Advance Access Publication Date: 13 May 2021 Original Paper



Structural bioinformatics

SPOT-1D-Single: improving the single-sequence-based prediction of protein secondary structure, backbone angles, solvent accessibility and half-sphere exposures using a large training set and ensembled deep learning

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Associate Editor: Dr. Pier Luigi Martelli

Received on November 6, 2020; revised on April 6, 2021; editorial decision on April 22, 2021; accepted on April 26, 2021

Abstract

Motivation: Knowing protein secondary and other one-dimensional structural properties are essential for accurate protein structure and function prediction. As a result, many methods have been developed for predicting these one-dimensional structural properties. However, most methods relied on evolutionary information that may not exist for many proteins due to a lack of sequence homologs. Moreover, it is computationally intensive for obtaining evolutionary information as the library of protein sequences continues to expand exponentially. Here, we developed a new single-sequence method called SPOT-1D-Single based on a large training dataset of 39 120 proteins deposited prior to 2016 and an ensemble of hybrid long-short-term-memory bidirectional neural network and convolutional neural network.

Results: We showed that SPOT-1D-Single consistently improves over SPIDER3-Single and ProteinUnet for secondary structure, solvent accessibility, contact number and backbone angles prediction for all seven independent test sets (TEST2018, SPOT-2016, SPOT-2016-HQ, SPOT-2018, SPOT-2018-HQ, CASP12 and CASP13 free-modeling targets). For example, the predicted three-state secondary structure's accuracy ranges from 72.12% to 74.28% by SPOT-1D-Single, compared to 69.1-72.6% by SPIDER3-Single and 70.6-73% by ProteinUnet. SPOT-1D-Single also predicts SS3 and SS8 with 6.24% and 6.98% better accuracy than SPOT-1D on SPOT-2018 proteins with no homologs (Neff = 1), respectively. The new method's improvement over existing techniques is due to a larger training set combined with ensembled learning.

Availability and implementation: Standalone-version of SPOT-1D-Single is available at https://github.com/ias-preet/ SPOT-1D-Single. Direct prediction can also be made at https://sparks-lab.org/server/spot-1d-single. The datasets used in this research can also be downloaded from GitHub.

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1 Introduction

The past two decades have seen many developments in the field of deep learning-based prediction of protein structure (Yang et al., 2018). Significant headway has been observed specifically for the protein secondary structure and contact map prediction (Fang et al., 2018; Hanson et al., 2019; Li et al., 2019; Wang et al., 2016; Wu et al., 2020). These improvements have ultimately led to a considerable improvement in protein tertiary structure prediction, as observed in CASP13 (Cheng et al., 2019). Particularly, the protein secondary structure prediction is approaching the theoretical upper bounds at 88-90% accuracy with SPOT-1D prediction of three-state secondary structure at 86.18% and eight-state secondary structure at 79% accuracy (Hanson et al., 2019; Yang et al., 2018).

However, most of the above-stated improvement came from evolutionary-profile-based methods (Hanson et al., 2019; Klausen SPOT-1D-Single 3465

et al., 2019; Wang et al., 2016; Xu et al., 2020). These methods employed the feature profiles generated by PSI-BLAST (Altschul et al., 1997), HHblits (Remmert et al., 2012) and the output of other predictors (Cuff and Barton, 2000). However, more than 90% of proteins have none or very few homologous sequences (Ovchinnikov et al., 2017). Their predicted secondary structure and other structural properties are substantially less accurate than those with many homologous sequences (Heffernan et al., 2018). Thus, it is necessary to develop single-sequence or no evolutionary information-based methods. Developing single-sequence-based methods is important because one can only claim that the problem of secondary structure prediction is solved if only a single sequence is utilized as an input for prediction. After all, proteins fold into secondary and tertiary structures from their single sequences only.

Unlike evolution profile-based methods, only a few methods were developed for single-sequence-based prediction of one-dimensional structural properties. McGuffin *et al.* (2000) proposed a profile-based secondary structure predictor along with a single-sequence-based predictor, which we will refer to as PSIPRED-Single. A method called accessible surface area (ASA)-quick employed single-sequence information to predict the Accessible Surface Area (Faraggi *et al.*, 2017).

Recently, we developed SPIDER3-Single that was dedicated to single-sequence-based prediction for not only secondary structure and solvent accessibility but also other structural properties such as backbone torsion angles and half-sphere exposures (HSE) (Heffernan et al., 2018). SPIDER3-Single took advantage of iterative learning on a two-layer Bidirectional Long Short-Term Memory Recurrent Neural Network trained on a training set of 9993 proteins (TR9993) which was also employed to train a profile-based method SPIDER3. This profile-based technique was substantially improved by SPOT-1D, which employs an ensembled learning from hybrid long-short-term-memory (LSTM) and Convolution based architectures (Hanson et al., 2019).

More recently, ProteinUnet (Kotowski et al., 2020) demonstrated a more computationally efficient deep learning-based method with comparable performance to SPIDER3-Single. This method employed an ensemble of Unet architectures popularly used in medical imaging tasks (Ronneberger et al., 2015). It was trained on the same training set as SPIDER3-Single after removing proteins longer than 1024 residues.

In this work, we examine the possibility of further improving single-sequence-based prediction of one-dimensional protein structural properties by employing an ensemble of hybrid LSTM-CNN model architecture trained on a large dataset of 39 120 proteins. We demonstrate that the larger dataset and the ensemble, to a lesser extent, allows a substantial and consistent improvement over SPIDER3-Single and ProteinUnet over several independent test sets.

2 Materials and methods

2.1 Datasets

To examine the effect of training on a large dataset, we utilized the benchmark dataset prepared by ProteinNet (AlQuraishi, 2019). It consists of 50 914 proteins submitted to PDB before 2016 with a high X-ray resolution (<2.5 Å) crystal structure and clustered at sequence identity cutoff of 95% according to MMseqs2 tool (Steinegger and Söding, 2017). ProteinNet provides the datasets at different sequence identity cutoffs, but we choose the dataset with the sequence identity cutoff of 95% to get more training data to harness the full capabilities of recent deep learning algorithms.

For independent testing and comparison, we downloaded all protein structures released between January 2016 and April 2020. Because the existence of remote homologs makes it insufficient to remove homologous sequences by a sequence identity cutoff, we removed potential homologs of the test data by comparing the Hidden Markov Models of all post-2016 proteins to the Hidden Markov Models of all pre-2016 proteins using the HHSEARCH tool (Steinegger *et al.*, 2019). Any proteins with an e-value cutoff of less than 0.1 were removed from the test set. This led to 1473

proteins as the stringent test set SPOT-2016. These 1473 proteins include all proteins without any resolution-based constraints applied. To create a high-resolution test set, we applied resolution constraints of <2.5 Å and R-free <0.25. This separated 295 proteins from SPOT-2016 and created a new test set SPOT-2016-HQ.

To further provide a fair comparison with models that would have possibly used proteins until 2018, we separated another subset of SPOT-2016 of proteins released after January 2018. We also performed a remote homolog search using their Hidden Markov Models against the Hidden Markov Models of all proteins released before 2018 with the same constraints as previous test sets. This led to 682 proteins forming the strict test set SPOT-2018. Also, based on resolution constraints, we separated 125 proteins at the resolution <2.5 Å and R-free <0.25 forming the test set SPOT-2018-HQ.

Apart from SPOT-2016, SPOT-2016-HQ, SPOT-2018 and SPOT-2018-HQ, we use three additional independent test sets: TEST2018, CASP12-FM and CASP13-FM. TEST2018 is a test set employed in SPOT-1D (Hanson *et al.*, 2019). It includes 250 proteins released between January 01, 2018 and June 17, 2018 with resolution <2.5 Å and R-free <0.25, filtered at a sequence identity of 25% using Blastclust against all pre-2018 proteins released on PDB. CASP12-FM is a test set of 22 protein targets constituting free modeling targets released during CASP12 (Schaarschmidt *et al.*, 2018). Similarly, CASP13-FM contains 17 free modeling targets released during CASP13 (Kryshtafovych *et al.*, 2019). Free modeling targets are those proteins without known structural templates in the protein databank at the time of release.

To minimize possible over-fitting, we separated 100 proteins from the training set and compared their Hidden Markov Models generated by HHblits with the Hidden Markov Models of all other proteins in the training set using HHSEARCH. Any training proteins, which had hits with the 100 validation proteins at an e-value cutoff of less than 0.1, were removed from the training set. This left us with the final 39120 proteins for training.

2.2 Outputs

For a classification task (multi-task prediction), our deep learning predictor has eight output nodes for eight-state secondary structure and three output nodes for three-state secondary structure prediction. We employed the Dictionary of Secondary Structure of Proteins (DSSP) (Kabsch and Sander, 1983) definition of assigning eight secondary structure classes according to protein 3D structures. These eight secondary structure states are: 3₁₀-helix (G), alpha-helix (H), pi-helix (I), beta-bridge (B), beta-strand (E), high curvature loop (S), beta-turn (T) and coil (C) states. The above eight states can be further simplified to the three-state labels of strand E (B and E in the eight-state definition), helix H (G, H, and I in the eight-state definition) and coil C (everything else in the eight-state definition).

Our method is not limited to the prediction of the secondary structure of the proteins. It also predicts the ASA, protein backbone angles, HSE and contact number (CN). ASA is a measure of the area of an amino acid in a protein that is exposed to a solvent molecule (Chothia, 1974). Here, we predict the relative ASA (rASA) to avoid any bias, and later it is converted to the absolute ASA. Similar to ASA, another 1D property is HSE, which is another measure of how buried an amino acid is in a protein according to the number of contacts. HSE separates an amino acid residue's contacting sphere into two half-spheres, up and down. Both these half-spheres have different HSE called HSE-up and HSE-down, and the sum of these two make the CN (Heffernan et al., 2016). The backbone angles that this model predicts are ψ , ϕ , θ and τ . The first two angles are the backbone torsion angles. The DSSP software was utilized to generate the ψ and ϕ angles from protein structures (Cornilescu et al., 1999). The other two angles θ and τ are also crucial as θ is the angle between the $C\alpha_{i-1}$ - $C\alpha_{i}$ - $C\alpha_{i+1}$ and τ is the dihedral angle rotated about the $C\alpha_i$ - $C\alpha_{i+1}$ vector (Lyons *et al.*, 2014). Protein backbone and dihedral angles are not predicted directly but as a function of sine and cosine of the angle due to their periodicity. In total, twelve regression output nodes are available, eight output nodes for the angles and four more for the ASA, HSE-d, HSE-u and CN.

3466 J.Singh et al.

2.3 Performance evaluation

The performance evaluation for different tasks has been divided into three categories: accuracy, correlation coefficient and mean absolute error (MAE). SS3 and SS8 prediction performance of the model was measured by the accuracy. Pearson's correlation coefficient (PCC) between the predicted values and the true values of the ASA, HSE-u, HSE-d and CN were calculated for each protein and then averaged for the dataset (Benesty *et al.*, 2009). To evaluate the model performance for backbone angles, we calculate the MAE between the true and the predicted values for all residues for all proteins concatenated together. To show the statistical significance of

improvement by SPOT-1D-Single over SPIDER3-Single and ProteinUnet, a paired *t*-test was used across SS3, SS8, ASA, HSE-u, HSE-d, CN and backbone angles to obtain *P*-value (Lovric, 2011).

2.4 Neural networks

Our deep neural network architecture shown in Figure 1 was inspired by the recent success of the deep neural network for protein 1D structural properties using evolutionary information. In particular, SPOT-1D (Hanson *et al.*, 2019) employed an ensemble of models by using variants of ResNet and bidirectional recurrent neural

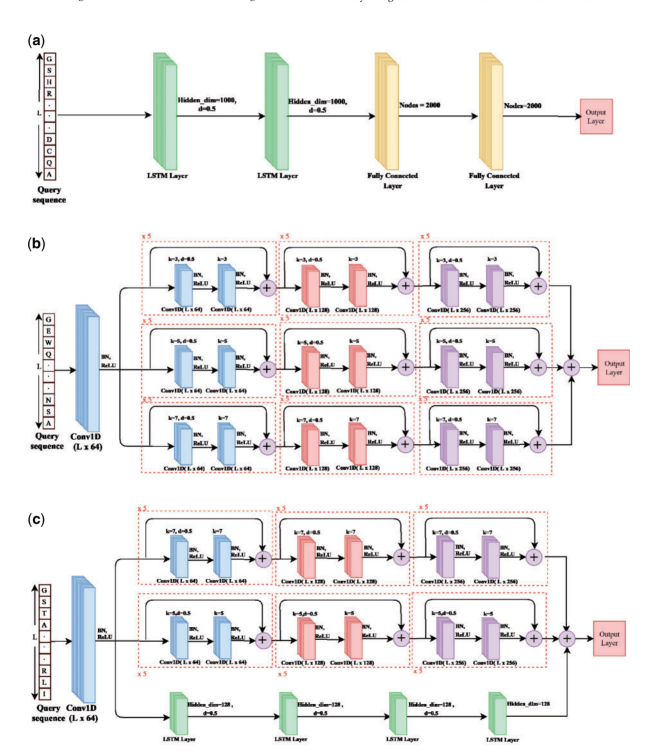


Fig. 1. Overview of the model architecture

SPOT-1D-Sinale 3467

networks (Schuster and Paliwal, 1997). By comparison, the previous single-sequence-based predictor (SPIDER3-single) employed two bidirectional LSTM layers and two fully connected layers only. Here, as in SPOT-1D, we explored many deep neural network models based on the architecture shown in Figure 1 to predict protein 1D structural properties using single-sequence only.

As shown in Figure 1a, the first model we trained consists of two bidirectional-LSTM stacks with hidden dimensions (Hidden_dim) of 1000 and a dropout (d) of 0.5 after each stack to avoid overfitting (Schuster and Paliwal, 1997; Srivastava *et al.*, 2014). After the LSTM layers, two fully connected layers of size 2000 are stacked. Similar models have been used in our previous profile-based predictors (Hanson *et al.*, 2018; 2019).

The second model (Fig. 1b), is a multi-scale parallel 1D ResNet model. It is a variation of ResNet architecture selected based on hyper-parameter tuning. Previously, ResNet architectures have exhibited high-performance accuracy, as shown in SPOT-1D (Hanson et al., 2019) and SPOT-Contact (Hanson et al., 2018). In this model, the data is passed through the first initial block with a 64 filter convolution layer of kernel size 7, followed by batch normalization and ReLU activation function (Agarap, 2018; Ioffe and Szegedy, 2015). After that, the output of the initial block is passed through three parallel stacks of ResNets. All three parallel stacks contain 15 blocks of ResNet stacked together. The first five stacks contain 64-filter convolutional layers, then the next five have 128 filters, and the last five have 256 filters. All residual blocks stacked in a parallel fashion (shown in the dashed red line in Figure 1b) containing two 1D convolutional layers, each followed by batch normalization and ReLU activation function (Agarap, 2018; Ioffe and Szegedy, 2015). After the first set of convolution, batch normalization, and ReLU activation function, we applied a dropout of 0.5 in each block. The only difference among three parallel stacks of ResNet blocks is their kernel sizes (the blocks in the first, second, and third stack have a kernel size of 3, 5 and 7, respectively). The output of the three parallel stacks is then concatenated at the end and passed to the output layer.

The third model as shown in Figure 1c is similar to the second model. It also contains three parallel stacks of layers. The only difference is that instead of a stack of ResNet models with three kernel size in the parallel stack, this model has 4 layers of bidirectional LSTM stacked together with a hidden size of 128 and dropout of 0.5. This model was inspired by the success of hybrid CNN and LSTM networks in SPOT-1D and SPOT-Contact.

The above three selected models were inspired by the neural network architecture used in SPOT-1D. The first SPOT-1D model is a two-layer bidirectional-LSTM. We tested the LSTM model for a hidden dimension varying between 64 and 2048, and a model with two layers and a hidden dimension of 1024 performs the best as shown in Supplementary Table S1. The second SPOT-1D model is a ResNet. We tried different hyperparameters for ResNet with the number of blocks varying between 10 and 50 and the kernel size ranging from 3 to 7. ResNet with 45 blocks performs the best with kernel size 7. Instead of using a vanilla ResNet, we experimented with multi-scale ResNet keeping the same 45 blocks and it performed better than the vanilla ResNet on the validation set. The third model, a ResNet-LSTM model is a hybrid of the two models above with the maximum number of the hidden dimensions of the LSTM layers that we could train on the GPU. Due to the limited availability of the computing power, we limit the hyperparameter search to optimize on the secondary structure classification only.

The above three models are used for both classification and regression tasks. The classification model is trained on a batch size of ten using Cross-entropy loss and Softmax as the output layer activation function while the regression model is trained using the L1 Loss on a batch size of ten with a Sigmoid output layer activation function. Instead of using the average loss, we use a sum loss for both tasks. In the end, the output of all the classification models are ensembled by calculating the mean of the classification predictions. For the regression models, we take the mean ensemble for the ASA, HSE-u, HSE-d, CN and median ensemble for the angles as done by SPOT-1D (Hanson *et al.*, 2019).

2.5 Method comparison

We compared SPOT-1D-Single with ProteinUnet and our previous predictor SPIDER3-Single. We also compare to PSIPRED-Single for three-state secondary structure prediction and ASA-quick in ASA prediction. All the above-stated methods have stand-alone programs available online from https://codeocean.com/capsule/2521196/tree/v1, https://servers.sparks-lab.org/downloads/SPIDER3-Single_np. tgz, http://bioinfadmin.cs.ucl.ac.uk/downloads/psipred/ and http://mamiris.com/GENN+ASAquick.tgz, respectively.

3 Results

3.1 Effect of using a large training dataset

To examine the effect of using a large dataset for training, we employed an ensemble of the same three models trained on the ProteinNet dataset (39 120 proteins) and SPOT-1D dataset (10 200 proteins). The results for three-state (SS3) and eight-state (SS8) secondary structure prediction on the seven test sets were shown in Table 1. On SPOT-2018, SPOT-1D-Single trained on the bigger training set improves over the one trained on the smaller dataset by 2.42% with an accuracy of 60.09%, compared to 58.67% on the eight-state prediction (SS8). Similar trends are observed on all other test sets and the three-state prediction (SS3), as well. Such improvement due to a larger training set can also be observed in other structural properties. Supplementary Table S2 shows that the ASA, HSE, and CN shows a consistent improvement across all seven test sets. Supplementary Table S3 also exhibits similar improvement trends for protein backbone angles across seven different test sets.

3.2 Ensemble learning performance

To demonstrate the advantage of ensemble learning over individual models, Table 2 presents the results of the selected three models and the ensemble of the three models on TEST2018. The ensemble performance for SS3 and SS8 is 1.10% and 1.15% better than the best performing single model for both categories. The error for angle prediction has also reported a drop of 2.27%, 1.16%, 1.42% and 1.93% in error for ψ , ϕ , θ and τ angle prediction by the best performing single sequence method in the respective categories. Similarly, an ensemble shows improvement in the PCC for ASA, HSE-U and CN predictions. Similar levels of improvement were also found in other test sets as shown in Supplementary Tables S4–S6.

The above result of an ensemble is obtained by the mean of all models (except median for angle prediction). We employed the mean after investigating three different ensemble techniques (mean, median, and the majority voting). As shown in Supplementary Table S7, the best option among the three techniques is to use the mean

Table 1. The effect of the size of the training set on the prediction accuracy of three-state (SS3) and eight-state (SS8) secondary structure on seven different test sets: TEST2018, SPOT-2016, SPOT-2016-HQ, SPOT-2018, SPOT-2018-HQ, CASP12-FM and CASP13

Training set	Test20	018	SPOT-	2016	SPOT-20	16-HQ	SPOT-	2018	SPOT-20	18-HQ	CASP12	2-FM	CASP13	3-FM
	SS3	SS8	SS3	SS8	SS3	SS8	SS3	SS8	SS3	SS8	SS3	SS8	SS3	SS8
SPIDER3-Single (9993)	72.57	59.81	72.04	58.85	72.16	59.93	71.29	57.38	70.76	57.95	69.152	55.146	71.264	56.995
SPOT-1D Training Set (10200)	73.26	61.26	73.02	60.12	72.93	61.12	72.32	58.67	71.07	59.05	70.87	56.62	70.71	58.27
ProteinNet Set (39120)	74.28	62.17	74.29	61.39	73.65	61.59	73.71	60.09	72.12	59.66	72.44	57.59	73.21	60.93

3468 J.Singh et al.

Table 2. Individual model performance as compared to the ensemble performance on Test2018 for prediction of secondary structure in three (SS3) and eight (SS8) states, solvent accessibility (ASA), half-sphere-exposure-up (HSE-u), half-sphere-exposure-down (HSE-d), contact number (CN), backbone angles (ψ, ϕ, θ) and τ)

Model	SS3	SS8	ASA	HSE-u	HSE-d	CN	ψ	ϕ	θ	τ
2 Layer LSTM	73.37	61.46	0.664	0.558	0.548	0.572	42.033	22.562	9.621	43.681
Multi-Scale ResNet	73.00	60.93	0.648	0.554	0.539	0.559	41.527	22.416	9.480	43.152
Multi-Scale ResNet LSTM	73.47	61.24	0.641	0.554	0.541	0.562	41.732	22.471	9.531	43.342
Ensemble (SPOT-1D-Single)	74.28	62.17	0.665	0.573	0.563	0.585	40.585	22.155	9.345	42.315

Note: Performance measures are accuracy for SS3 and SS8, correlation coefficient for ASA, HSE-u, HSE-d and CN, and mean absolute errors for the angles.

Table 3. Performance comparison of SPOT-1D-Single with other predictors for seven different test sets TEST2018, SPOT-2016, SPOT-2016-HQ. SPOT-2018-HQ. CASP12-FM and CASP13-FM

Model	Test20	018	SPOT-	2016	SPOT-20	016-HQ	SPOT-	2018	SPOT-20)18-HQ	CASP12	2-FM	CASP1	3-FM
	SS3	SS8	SS3	SS8	SS3	SS8	SS3	SS8	SS3	SS8	SS3	SS8	SS3	SS8
PSIPRED-Single	68.91	_	70.29	_	69.45	_	68.00	_	68.00	_	68.13	_	68.05	_
SPIDER3-Single	72.57	59.81	72.04	58.85	72.16	59.93	71.29	57.38	70.76	57.95	69.152	55.146	71.264	56.995
ProteinUnet	72.57	60.30	_	_	_	_	_	_	_	_	71.33	57.56	70.63	58.07
SPOT-1D-Single (this work)	74.28	62.17	74.29	61.39	73.65	61.59	73.71	60.09	72.12	59.66	72.44	57.59	73.21	60.93
SPOT-1D (profile)	86.18	75.41	81.73	69.32	83.06	71.72	80.39	67.43	82.02	70.51	79.53	65.92	83.55	71.22
Short length sequence (less than 1024))													
PSIPRED-Single	_	_	70.38	_	69.40	_	67.94	_	67.94	_		_	_	_
SPIDER3-Single	_	_	72.14	59.14	72.25	60.03	71.31	57.57	71.02	58.23	_	_	_	_
ProteinUnet	_	_	73.00	60.14	72.72	60.55	72.20	58.71	71.28	58.74	_	_	_	_
SPOT-1D-Single (this work)	_	_	74.44	61.71	73.69	61.58	73.80	60.35	72.22	59.70	_	_	_	_
SPOT-1D (profile)	_	_	82.08	69.88	83.15	71.78	80.52	67.76	81.97	70.41	_	_	_	_

Note: The values provided below are the percentage accuracy of three-state secondary structure (SS3) and eight-state secondary structure (SS8) prediction.

ensemble for Secondary Structure, ASA, HSE and CN. Although the mean ensemble is also the best for angle prediction, we employed median to prevent angles from locating in the low probability or forbidden region.

3.3 Method comparison

Table 3 compares the performance of SPOT-1D-Single (this work) with PSIPRED-Single, SPIDER3-Single, ProteinUnet, and SPOT-1D (profile-based) for seven different test sets (TEST2018, SPOT-2016, SPOT-2016-HQ, SPOT-2018, SPOT-2018-HQ, CASP12-FM and CASP13-FM). As ProteinUnet does not predict a sequence with more than 1024 amino acids, the sub-table shows the results for SPOT-2016, SPOT-2016-HQ, SPOT-2018 and SPOT-2018-HQ excluding the proteins longer than 1024 amino acids. SPOT-1D-Single consistently performs better than other single-sequence-based method on secondary structure prediction for three (SS3) or eight (SS8) states classification. For TEST2018, SS3 and SS8 were predicted with 74.28% and 62.17% accuracies, respectively, by SPOT-1D-Single, which are 2.35% and 3.1% better than the next best ProteinUnet, respectively. On the new test set SPOT-2016, SPOT-1D-Single improves over SPIDER3-Single by 3.12% for SS3 and 4.3% for SS8, respectively. Consistent improvement for SPOT-2016-HQ, SPOT-2018, SPOT-2018-HQ, CASP12-FM and CASP13-FM sets by SPOT-1D-Single over ProteinUnet and SPIDER3-Single are also observed for SS3 and SS8, indicating that the improvement of SPOT-1D-Single over other predictors is robust. This is further confirmed by the statistical significance analysis for the results in Supplementary Table S8.

It is of note that using profiles (SPOT-1D) has an advantage of >10% improvement over SPOT-1D-Single. This large difference highlights the challenge facing single-sequence-based prediction. However, SPOT-1D has a larger performance fluctuation across different test datasets (79.5–86.2% for SS3), whereas SPOT-1D-Single's performance is more robust (72.4–74.3% for SS3). This is

likely because the number of homologous sequences for different datasets is different. In fact, the average Neff values (the number of the effective homologous sequences from HHblits) are 6.92, 4.82, 5.07, 4.38, 4.7, 5.73 and 6.99 for TEST2018, SPOT-2016, SPOT-2016-HQ, SPOT-2018, SPOT-2018-HQ, CASP12-FM and CASP13-FM, respectively.

Figure 2 shows the accuracy of secondary structure prediction as a function of Neff on SPOT-2018. At low Neff values, SPOT-1D-Single exhibits better performance than SPOT-1D for both three (SS3) and eight (SS8) state secondary structure prediction. Table 4 further examined the method performance for all Neff = 1 proteins (i.e. proteins with no homologs). SPOT-1D-Single predicts SS3 and SS8 at 76.57% and 65.24% respectively, which is 6.24% and 6.98%, respectively higher than SPOT-1D. In fact, if we treated all proteins in SPOT-2018 as proteins without homologs by using a single MSA and generated the results for SPOT-1D, there is a significant drop in the performance of SPOT-1D as shown in Table 4. This confirms that SPOT-1D achieves higher accuracy only for those proteins with sufficient evolutionary information.

Table 5 examines the performance of different predictors across different datasets for ASA, HSE-U, HSE-D and CN prediction. For TEST2018, SPOT-1D-Single predicts ASA with a 2.78% higher PCC than SPIDER3-Single and 7.25% higher than ProteinUnet and ASA-quick. Similar improvement trends are observed for other structural properties (HSE-U, HSE-D, CN, ψ , ϕ , θ , τ) as shown in Tables 5 and 6. The difference is statistically significant as shown in Supplementary Table S8. SPOT-1D using homologous information performs significantly better than SPOT-1D-Single for all regression tasks, but the difference in performance measure for SPOT-1D varies largely across different test sets as compared to SPOT-1D-Single. Similar to secondary structure prediction, Figure 3 shows that SPOT-1D performs worse than SPOT-1D-Single at low Neff. At Neff = 1, Table 4 shows that SPOT-1D-Single improves over SPOT-1D by of 7.57%, 6.53%, 7.86%, and 9.66% in the prediction of ψ , ϕ , θ and τ . Interestingly, SPOT-1D is slightly better for

3469

(a) SS3 vs Neff plot 100 SS8 v

SPOT-1D-Single

Fig. 2. Prediction accuracy as a function of the number of effective homologous sequence (Neff) by SPOT-1D-Single compared with other methods on SPOT-2018-short as labeled for (a) three-state secondary structure and (b) eight-state secondary structure

predicting HSE-u and HSE-d but SPOT-1D-Single is still better for ASA and CN.

To further understand where is the improvement of SPOT-1D-Single over SPIDER3-Single and ProteinUnet, we examined the dependence of the accuracy of the secondary structure as a function of the number of local (|i-j| < 20) (Fig. 4) and non-local contacts ($|i-j| \ge 20$) (Fig. 5) per residue for each protein, where i and j is the sequence position of the amino acid residues. It seems that SPOT-1D-Single improves over SPIDER3-Single most for those residues with few local contacts whereas the former consistently improves over the latter for proteins with a different number of non-local contacts. In other words, SPOT-1D-Single captures long-range interactions better than existing techniques.

To test the performance of different predictors for proteins in different structural folds, we clustered SPOT-2018 into different evolutionary classifications based on ECOD (Cheng et al., 2014). SPOT-2018 proteins were selected to be unique from existing structures based on HMM similarity. As a result, this set is not well covered by automated structural classifications. Out of 682 proteins of SPOT-2018, 147 are classified into 17 different categories and the remaining 535 are marked as unclassified. Unclassified proteins include some naturally disordered obligate multimer fragments such as 6S29_D but also several bona fide structured domains that are missed by automated classification schemes such as the archetypal designed beta-barrel found in 6OHH_B. Supplementary Table S9 shows the performance comparison of SPOT-1D-Single and SPIDER3-Single based on the available ECOD classifications. In general, protein structures with more beta sheets appear more difficult to predict than proteins with more alpha helices.

4 Discussion

In this article, we have developed a new single-sequence-based method for predicting one-dimensional structural properties of proteins, including secondary structure, solvent accessible surface area, and backbone torsion angles. We employed an ensemble of hybrid LSTM-CNN network architecture and a large training set of approximately 40 000 proteins with validation and test sets that are non-redundant from the large training set according to HHSEARCH. The improvement of SPOT-1D-Single over ProteinUnet, SPIDER3-Single and ASA-quick is consistent across all seven test sets (TEST2018, SPOT-2016, SPOT-2016-HQ, SPOT-2018, SPOT-2018-HQ, CASP12-FM and CASP13-FM). The accuracy of SPOT-1D-Single is higher than the evolutionary-profile-based SPOT-1D when the number of effective homologous sequences is low. This highlights that SPOT-1D-Single can be used as a reasonably accurate screening tool for protein one-dimensional structural properties.

In the interest of profiling our method in terms of computational time, we have measured the time taken by our method SPOT-1D-Single to predict the secondary structure and other 1D properties. As shown in Table 7, it takes 155 s and 20 s to predict 250 proteins in TEST2018 by our local machine on both

Table 4. Performance comparison of SPOT-1D, SPOT-1D-Single and SPOT-1D(Single MSA) on SPOT2018 and all proteins with Neff = 1 in SPOT-2018 for prediction of secondary structure in three SS3) and eight (SS8) states, solvent accessibility (ASA), half-sphere-exposure-up (HSE-u), HSE-down (HSE-d), contact number (CN), backbone angles (ψ , ϕ , θ and τ)

Model	SPOT-2	SPOT-2018 (NEFF=1)	FF = 1)								SPOT-2018	1018								
	SS3	888	ASA	HSE-u	HSE-d	$^{\rm CN}$	SS3 SS8 ASA HSE-u HSE-d CN ψ	ϕ	θ	2	SS3	888	ASA	HSE-u	SS3 SS8 ASA HSE-u HSE-d CN ψ	CN	ϕ	ϕ	θ	2
SPOT-1D-Single 76.57 65.24 0.641 0.370 0.522 0.547 (this work)	76.57	65.24	0.641	0.370	0.522	0.547	43.106	43.106 21.047	9.495	9.495 41.644 73.71 60.09 0.620 0.403	73.71	60.09	0.620	0.403	0.479	0.487	44.407	0.479 0.487 44.407 22.864	9.851	9.851 43.787
SPOT-1D (profile)		72.07 60.98	0.616	0.373	0.616 0.373 0.523	0.535	46.640	22.519	10.305	46.097	80.39	67.43	0.691	0.518	0.595	909.0	34.790	20.390	8.515	34.011
SPOT-1D (profile)	I	I		I	I	I	I	I	I	I	69.62	56.23	0.594	0.365	0.452	0.462	49.208	24.171		48.581
(Single MSA)																				

Note: Performance measures are accuracy for SS3 and SS8, correlation coefficient for ASA, HSE-u, HSE-d and CN, and mean absolute errors for the angles

Table 5. Performance comparison of SPOT-1D-Single with other predictors for seven different test sets TEST2018, SPOT-2016, SPOT-2016, SPOT-2018, SPOT-2018, SPOT-2018, DAT-2018, DAT-2018, SPOT-2018, DAT-2018, DAT-2018

0.495

0.462 0.480 0.489 0.683

0.572 0.565

0.554 0.599 0.692

0.467 0.482 0.522 0.621

> 0.516 0.523 0.556 0.660

0.504 0.586 0.540 0.612

0.469 0.530 6.679

0.452 0.492 0.626

0.586 0.435 0.612

0.418

0.361 0.403 0.518

0.513 0.596 0.549 0.620

0.486

0.474 0.516

0.619

0.447 0.499

0.436

0.374

909.0 0.633 0.704

0.523 0.573 0.537

0.620 0.6470.620 0.665

SPIDER3-Single

0.590 0.547

0.596

0.586

0.573

0.582

 $\frac{Z}{C}$

ASA

 $\frac{Z}{C}$

HSE-U HSE-D

ASA

 C_{N}^{N}

HSE-U HSE-D

ASA

 $^{\rm Z}$

HSE-U HSE-D

ASA

 $\frac{Z}{C}$ 1

HSE-D

ASA

 $\frac{Z}{Z}$

HSE-U HSE-D

 $\frac{Z}{C}$

HSE-D

HSE-U

ASA

Model

CASP12-FM

SPOT-2018-HQ

SPOT-2018

SPOT-2016-HQ HSE-U

SPOT-2016 ASA

0.531 0.433 0.513

0.464 0.632

0.572

0.556 0.571

0.704

0.701

0.706 0.667

0.720

0.705 0.691

0.683

0.650

0.622

0.615

0.777

0.732

SPOT-1D (Profile) 0.787

SPOT-1D-Single

(This work)

Short length sequence(less than 1024)

0.627

0.487 909.0

0.479 0.595

0.538

0.637 0.726

0.496

0.417 0.537

0.585 0.545

0.510 0.563 0.737 $I \quad I \quad I \quad I$

I I I I

 $| \cdot |$

 $| \cdot |$ | |

 $| \cdot |$ | |

0.504 0.499 0.540 0.706

0.469

0.451 0.459 0.491 0.625

- 0.586 0.434 0.612

0.417

0.358 0.366

- 0.582 0.513 0.596

0.486

0.473

0.595 0.452 0.585

0.446

0.435

0.372 0.378 0.415 0.535

909.0 0.563 0.633 0.704

 $| \cdot |$

0.426 0.478 0.594

0.512 0.555

0.493

0.481 0.515

0.445

0.477

0.530 0.679

0.627 0.441 0.573

0.485 0.604

0.400

0.548 0.621

0.538

0.637

0.498

0.495

0.720

0.683 0.705 0.691 0.516

0.650

0.615 0.621 0.727

SPOT-1D (profile)

SPOT-1D-Single

ProteinUnet

SPIDER3-Single

Note: The values provided below are the Pearson's Correlation Coefficient (PCC) for the predicted ASA, HSE-U, HSE-D and CN.

Table 6. Performance comparison of SPOT-1D-Single with other predictors for seven different test sets TEST2018, SPOT-2016, SPOT-2016, SPOT-2018, SPOT-2018, SPOT-2018, DAT-2018, DAT-2018,

Model	TEST2018	018			SPOT	SPOT-2016			SPOT	SPOT-2016-HQ	O		SPOT-2018	.2018			SPOT-2	SPOT-2018-HQ	~	J	CASP12-FM	FM		J	CASP13-FM	FM		
	ψ	φ	θ ϕ h z θ ϕ h	2	*	φ	θ	2	*		θ	ь	*	φ	θ	ı	ψ	φ	θ	ь	*	φ	θ	Þ	ψ	φ	θ	t,
SPIDER3-Single 43.054 23.779 11.075 45.384 44.37 23.483 11.331 44.799 42.660 23.796 12.702 46.028 45.713 23.436 11.452 46.033 44.184 24.195 12.995 47.502 47.462 26.168 11.639 47.591 46.164 25.315 11.076 46.348	43.054	23.779	11.075	45.38	4 44.37	7 23.48	3 11.33	1 44.79	9 42.66	0 23.796	; 12.702	46.028	45.713	3 23.436	11.452	46.033	44.184	24.195	12.995	47.502 4	17.462 2	6.168 1	1.639	7.591 4	16.164 2	5.315 1	1.076 4	6.348
ProteinUnet	42.932	23.422	42.932 23.422 10.282 44.941 — —	44.94	1				1	I	I	I	I	1	I	I	I	1		- 46.527 25.942 10.949 46.259 46.884 25.036 10.380 46.093	16.527 2	5.942 1	0.949	6.259 4	16.884	5.036 1	0.380 4	6.093
SPOT-1D-Single 40.583 22.155 9.345 42.315 42.818 22.716 9.648 42.371 40.879 22.299 10.986 43.737 44.407 22.864 9.851 43.787 42.587 22.889 11.366 45.320 43.457 25.426 10.278 44.022 45.231 25.125 9.889 44.903	40.583	22.155	9.345	42.31	5 42.81	8 22.71	6 9.648	8 42.37	71 40.87	9 22.299	9 10.986	43.737	7 44.407	7 22.864	9.851	43.787	42.587	22.880	11.366	45.320 4	13.457 2	5.426 1	0.278	4.022 4	15.231 2	5.125	9.889	4.903
(this work)																												
SPOT-1D (profile) 24.871 16.886 6.914 25.944 32.725 20.030 8.211 32.085 27.971 18.285 9.132 31.234 34.790 20.390 8.515 34.011 28.824 18.688 9.428 32.224 33.962 21.844 8.700 33.114 28.489 20.238 7.551 27.867	24.871	16.886	6.914	25.94	4 32.72	5 20.03	0 8.211	1 32.08	15 27.97	1 18.285	5 9.132	31.234	1 34.790) 20.396	8.515	34.011	28.824	18.688	9.428	32.224	3.962 2	1.844	8.700	3.114 2	28.489 2	0.238	7.551 2	7.867
Short length sequence (less than 1024)	suce (less	than 10)24)																									
SPIDER3-Single	I	I	I	I	44.17	0 23.44	0 11.37	73 44.65	6 45.69	44.170 23.440 11.373 44.656 42.695 23.933 12.798 46.111 45.640 23.480 11.520 46.043 44.051 24.264 13.061 47.424	3 12.798	46.111	45.640) 23.480	11.520	46.043	44.051	24.264	13.061	47.424	1	1	I	I	I	I	I	1
ProteinUnet	I	I	I	I	43.38	4 23.06	7 10.28	6 43.61	2 41.83	43.384 23.067 10.286 43.612 41.837 23.278 11.777 0.512 44.866 23.189 10.493 44.948 43.282 23.699 12.120 46.484	3 11.777	7 0.512	44.866	5 23.189	10.493	44.948	43.282	23.699	12.120	46.484	I	I	I	I	I	I	I	ı
SPOT-1D-Single	I	I	I	I	42.52	8 22.66	9 9.645	9 42.13	13 40.93	42.528 22.669 9.649 42.133 40.931 22.450 11.102 43.858 44.252 22.921 9.884 43.670 42.624 23.011 11.466 45.393) 11.102	43.858	3 44.252	2 22.921	9.884	43.670	42.624	23.011	11.466	45.393	I	I	I	I	I	I	I	I
(this work)																												
SPOT-1D (profile)	1	I	I	I	32.08	4 19.83	7 8.153	3 31.45	0 27.86	32.084 19.837 8.153 31.450 27.865 18.367 9.209 31.198 34.445 20.327 8.498 33.640 28.875 18.782 9.512 32.311	7 9.209	31.198	34.445	5 20.327	8.498	33.640	28.875	18.782	9.512	32.311	I	I	ı	I	I	I	ı	I

Note: The values provided below are the mean absolute errors (MAE) for the predicted ψ , ϕ , θ and τ .

SPOT-1D-Single 3471

Table 7. Inference time comparison of SPOT-1D-Single, ProteinUnet and SPIDER3-Single for prediction on 250 proteins of TEST2018

Computational specifications	SPIDER3-Single	ProteinUnet	SPOT-1D-Single
16 CPU threads on Intel(R) Xeon(R) CPU E5-2620 v4 @ 2.10 GHz	311 s	109 s	155 s
GeForce GTX 1080 Ti		121 s	56 s

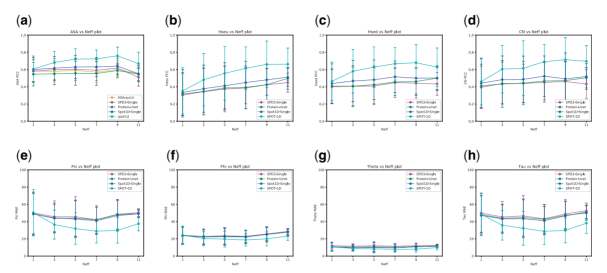


Fig. 3. Prediction accuracy as a function of the number of effective homologous sequence (Neff) by SPOT-1D-Single compared with other methods on SPOT-2018-short as labeled for (a) solvent accessibility (ASA) prediction, (b) half-sphere-exposure-up (HSE-u) prediction, (c) half-sphere-exposure-down (HSE-d) prediction, (d) contact number (CN) prediction, (e) ψ angle prediction, (f) ϕ angle prediction, (g) θ angle prediction and (h) τ angle prediction

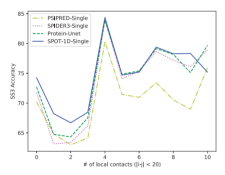


Fig. 4. Performance comparison between SPOT-1D-Single and SPIDER3-Single for secondary structure prediction (SS3) on all four test sets combined (TEST2018, SPOT-2016, CASP12-FM and CASP13-FM) as a function of the number of locally (short range) contacting residues ($|i\text{-}i| \leq 20$)

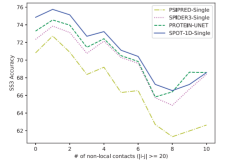


Fig. 5. Performance comparison between SPOT-1D-Single and SPIDER3-Single for secondary structure prediction (SS3) on all four test sets combined (TEST2018, SPOT-2016, CASP12-FM and CASP13-FM) as a function of the number of non-locally (long range) contacting residues $(|i \cdot j| \geq 20)$

CPU and GPU, respectively. By comparison, SPIDER3-Single's standalone version which only runs on the CPU takes nearly doubled time. ProteinUnet is faster on CPU but slower on GPU than SPOT-1D-Single.

SPOT-1D-Single predicts the secondary structure and 1D properties without using evolutionary features. The next improvement in protein secondary structure prediction and 1D prediction without using evolutionary features may come from using recent developments in feature generation using deep learning-based unsupervised learning features or protein embedding (Heinzinger *et al.*, 2019; Rao *et al.*, 2019; Rives *et al.*, 2021). It can be of interest to see how our models perform when trained on embedding instead of single-sequence.

Acknowledgements

We gratefully acknowledge the use of the High Performance Computing Cluster Gowonda to complete this research, and the aid of the research cloud resources provided by the Queensland Cyber Infrastructure Foundation (QCIF). We also gratefully acknowledge the support of NVIDIA Corporation with the donation of the Titan V GPU used for this research.

Funding

This work was supported by the Australian Research Council (DP180102060 and DP210101875 to Y.Z. and K.P.).

Conflict of Interest: none declared.

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3472 J.Singh et al.

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