address the issue of interpretability via the concept of disentanglement. Recently, a variety of disentangled representation learning algorithms have been proposed for VAEs. Assuming that the data is generated from independent factors of variation (e.g., object orientation and lighting conditions in images of objects), disentanglement as a meta-prior encourages these factors to be captured by different independent variables in the representation. Learning via disentanglement should result in a concise abstract representation of the data useful for a variety of downstream tasks and promises improved sample efficiency. This has motivated a search for ways of learning disentangled representations, where perturbations of an individual dimension of the latent code z perturb the corresponding x in an interpretable manner. A well-known example is the $\beta - VAE[20]$. This has prompted a number of approaches that modify the VAE objective by adding, removing, or altering the weight of individual terms [3, 9, 14, 24, 30, 38, 59].

Marrying recent developments in disparate communities, we present here a VAE that generates contact graphs (of tertiary protein structures) and learns disentangled representations that allow interpreting how the learned latent factors control changes at the contact graph level. By utilizing known 3D reconstruction tools, we additionally show

how the factors control changes in Cartesian space. The following Section provides further details into the DECO-VAE and other models we design and assess for their ability to generate physically-realistic tertiary protein structures.

3 RESULT

Comparison Methods

We utilize three deep graph generation models to validate the superiority of our proposed model. VGAE [27] is a framework for unsupervised learning on graph-structured data based on the variational auto-encoder (VAE). This model makes use of latent variables and is capable of learning interpretable latent representations for undirected graphs. Graphite [16] is an algorithmic framework for unsupervised learning of representations over nodes in large graphs using deep latent variable generative models. This model parameterizes variational autoencoders (VAE) with graph neural networks, and uses a novel iterative graph refinement strategy inspired by low-rank approximations for decoding. GraphRNN [55] is a deep autoregressive model to approximate any distribution of graphs with minimal assumptions about their structure. GraphRNN learns to generate graphs by training on a representative set of graphs and decomposes the graph generation process into a sequence of node and edge formations, conditioned on the graph structure generated so far.

Training and Testing Dataset

The evaluation is carried out on 13 proteins of varying lengths (53 to 146 amino acids long) and folds (α , β , $\alpha + \beta$, and coil) that are used as a benchmark to evaluate structure prediction methods [57, 58]. On each protein, given its amino-acid sequence in fasta format, we run the Rosetta AbInitio protocol available in the Rosetta suite [31] to obtain an ensemble of 50,000 - 60,000 structures. For each structure, we only retain the central carbon (CA) atom for each amino acid (effectively discarding the other atoms). Each such CA-only structure is converted into a contact graph, as described above. The contact graph dataset of each protein is split into a training and testing data set in a x:y split. The deep models described above are separately trained on each protein. An overview of Disentanglement-enhanced Contact-VAE (DECO-VAE) is illustrated in Fig.6. By obeserving many decoys contact maps for a protein sequence, it learns the distribution of these decoys and generate many more decoys for that protein sequence. DECO-VAE has two main properties: contact generation and interpretation. The contact generation is an implementation of a deep graph varitional auto-encoder. The graph encoder learns the distribution of the decoys of a certain sequence and generate new decoys based on a low dimension latent code. And then the contact decoys can be recovered into 3D structure by the existing methods. The contact interpretation ensures each element in the latent representation has a control on an interpretable factor of the generated protein structure. By changing the element value continuously in the latent code, we can control the different properties of the generated protein structure. The proposed method also has a variant which does not consider the disentanglement but only target on the generation of contact maps, named as Contac-VAE (CO-VAE). The CO-VAE has the same structure with the DECO-VAE, except the objective function when training a generative model. The details of the methods are illustrated in the Section of Methods

Comparing the generated contact maps with the native one

The proposed methods have been evaluated on 13 datasets, each of which corresponds to a protein. For each protein that is a sequence if amino acids, we use thousands of its contact map decoys to train the CO-VAE and DCO-VAE model. Then based on this well-trained model, the same number of contact maps are generated and compared with the only

native one. For each generated contact map and native one, we compute the precision, recall, coverage and and F1 score between them, as shown in Table 1. In addition, the performance of our methods are also compared with the other graph generation methods in machine learning domain (Graphite [16], GVAE [27], GraphRNN [55]). The algorithm and parameter settings of the proposed CO-VAE, DCO-VAE model and the comparison methods are described in details in the Section of Methods.

As shown in Table 1, the graphs generated by CO-VAE and DCO-VAE both have very high F1_score, and the coverage and recall are all around 0.34, which outperforms the other methods by a large margin. Specifically, our precision is 0.67 and 0.71 on average, over 53.4% and 56.7% higher than the highest performance of comparison methods; our coverage is 0.55 and 0.58 on average, over 12.7% and 24.8% higher than the highest performance of comparison methods; our recall is 0.57 and 0.59 on average, over 12.2% and 15.2% higher than the highest performance of comparison methods; our F1_score is 0.61 and 0.64 on average, over 52.4% and 54.7% higher than the highest performance of comparison methods; Moreover, the proposed methods (CO-VAE and DCO-VAE) obtain the best performance in four metrics in almost 76% of the proteins. This indicates that the P-GraphVAE and PD-GraphVAE truly learn the underlined distribution for the contact map decoys for the amino acid sequence. The good performance of the proposed methods rely on its special architecture, where each amino acid has its unique parameters in extracting features from its contacts and the structure similarity is considered among different amino acids in generation process.

Table 1. Evaluation of Generated Contact Maps by Comparing to Native One for different proteins

Protein	Model	Precision	Coverage	Recall	F1_score	Protein	Model	Precision	Coverage	Recall	F1_score
1DTJA	Graphite	0.1320	0.5000	0.5000	0.2031		Graphite	0.1526	0.4893	0.5002	0.2338
	GVĀE	0.1390	0.5109	0.5000	0.2101		GVĀE	0.1526	0.4892	0.5001	0.2339
	GraphRNN	0.3153	0.1301	0.1301	0.1733	1DTDB	GraphRNN	0.2974	0.2696	0.2755	0.2858
	CO-VAE	0.7432	0.3310	0.3312	0.4607	İ	CO-VAE	0.6144	0.5961	0.6093	0.6118
	DCO-VAE	0.7429	0.3347	0.3346	0.4618		DCO-VAE	0.6136	0.6014	0.6148	0.6142
1AIL	Graphite	0.1277	0.4912	0.4994	0.2034		Graphite	0.0748	0.4976	0.5004	0.1302
	GVĀE	0.1276	0.4911	0.4992	0.2032		GVĀE	0.0747	0.4971	0.4998	0.1300
	GraphRNN	0.3900	0.3718	0.37802	0.3836	2H5ND	GraphRNN	0.2846	0.2829	0.2845	0.2844
	CO-VAE	0.8244	0.8391	0.8531	0.8385		CO-VAE	0.7127	0.6393	0.6428	0.6759
	DCO-VAE	0.8231	0.8393	0.8533	0.8379	İ	DCO-VAE	0.7248	0.6354	0.6389	0.6791
1ALY	Graphite	0.0684	0.4981	0.5002	0.1203	1HHP	Graphite	0.0902	0.4979	0.5003	0.1529
	GVÂE	0.0683	0.4989	0.5011	0.1206		GVÂE	0.0903	0.4978	0.5002	0.1529
	GraphRNN	0.2356	0.1979	0.1987	0.2155		GraphRNN	0.2804	0.2582	0.2594	0.2693
	CO-VAE	0.8034	0.4463	0.4482	0.5754	İ	CO-VAE	0.6494	0.6612	0.6643	0.6567
	DCO-VAE	0.8368	0.4408	0.4427	0.5791		DCO-VAE	0.6544	0.6591	0.6622	0.6583
	Graphite	0.1065	0.4963	0.4995	0.1756	1BO9	Graphite	0.1566	0.4809	0.4994	0.2384
	GVÂE	0.1066	0.4964	0.4996	0.1757		GVÂE	0.1568	0.4815	0.5001	0.2387
1AOY	GraphRNN	0.3717	0.2959	0.2978	0.3299		GraphRNN	0.3659	0.3232	0.3356	0.3494
	CO-VAE	0.7620	0.7363	0.7409	0.7511	~	CO-VAE	0.4658	0.4451	0.4622	0.4639
	DCO-VAE	0.8109	0.7299	0.7345	0.7708		DCO-VAE	0.4638	0.4363	0.4531	0.4584
1C8CA	Graphite	0.1322	0.4907	0.5002	0.2091		Graphite	0.1234	0.4843	0.4997	0.1980
	GVÂE	0.1319	0.4900	0.4995	0.2088		GVÂE	0.1234	0.4846	0.4999	0.1979
	GraphRNN	0.3381	0.3065	0.3124	0.3239	1CC5	GraphRNN	0.3489	0.2733	0.2820	0.3117
	CO-VAE	0.6513	0.6707	0.6837	0.6670		CO-VAE	0.8972	0.5878	0.6064	0.7237
	DCO-VAE	0.6538	0.6695	0.6824	0.6678		DCO-VAE	0.8965	0.5864	0.6051	0.7225
	Graphite	0.1500	0.4933	0.5000	0.2308		Graphite	0.1534	0.4891	0.4996	0.2348
1HZ6A	GVÂE	0.1500	0.4933	0.5000	0.2307	1ISUA	GVÂE	0.1537	0.4898	0.5003	0.2351
	GraphRNN	0.1849	0.3543	0.3592	0.2441		GraphRNN	0.2837	0.2546	0.2601	0.2711
	CO-VAE	0.5960	0.4519	0.4581	0.5179		CO-VAE	0.7542	0.4738	0.4839	0.5886
	DCO-VAE	0.6067	0.4545	0.4607	0.5237		DCO-VAE	0.8435	0.4738	0.4839	0.6151
1SAP	Graphite	0.1329	0.4958	0.4993	0.2099		Graphite	0.1055	0.4945	0.5008	0.1742
	GVÅE	0.1332	0.4964	0.5000	0.2103		GVÅE	0.1054	0.4939	0.5001	0.1740
	GraphRNN	0.3021	0.3178	0.3202	0.3107	1TIG	GraphRNN	0.2787	0.2687	0.2721	0.2752
	CO-VAE	0.3021	0.3179	0.3201	0.3106		CO-VAE	0.5797	0.5691	0.5764	0.5780
	DCO-VAE	0.6681	0.6838	0.6888	0.6783	l	DCO-VAE	0.5799	0.5689	0.5761	0.5780

Percentage of native and non-native contacts in a structure

For each of the generated contact map, we calculate the percentage of its contacts that are indeed native contacts, namely those contacts in a native structure. Specifically: Given contacts for a structure (each model) and the known

native structure, we calculate the percentage of contacts that are in the native structure that are also in the reconstructed structure. Percentage for each reconstructed structure (model) is calculated for each protein per each method and then an average over all reconstructed structures is reported per protein per method. In addition, we also calculate the percentage of non-native contacts: the contacts that are in a reconstructed structure but not in the structure. For each model, we calculate the number of non-native contacts and divide it by the number of amino acids in the protein. Then we report the average percentage over all structures per protein per method. The comparison methods are the same ones that mentioned above.

As shown in Table 2, the proposed CO-VAE and DCO-VAE both have the highest percentage of the native contacts and the lowest non-native contacts among around 86% of all the proteins, especially with a very large superiority (e.g. 33% on average for native contacts and 61% on average for non-native contacts comparing to the highest performance of comparison methods). The results demonstrate that the proposed methods can accurately generate the contacts like the native one and the generated contact maps have the similar balance between the contacts and non-contacts in the structure. Moreover, even the DCO-VAE enhanced the disentanglement in the objective function of the model, its performance on contact map generation still remains infected, and sometimes even better than the CO-VAE.

Table 2. Evaluation of the contacts and non-contacts percentage in the generated structures

Protein ID	dimension	metric-type	VGAE	Graphite	GraphRNN	CO-VAE	DCO-VAE
1AIL	73	contacts	49.41	49.47	35.94	97.06	97.00
IAIL	73	non-contacts	87.86	87.84	62.26	5.14	5.00
1ALY	146	contacts	49.46	49.60	28.86	65.00	63.00
IALI	146	non-contacts	98.98	98.98	95.28	85.07	85.02
1AOY	78	contacts	49.46	49.44	27.39	82.73	83.00
IAOI	78	non-contacts	89.39	89.39	65.10	14.54	8.01
1PO0	54	contacts	49.57	49.56	28.64	83.41	83.34
1BQ9	54	non-contacts	84.57	84.53	66.92	12.73	11.43
1C8CA	64	contacts	49.44	49.42	27.94	92.03	93.00
ICOCA	64	non-contacts	86.68	86.68	68.11	9.65	8.08
1CC5	83	contacts	49.51	49.44	26.49	59.83	59.52
iccs	83	non-contacts	88.66	88.67	69.09	18.62	19.06
1DTDB	61	contacts	49.52	49.57	25.65	37.80	38.00
מטוטו	61	non-contacts	86.69	86.69	74.63	65.61	66.00
1DTJA	76	contacts	49.56	49.49	10.56	27.97	27.02
IDIJA	76	non-contacts	88.18	88.19	75.10	46.86	43.71
1HHP	99	contacts	49.48	49.55	24.32	90.95	91.11
111111	99	non-contacts	90.82	90.83	72.88	10.77	9.18
1HZ6A	72	contacts	49.48	49.51	35.11	45.53	46.00
IIIZOA	72	non-contacts	88.50	88.50	85.53	55.72	55.00
1ISUA	62	contacts	49.59	49.43	24.64	51.83	52.00
1130A	62	non-contacts	85.15	85.19	72.65	19.65	9.34
1SAP	66	contacts	49.52	49.43	29.21	91.70	92.18
ISAF	66	non-contacts	86.93	86.96	71.81	9.85	10.47
1TIG	94	contacts	49.75	50.11	43.70	100.00	100.00
111G	94	non-contacts	99.00	99.00	98.73	98.00	98.00
2H5ND	133	contacts	49.53	49.50	25.22	82.61	81.15
2113ND	133	non-contacts	93.56	93.61	76.86	19.92	19.40

Evaluating the learned graph distribution

Since the successful generation of contact maps rely on successfully learning the distribution of the contact maps, we evaluate whether the generated contact maps follow the learned distributions. By regarding each contact map as a graph,

we can calculate four properties of each graphs: density, number of edges, the degree, and transitivity. The density of a graph is the ratio of the number of edges and the number of possible edges. The average degree coefficient measures the similarity of connections in the graph with respect to the node degree. Transitivity is the overall probability for the graph to have adjacent nodes interconnected. All these properties can be calculated by the open source API NetworkX. Then the distance between the distribution of the generated contact maps and the distribution of the training sets in terms of the four properties is measured by three metrics: Pearson correlation coefficient, Bhattacharyya distance and Earth Mover's Distance (EMD). In statistics, the Pearson correlation coefficient (PCC) is a measure of the linear correlation between two variables *X* and *Y*. Here the *X* and *Y* refers to the a kind of graph property of the generated graphs and training graphs respectively. The Bhattacharyya distance and EMD both measures the similarity of two probability distributions and the smaller the better. The results for one example protein are shown as Table 3. In addition, we also compare our proposed methods with the comparison methods. The results for other proteins can be seen in the supplemental materials.

As shown in Table 3, considering EMD distance, the proposed P-GraphVAE has the best performance regarding all the graph properties. For the Bhattacharyya distance, the proposed P-GraphVAE outperformed other comparison methods by 33.38% and the disentangle version outperformed others by 33.68%. The Pearson similarity is very small, which demonstrate that the generated graphs are various in some degree than the training set, which ensure the diversity of the generated contact maps. Specifically, in terms of EMD distance, the proposed CO-VAE and DCO-VAE achieve 1.4 and 2.9 on average, which is 65.8% and 29.2% smaller than the best performance comparison method.

Table 3. Evaluation the learned distributions by comparing to the training sets for Protein 1DTJA

Graph Property	Methods	Pearson	Bhattacharyya	EMD
	Graphite	0.008	5.410	0.465
	GVAE	0.007	5.415	0.466
Density	GraphRNN	0.022	2.450	0.004
,	CO-VAE	0.018	4.941	0.002
	DCO-VAE	0.004	3.740	0.004
	Graphite	0.007	5.410	1327
	GVĀE	0.007	5.415	1329
Number of Edge	GraphRNN	0.022	2.450	12.18
O	CO-VAE	0.019	4.941	5.423
	DCO-VAE	0.004	3.741	11.29
	Graphite	0.030	5.056	0.136
	GVĀE	0.009	5.401	0.030
Ave-Degree Cor	GraphRNN	Nan	3.413	Nan
_	CO-VAE	0.005	5.310	0.017
	DCO-VAE	0.004	3.361	0.092
	Graphite	0.008	5.436	0.244
	GVAE	0.008	5.348	0.307
Transitivity	GraphRNN	0.009	2.886	0.122
•	CO-VAE	0.005	5.159	0.004
	DCO-VAE	0.005	3.571	0.027

Visualizing the 3D Structure from Generated Contact Maps

Given the generated contact maps by the proposed CO-VAE and DCO-VAE, we use the existing tool CONFOLD [1] to reconstruct the contact map to its responding 3D structure. To visualize and evaluate the real 3D structure of the generated proteins, we also visualize the protein structure by the tool PyMol [11] which is widely used and user-sponsored molecular visualization system. We show the results both for the proposed CO-VAE model and the DCO-VAE,

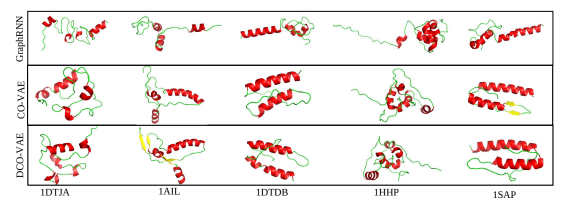


Fig. 1. Reconstructing the 3D structure from the generated contact maps: we reconstruct the 3D structure based on the generated contact maps of five proteins (i.e.,1DTJA, 1AIL, 1DTDB, 1HHP, and 1SAP) shown as examples; The generated contact maps are generated by the proposed CO-VAE and DCO-VAE, as well as the comparison method GraphRNN.

which is disentangled. And we also compare our proposed model with the graph generative model graphRNN. As shown in Fig. 1, the proposed CO-VAE and DCO-VAE have better performance in finding both the secondary and tertiary structure.

Interpreting the disentangled latent representation

It is important to be able to measure the level of disentanglement achieved by different models. In this subsection, we qualitatively demonstrate that our proposed DCO-VAE framework consistently discovers more latent factors and disentangles them in a cleaner fashion. By learning a latent code representation of a protein structure, we assume each variable in the latent code corresponds to a certain factor or property in generating the protein structure. Thus, by changing the value of one variable continuously and fixing the remaining variables, we could visualize the corresponding change of the generated contact map and 3D protein structure.

First, an example protein IDTJA is analyzed and displayed. the As shown in Fig 3 and Fig 2, we use the convention that in each one latent code varies from left to right while the other latent codes and noise are fixed. The different rows correspond to different random samples of fixed latent codes and noise. For instance, in Fig 2, one column contains four generated contact map samples regarding each variables, and a row shows the generated contact maps for 5 possible values among 1 and 10000 in a certain variable of latent code with other noise fixed. From the changing contact maps in each row, it is easily to find out which part of the contact map is related or controlled by the certain row. We highlight the part that controlled by each variable in red circle in Fig 2. The corresponding 3D structure visualization is shown in Fig 2. On this way, we could easily interpret the role of each variable in the latent code in controlling the generation of the contact maps and the 3D structure.

Second, some other proteins are also shown in Fig 5 and Fig 4, which demonstrate that the interpretable model can be generalized to any proteins.

4 DISCUSSION

In summary, our study aims at utilizing the deep neural network-based generative model for generating tertiary protein structure as well as interpreting the generation process. To the best of our knowledge, we showed for the first time the development and application of an interpretative graph variational auto-encoder for the problem of protein structure

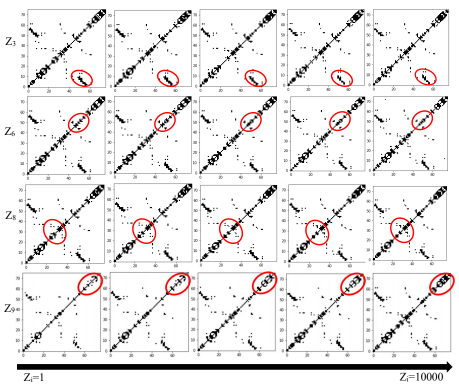


Fig. 2. Contact map interpretation for protein 1DTJA: Four semantic factors are discovered in the latent variables (i.e., Z_3 , Z_6 , Z_8 , and Z_9) which can control the local structural features of the contact maps; the value of latent variables travels from 1 to 10000 and five segments are selected to visualize the local structural feature variations.

generation and interpretation. This demonstrated the promise of generative models in directly revealing the latent space for sampling novel tertiary structures, highlighting axes/factors that carry structural meaning, and opening the black box of deep models.

By treating tertiary protein structure construction problem as the generation of contact maps, we proposed a new disentangled VAE, named Contact Map VAE (CO-VAE) with new graph encoder and decoder. It first learns the underlying distribution of contact maps and then generates additional contact maps by sampling from the learned distribution. The similarity of the structure (e.g., in terms of precision, recall, F1-score, and coverage) as well as positive contact percentage between the generated contact maps and the native ones demonstrated the quality of the generated contact maps. The proposed methods(CO-VAE and DCO-VAE) show great advantages than the existing models by obtaining the best performance in four metrics in almost 76% of the proteins and the highest percentage of the native contacts as well as the lowest non-native contacts among around 86% of all the proteins. Furthermore, we reconstructed the generated contact maps into 3D protein structures. The visualization of the constructed 3D protein illustrated that the generated contact maps are valid and useful for further generating the 3D tertiary protein structure.

To further investigate whether the proposed CO-VAE indeed learns the distribution of the observed contact map samples, we generated a set of contact maps and compared their underlying distribution and that of the real contact maps. Since it is difficult to directly evaluate the graph distribution, we resorted to evaluating the distribution of the properties of graphs, such as density, edge numbers, average degree correlation, and transitivity. The small EMD and

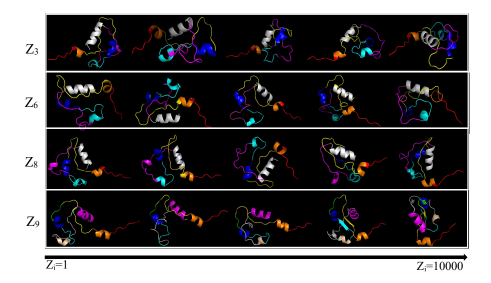


Fig. 3. Tertiary protein structure interpretation for protein 1DTJA: Tertiary protein structures are reconstructed based on the generated contact maps in Fig. 2; the amino acids within the same secondary structure are shown in the same color

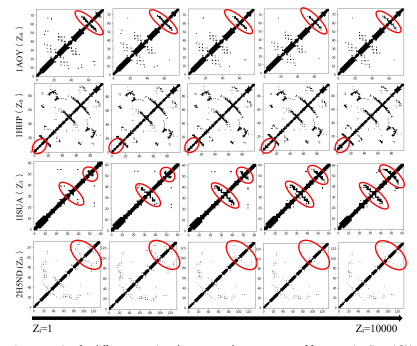


Fig. 4. Contact map interpretation for different proteins: the generated contact maps of four proteins (i.e.,1AOY, 1HHP, 1ISUA, and 2H5ND) are visualized as examples; for each protein, one of the semantic factors that are discovered in the latent variables is shown to control the local structural features of the contact maps; the value of latent variables travels from 1 to 10000 and five segments are selected to visualize the local structural feature variations.

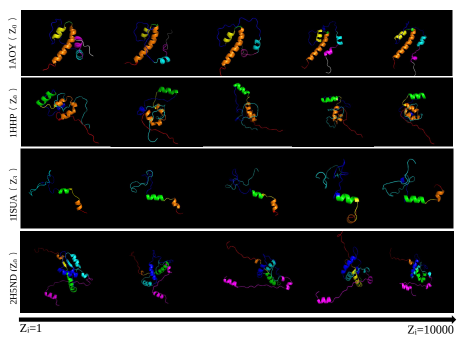


Fig. 5. Tertiary protein structure interpretation for different proteins: Tertiary protein structures are reconstructed based on the generated contact maps in Fig. 4; the amino acids within the same secondary structure are shown in the same color

Bhattachryya distances between the learned and real distribution in terms of all four properties of graphs generated by CO-VAE and DCO-VAE validated that the underlying graph distribution is effectively discovered, which outperformed the distances calculated from the graphs that generated by comparison methods by 33.4% and 47.5% on average. Though the Pearson correlation score was very low, it showed that the diversity of the generated graphs was ensured since it measures the strength and direction of a linear relationship between two variables other than the similarity of two distributions.

Next, to explore our generative model's capability of interpreting the process of contact map generation, we enhanced our CO-VAE by enforcing the weights of the second term of the training objective, leading to the disentanglement among variables and hence a new interpretable model named DCO-VAE. The learned latent representation variables in DCO-VAE is expected to be related to the factors that influence the formation of the contact maps. As a result, for each latent variable, by varying the value of this variable from 1 to 10,000 and fixing the values of others at the same time, a certain local part of the generated contact maps showed obvious trends (e.g., contracting or sketching). This demonstrated that the learned latent variables are effectively disentangled, which indicated the potential semantic factors in the formation of contact maps and the corresponding protein structures.

From one aspect, though there have been some deep generative models applied to the protein contact map generation, they cannot utilize the relationships among amino acids by treating the contact maps as the graph-structured data with graph generative models. From another aspect, though many interpretable learning models have been explored and applied into the image generation, there are no interpretable models that can discover the semantic factors that can control the process of protein folding. In summary, to the best of our knowledge, this is the first time an interpretable deep generative model for graphs have been applied for protein structure prediction and its effectiveness in generating

good-quality protein structures is demonstrated. In addition, the proposed DCO-VAE can also be applied to other real-world applications where there is a need for graph-structured data generation and interpretation.

There are much more promising and challenging topics that can be originated from the research in this paper for future exploration. For example, it would be interesting and potentially beneficial to develop an end-to-end tertiary protein structure generation model directly for 3D structure instead of contact maps. This is because there is a gap between the contact map generation and 3D structure formulation process, and the learned variables can only explain well of the formation of contact maps instead of the 3D structure. In addition, the exploration of the node-edge joint generative model would also be highly interesting. The proposed DCO-VAE model focuses on generating the graph topology (i.e., via contact maps) instead of node features (e.g., properties and types of amino acid). Jointly generating both graph topology and node features could be important in some cases, such as when directly generating the 3D structure where the node features can be the 3D positions.

5 METHODS

Problem Formulation

Recoverable tertiary structure formulation requires to preserve information not only on which atom is bonded with which else, but, more importantly, on which atoms are in proximity of each-other in three-dimensional space. As shown in Fig. 6, to address this issue, we employ contact map graph which can be trivially computed from tertiary structures as the input to our model. And to recover a tertiary structure from a contact map is comparatively trivial [1].

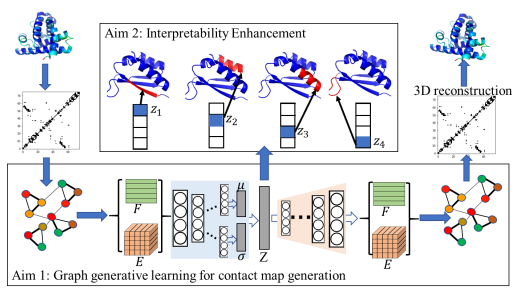


Fig. 6. Overall schematic of proposed generative learning framework.

Hence, the contact map is a graph-based representation of a tertiary structure that effectively embeds the three-dimensional Cartesian space into a two-dimensional space. Specifically, in our approach, let the contact graph G = (E, F) now be associated with its edge attribute tensor $E \in \mathbb{R}^{N \times N \times L_1}$, which denotes the adjacent matrix A when $L_1 = 1$; and node attribute matrix $F \in \mathbb{R}^{N \times L_2}$ (as shown in Fig. 6) denoting the identity of each atom by hot vector embedding, where N is the number of atoms over which contacts are computed; one can do so for all atoms in a molecule, or

representative atoms to control the size of the input space. The edge and node attributes are rich mechanisms to add additional information. For instance, node attributes can store not just the identities of the amino acids but also their PSSM profile (derived from the Position Specific Scoring Matrix), as well as their solvent accessibility and secondary structure as derived from a given tertiary structure. The edge attributes can encode additional information about contacts, such as their exact distance and/or contact predicted for that pair of amino acids (vertices) from sequence information alone. In our experiment, we are given an undirected, unweighted graph G as mentioned above, and we only use the adjacency matrix A of G (we assume diagonal elements set to 1, i.e. every node is connected to itself) and node attributes F store the identities of the amino acids with $L_2 = 20$ (the number of all the atom types).

Deep Contact Map Variational Auto-encoder

The task in this paper requires to learn the disentangled generative models on contact map that encodes graph information. Although disentanglement enhancement and deep graph generative models are respectively attracting fast-increasing attention in recent years, the synergistic integration of them has rarely been explored yet. To learn the distribution of the contact map decoys of a amino acid sequence, we first propose a new graph variational auto-encoder framework that consists of powerful graph encoder, graph decoder, and latent graph code disentanglement mechanism. First we introduce the model without disentanglement mechanism, named as The Deep Contact Map Variational Auto-encoder (CO-VAE).

Specifically, given the adjacent matrix A of a contact map, we further introduce stochastic latent variables z_i , summarized in an $1 \times H$ vector Z. The overall architecture contains a graph encoder and decoder which are trained by optimizing the variational lower bound Loss w.r.t. the variational parameters W_i :

$$\mathcal{L} = \mathbf{E}_{q(Z|F,A)}[logp(A|Z,F)] - KL[q(Z|F,A)||p(Z)]$$
(1)

where $KL[q(\cdot)||p(\cdot)]$ is the Kullback-Leibler divergence between $q(\cdot)$ and $p(\cdot)$. The first item is the reconstruction loss of the generated contact maps and the second term enforces the inferenced latent vector close to the real latent vector distribution. We take a Gaussian prior $p(Z) = \prod_i p(z_i) = \prod_i N(z_i|0,I)$

For the encoder part, we take a simple inference model parameterized by a two-layer Graph Convolution Nueral Network (GCN) [26] as the encoder of our CO-VAE:

$$q(Z|F,A) = \prod_{i=1}^{N} q(z_i|F,A), \quad \text{where} \quad q(z_i|F,A) = \mathcal{N}(z_i|\mu_i, diag(\sigma^2))$$
 (2)

Here $\mu = GCN_{\mu}(F, A)$ is the mean of the latent vectors, which is inferenced by GCN; similarly $log\sigma = GCN_{\sigma}(F, A)$ is the standard of the latent vector that is inferenced by another GCN. Thus, it is possible to sample Z from the distributions of the latent vectors q(Z|F, A).

For the generative model, we utilize the graph decoder network proposed in our previous works[17]. This work first proposes the graph deconvolution based graph decoders, which achieve the best performance in the graph generation task when the node set of the graph is fixed. Thus, we choose this graph decoder as part of our CO-VAE.

$$p(A|Z,F) = \prod_{i=1}^{N} \prod_{i=1}^{N} p(A_{ij}|Z)$$
(3)

where A_{ij} are the elements of A.

We perform full-batch gradient descent and make use of the re-parameterization trick [25] for training. The architecture and mathematical operations of the graph encoder for modelling the q(Z|A, F) and decoder for modelling the p(A|Z, F) are detailed in the supplemental materials.

Disentangled Contact Map VAE

Then next we introduce the Contact Map VAE with the disentanglement mechanism, which is inpired by the β –VAE [20]. β –VAE is proposed for automated discovery of interpretable factorised latent representations from data in a completely unsupervised manner, and has been currently used in image domain [21] and Natural Language Processing [54]. It is a modification of the variational autoencoder (VAE) framework by introducing an adjustable hyperparameter β that balances latent channel capacity and independence constraints with reconstruction accuracy. Thus, we proposed the Disentangled Contact Map VAE (DCO-VAE) to largely interpret the latent representations for the protein generation. We propose augmenting the CO-VAE framework with a single hyperparameter β that modulates the learning constraints applied to the model. These constraints impose a limit on the capacity of the latent information channel and control the emphasis on learning statistically independent latent factors. Eq. 1 can be re-written to arrive at the DCO-VAE formulation, but with the addition of the β coefficient:

$$\mathcal{L} = \mathbf{E}_{q(Z|F,A)}[logp(A|Z,F)] - \beta KL[q(Z|F,A)||p(Z)]$$
(4)

where with the $\beta = 1$ corresponds to the CO-VAE framework mentioned above; and with $\beta > 1$ the model is pushed to learn a more efficient latent representation of the data, which is disentangled if the data contains at least some underlying factors of variation that are independent.

DATA AVAILABILITY

All data, models, and source code are freely available to readers upon contacting the authors.

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AUTHOR CONTRIBUTIONS

XG implemented and evaluated the proposed methodologies, as well as drafted the manuscript. ST assisted with preparation of the input data and the evaluation of reconstructed structures. LZ and AS conceptualized the methodologies and provided guidance on implementation and evaluation, as well as edited and finalized the manuscript.

COMPETING INTERESTS

The authors declare no competing interests.