



# DeepDISOBind: accurate prediction of RNA-, DNA- and protein-binding intrinsically disordered residues with deep multi-task learning

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## Abstract

Proteins with intrinsically disordered regions (IDRs) are common among eukaryotes. Many IDRs interact with nucleic acids and proteins. Annotation of these interactions is supported by computational predictors, but to date, only one tool that predicts interactions with nucleic acids was released, and recent assessments demonstrate that current predictors offer modest levels of accuracy. We have developed DeepDISOBind, an innovative deep multi-task architecture that accurately predicts deoxyribonucleic acid (DNA)-, ribonucleic acid (RNA)- and protein-binding IDRs from protein sequences. DeepDISOBind relies on an information-rich sequence profile that is processed by an innovative multi-task deep neural network, where subsequent layers are gradually specialized to predict interactions with specific partner types. The common input layer links to a layer that differentiates protein- and nucleic acid-binding, which further links to layers that discriminate between DNA and RNA interactions. Empirical tests show that this multi-task design provides statistically significant gains in predictive quality across the three partner types when compared to a single-task design and a representative selection of the existing methods that cover both disorder- and structure-trained tools. Analysis of the predictions on the human proteome reveals that DeepDISOBind predictions can be encoded into protein-level propensities that accurately predict DNA- and RNA-binding proteins and protein hubs. DeepDISOBind is available at <https://www.csuligroup.com/DeepDISOBind/>

**Keywords:** intrinsic disorder, protein–protein interactions, protein–nucleic acids interactions, deep learning

## Introduction

Intrinsically disordered regions (IDRs) lack stable tertiary structures and form dynamic conformational ensembles under physiological conditions [1, 2]. Recent bioinformatics studies reveal that disorder is highly abundant in nature [3], with about 20% of residues in eukaryotic proteins estimated to be disordered [4]. Proteins with IDRs are involved in a variety of cellular functions [5, 6]. Many IDRs interact with partner molecules, including deoxyribonucleic acid (DNA), ribonucleic acid (RNA) and proteins [7–13]. More specifically, the version 8.1 of the DisProt database [14], the primary repository of the intrinsic disorder, includes 1652 interacting IDRs, which constitute 42% of the IDRs annotated in this resource. Close to 90% (1473 out of 1652) of the interacting IDRs bind to proteins and nucleic acids. However, DisProt altogether

covers only about 1700 proteins, while millions of protein sequences await annotation of the interacting IDRs.

Computational predictors of interacting IDRs assist with closing this huge and growing annotation gap [15]. Based on an extensive literature search [15–19], we identified 22 predictors of the interacting IDRs. Nearly all of them (19 out of 22) predict a subfamily of the protein-binding IDRs called molecular recognition features (MoRFs) [20]. MoRFs are short IDRs that undergo folding upon interaction with protein partners. Some of the popular MoRF predictors include MoRFpred [21, 22], fMoRFpred [20], DISOPRED3 [23], MoRFChiBi [24], MoRFChiBiLight [25], OPAL+ (2018) [26] and SPOT-MoRF [27]. The other three methods, ANCHOR [28], DisORDPbind [29, 30] and ANCHOR2 [31] predict a broad family of the protein-binding IDRs that encompass

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MoRFs. Moreover, DisoRDPbind is the only current tool that predicts IDRs that interact with DNA and RNA. These tools are frequently used to guide experimental studies and reveal novel functional insights. Just as an example, DisoRDPbind was recently used to study the SARS-CoV-2 proteome [32], decipher functions of genes from animal pathogens [33] and investigate specific proteins, such as CS-like zinc finger (FLZ) [34], spindle-defective protein 2 (SPD-2) [35], mixed lineage leukemia 4 [36] and heat shock factor 1 [37], some of which are associated with cancers and neurodegenerative diseases. The importance of these predictors is further underscored by the fact that Critical Assessment of protein Intrinsic Disorder (CAID) experiment, which is an equivalent of Critical Assessment of protein Structure Prediction, but for the disordered proteins, included assessment of methods that predict interacting (in a partner-agnostic way) IDRs [38]. The top performing tools in the recent CAID were ANCHOR2, DisoRDPbind and MoRFChiBiLight, but the organizers also noted that ‘substantial room for improvement remains’ [38], suggesting the need to develop more accurate predictors of the interacting IDRs.

The methods that offer the most relevant and accurate predictions of the interacting IDRs, ANCHOR2 and DisoRDPbind, rely on relatively simple predictive models. DisoRDPbind utilizes logistic regression, while ANCHOR2 uses biophysics-based scoring functions. Moreover, DisoRDPbind that predicts interactions with proteins, DNA and RNA applies three independent/concurrent regressors. This way, it misses the opportunity to model relations between the three types of interactions. For instance, residues that bind nucleic acids and proteins have higher relative solvent accessibility compared to the non-binding residues, while the nucleic acid-binding residues are often positively charged and more evolutionarily conserved than the protein-binding residues [39]. The fact that DisoRDPbind is the only tool that predicts nucleic acid-binding IDRs, combined with modest accuracy of the current predictors of interacting IDRs, motivate the development of more accurate solutions.

Furthermore, we note that some protein- and nucleic-interacting residues are located in the structured protein regions. Numerous methods target the prediction of structured interacting regions and they rely on the training data extracted from Protein Data Bank [39–45]. Recently published structure-trained tools include SPRINT [46], SSWRF [47], EL-SMURF [48] and SCRIBER [49], which predict protein-binding residues; RNABindRplus [50] and FastRNABindR [51] which predict RNA-binding residues; TargetDNA [52] and DNAPred [53] which predict DNA-binding residues; DRNAPred [54], NCBRPred [55] and BindN+ [56] which predict interactions with RNA and with DNA; and ProNA2020 [57] and MTDsites [58] which identify protein, DNA- and RNA-interacting regions. Interestingly, recent study reveals that the structure-trained predictors of protein-binding regions perform

poorly when used to predict protein-binding IDRs [59]. We further investigate this finding by evaluating the results produced by several recent and well-performing structure-trained predictors of the protein, DNA- and RNA-interacting residues on the corresponding disordered binding regions.

We introduce DeepDISOBind, a custom-designed multi-task deep neural network that accurately predicts DNA-, RNA- and protein-binding IDRs. Multi-task learning aims to improve predictive performance by using shared representations (i.e. common parts of the model) to predict related learning tasks (i.e. binding to different partners) [60, 61]. Recently, the multi-task models were shown to improve the predictive quality for bioinformatics problems, including prediction of cleavage sites [62] and inter-residue distances [63], when compared to the single-task models. We devise the multi-task architecture where subsequent layers progressively specialize to predict interactions with different partner types. We empirically compare this topology against a single-task implementation and a representative selection of the existing predictors. We compare DeepDISOBind against representative methods that predict protein and nucleic acid-binding IDRs as well as the structure-trained methods. We also assess the DeepDISOBind’s predictions on the human proteome and release our tool as a convenient webserver.

## Methods

### Datasets

We source the data for training and comparative assessment of our predictive model from DisProt [14]. DisProt annotates proteins with the experimentally validated IDRs, including IDRs that interact with proteins, DNA and RNA. We manually checked IDRs that were annotated in DisProt as nucleic acids, DNA- and RNA-binding using the underlying publication data listed in DisProt in order to classify them as DNA- and/or RNA-binding. This annotation work follows from parsing DisProt for a recent comparative survey [64]. We divide these proteins into three subsets that constitute training, validation and test datasets. We ensure that sequences in each dataset share low (<30%) similarity with the other datasets. We use training and validation datasets to design and optimize the predictive model and the set-aside (during design and optimization) test dataset to comparatively assess this model against other solutions. Using protocol from [64], we cluster the original set of proteins with CD-HIT [65] at 30% sequence similarity and we place the entire protein clusters into training, validation and test datasets. The test and combined training/validation datasets share similar size, while the training dataset is set to be twice the size of the validation dataset. This procedure adheres to the commonly used practice in this field [64] and ensures a proper level of separation between the training/validation and test datasets (<30% sequence similarity). Detailed statistics, which cover distribution of

**Table 1.** Summary of datasets

Dataset	Number of proteins	Number of disordered residues				Number of all residues
		Protein-binding	DNA-binding	RNA-binding	All disordered	
Training	238	15 341 (14.5%)	2913 (2.7%)	1437 (1.4%)	27 304 (25.9%)	105 601
Validation	118	6464 (14.7%)	1284 (2.9%)	608 (1.4%)	11 716 (26.8%)	43 776
Test	394	17 540 (8.4%)	2377 (1.1%)	1518 (0.7%)	46 041 (22.2%)	207 743

RNA-/DNA-/protein-binding residues in the three datasets, are shown in Table 1. The datasets, including annotations of the DNA-, RNA- and protein-interacting IDRs, are freely available at <https://www.csuligroup.com/DeepDISOBind/>. We note that these datasets are larger than the datasets that are used to train and test DisoRDPbind [29] and are on par with the size of datasets utilized in CAID [38].

### Evaluation criteria

DeepDISOBind and other related tools produce putative propensities for the disordered DNA-, RNA- and protein-binding interactions for each residue in the input protein sequences. These real-valued propensities are accompanied by binary predictions, i.e. residues are classified as either DNA-/RNA-/protein-interacting or non-DNA-/RNA-/protein-interacting. The binary predictions are derived from the propensities by thresholding, i.e. residues with propensities greater than threshold are assumed to interact, while the remaining residues are assumed not to interact. Following related works [29, 59], we calibrate the thresholds for all considered predictors such that their binary predictions produce to the same specificity=0.8. Specificity is the rate of predictions of the interacting residues among the native non-interacting residues. We select 0.8 since it approximates the combined rate of the interacting residues across the three partner types. This calibration facilitates direct comparison of the binary predictions across different methods. Moreover, Table 1 reveals that the rates of the DNA- and RNA-interacting residues are much smaller than the rates of the protein-interacting residues. Thus, we further calibrate the evaluation between the three partner types by randomly undersampling the non-binding residues when evaluating performance for the RNA and DNA interactions so that their rate is the same as for the protein interactions. We assess the binary predictions with two popular metrics:  $F1 = (2 * TP) / (2 * TP + FN + FP)$  and  $sensitivity = TP / (TP + FN)$ , where TP is the number of correctly predicted protein-/RNA-/DNA-interacting residues, TN is the number of correctly identified non-protein-/RNA-/DNA-interacting residues, FN is the number of protein-/RNA-/DNA-interacting residues incorrectly predicted as non-interacting and FP is the number of the non-interacting residues incorrectly

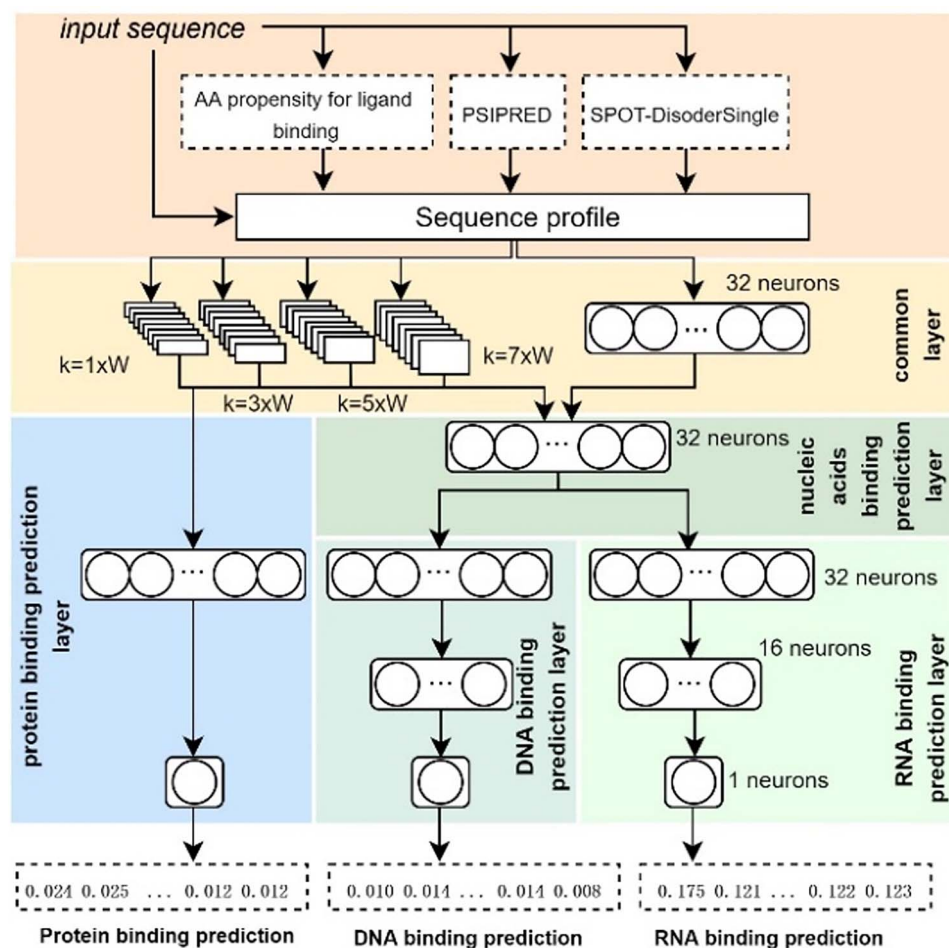
predicted as protein-/RNA-/DNA-interacting. We assess the predicted propensities with a commonly used area under the receiver operating characteristics (ROC) curve (AUC) that plots sensitivity against  $FPR = FP / (FP + TN)$ . Higher values of the three metrics (F1, sensitivity and AUC) indicate better predictive quality. In addition, since some residues interact with more than one partner, we evaluate predictors that provide protein-, DNA-, and RNA-binding predictions with the macro-average and micro-average metrics which are used in related multi-label predictions' studies [66–68]:

$$\begin{aligned} \text{micro - sensitivity} &= \frac{TP_{\text{avg}}}{TP_{\text{avg}} + FN_{\text{avg}}}, \\ \text{micro - F1} &= \frac{2 * TP_{\text{avg}}}{2 * TP_{\text{avg}} + FP_{\text{avg}} + FN_{\text{avg}}}, \\ \text{macro - sensitivity} &= \frac{1}{N} \sum \frac{TP_i}{TP_i + FN_i}, \\ \text{macro - F1} &= \frac{2}{N} \frac{\sum TP_i / TP_i + FN_i * \sum TP_i / TP_i + FP_i}{\sum TP_i / TP_i + FN_i + \sum TP_i / TP_i + FP_i}, \end{aligned}$$

where  $TP_{\text{avg}}$  is the average number of correctly identified protein-, DNA- and RNA-interacting residues,  $FN_{\text{avg}}$  is the average number of protein-/RNA-/DNA-interacting residues incorrectly predicted as non-interacting,  $FP_{\text{avg}}$  is the average number of the non-interacting residues incorrectly identified as protein-/DNA-/RNA-interacting,  $TP_i$  is the number of correctly predicted protein-, DNA- or RNA-binding residues,  $FN_i$  is the number of protein-/RNA-/DNA-interacting residues incorrectly predicted as non-interacting,  $FP_i$  is the number of incorrectly identified as protein/DNA/RNA interactions and  $i$  represents RNA, DNA and protein interaction labels.

### The DeepDIOSBind predictor

DeepDISOBind is a multi-task deep neural network that concomitantly predicts IDRs that interact with proteins, DNA and RNA (Figure 1). We use a custom-defined sequence profile that is extracted directly from the protein sequence as the input. Subsequent layers of the DeepDISOBind's network progressively specialize to predict interactions with different partner types. Correspondingly, the network is composed of five major elements (Figure 1): the common layer, the nucleic acid-binding layer, the protein-binding layer, the DNA-binding



**Figure 1.** The multi-task topology of the DeepDISOBind predictor.

layer and the RNA-binding layer. Following, we provide a more detailed description of the sequence profile and network topology.

### Sequence profile

Inspired by other recent models in this area [23, 27, 29, 69], the input protein sequence is first converted into a multi-dimensional profile. The profile covers the sequence itself together with relevant sequence-derived structural and functional properties that include relative amino acid propensities (RAAPs) for ligand-binding and predicted secondary structure and disorder. We use the one-hot encoding to represent the sequence. More specifically, each amino acid in the input sequence is represented by the 20-dimensional vector, where the position of the corresponding amino acid type is set to 1 while the other positions are set to 0. Moreover, we compute the maximum, minimum and average of the sequence embedding vectors that are defined in [70]. Inspired by recent studies that introduce novel predictors of the protein-binding residues from structured/ordered proteins [49, 71], we use RAAP for ligand-binding. These scores are derived empirically from binding data and quantify propensities of each amino acid type to bind a specific type of ligand. We use the five RAAP scales for

the protein- and nucleic acid-binding, which have been described in Table 3 in [39]. Finally, we use popular and fast predictors of the secondary structure, the single-sequence version of PSIPRED [72], and of the intrinsic disorder, SPOT-Disorder Single [73]. PSIPRED generates the three-state secondary structures (helix, strand and coil), which we represent with the one-hot encoding. SPOT-Disorder Single produces real-valued propensities and binary predictions of disorder. Altogether, the profile includes 33 dimensions: 20 for one-hot encoding of sequence +3 sequence embedding values +5 RAAP values +3 secondary structure predictions +2 disorder predictions. Similar to the other solutions in this area [27, 29, 69, 73–75], we use sliding windows to predict the interaction propensity for the residues in the middle of the windows. We pad the windows at the sequence termini with zeros.

### Architecture of the DeepDISOBind network

The underlying idea is to initially model a generic set of interacting residues and progressively specialize the network to more specific interacting partners. To this end, the partner-agnostic common layer (yellow block in Figure 1) links to layers that discriminate protein- and nucleic acid-binding (blue and green blocks in Figure 1),



while the latter layer further connects to layers that distinguish between DNA and RNA interactions.

The first, common layer consists of convolutional neural network (CNN) and feed-forward neural network (FNN) modules. The CNN module is composed of four different kernels that differ in size ( $k=1, 3, 5$  and  $7$ ). The variable kernel size designs were shown to be effective to reproduce the sequential nature of the protein sequences by accommodating for varying sizes of the residue neighborhoods, leading to improvements in predictive performance when compared to more traditional network architectures [70, 76–78]. We use eight channels for each kernel which are followed by ReLU activation units and a 1D max-pooling layer. We utilize the 1D max-pooling layer to reduce the dimension of the latent feature spaces before they are passed to the subsequent layers. Since the CNN module focuses specifically on local information (in a small sequence neighborhood around the predicted residue), we supplement it with the FNN module that extracts information from a larger window. This module uses a layer of  $n=32$  ReLU activation units that work in parallel to the CNN module. The outputs of the CNN and FNN modules are combined and fed into the subsequent FNN layers that aim to specialize the latent feature space produced in the common layer to specific types of interactions. We use four of these layers. First, the common layer is linked to the protein-binding and the nucleic acid-binding layers. Next, the nucleic acid-binding layer is linked to the DNA-binding and RNA-binding layers. We fix the sizes of the protein, DNA and RNA layers to  $n=32$  units, and we add additional sub-layers (smaller by a factor of 2) into the DNA and RNA layers. Consequently, RNA and DNA elements consist of two fully connected sub-layers,  $n=32$  and  $n/2=16$  units. The latter is motivated by the fact that DNA and RNA interactions are harder to differentiate compared to the nucleic acids and protein interactions [39]. Finally, the output layer that generates putative propensities for disordered RNA, DNA and protein interactions consists of three neurons implemented with the sigmoid transfer function.

Learning of the multi-task network requires a more specialized strategy compared to classical single-task networks. This is because some of the tasks (interactions) could be easier to optimize compared to the other tasks. This can be solved by relative weighting between tasks. We use a recently proposed tuning that relies on estimating uncertainty of each task [79]. Under this approach, if the performance of two tasks improves and the reduction of the other task gets worse by no more than  $\varepsilon$  (we set  $\varepsilon$  to a small value of 0.1), then we continue training the model. Otherwise, we stop the training process. Moreover, we adopt early stopping approach to avoid overfitting the training dataset.

We empirically investigate the impact of the selection of the hyperparameter  $n$  (size of the FNN modules in the common, protein, nucleic acids, DNA and RNA

layers) on the predictive performance. We consider networks with  $n=16$  (small size),  $n=32$  (medium),  $n=64$  (large) and  $n=256$  (very large). We summarize the corresponding topologies in [Supplementary Table S1](#) available online at <http://bib.oxfordjournals.org/>. We also empirically compare learning of the complete networks with the dropout learning [80] across the different network sizes. We set the dropout rate to 0.2. The dropout is meant to prevent overfitting, which would be apparent if the dropout-based learned networks would provide superior results. We compare the results on the validation dataset across different network sizes and when learning with and without the dropout on the training dataset in [Supplementary Table S2](#), available online at <http://bib.oxfordjournals.org/>. The average (across the three interaction types and three training runs) AUC ranges between 0.759 (small network with dropout) and 0.791 (medium network without dropout). Similarly, the average F1 varies between 0.238 (small network with dropout) and 0.271 (medium network without dropout). We observe that the averaged AUC and F1 scores are highly correlated (Pearson correlation of 0.95), which means that the considered networks produce high-quality propensities that are used to generate similarly accurate binary predictions. The medium size networks produce slightly better results than the small and large networks. Further increasing the size to the very large does not improve over the large-size networks. This means that the medium size networks are sufficiently large for this prediction. Lastly, we find that use of dropout does not lead to improvements. This together with the observation that modest-sized network produces the best results and outperforms the very large network suggest that our design does not overfit the training dataset. Consequently, we implement DeepDISOBind based on the medium network size ( $n=32$ ) and using training without dropout.

We also compare the above architecture that combines CNN and FNN modules with a design that relies on the graph neural network (GNN). GNNs were recently used in related projects that target prediction of protein–protein interactions (PPIs) at the protein level [81] and PPIs at the residue level from protein structure [82]. The corresponding underlying graphs represent the PPI networks and the spatial arrangement of amino acids in the protein structures. We use the graph to represent our input protein sequence, and more specifically, the sequential nature of connection between the residues in the input sliding window. The architecture of the GNN model draws from the best-performing medium size CNN/FNN network (i.e. DeepDISOBind) where we replace the CNN-based common layer with two graph convolutional layers, where nodes correspond to amino acids linked by peptide bonds, and we retain the other layers. [Table S2](#), available online at <http://bib.oxfordjournals.org/>, compares the results produced by this GNN model with the DeepDISOBind. The average AUC and F1 of the GNN-based design are modestly lower than the results

produced by the CNN-based DeepDISOBind; AUC of 0.756 versus 0.791 and F1 of 0.234 versus 0.271. This could be explained by the fact that the underlying graph is rather simple as it can only represent corrections between residues in the protein sequence compared to the CNN architecture that models these sequential relations more effectively. The more successful application of GNNs for the above-mentioned prediction of PPI networks and PPIs from protein structure stems from a more informative structure of the corresponding graphs.

## Results

### Ablation analysis of the network design

DeepDISOBind relies on two major elements: the multi-element sequence profile and the multi-task architecture. We investigate the relation between the specific formulation of these elements and the resulting predictive performance. We run ablation analysis where we measure predictive performance when removing certain parts of the profile and when we implement the topology as the collection of three single-task networks. The corresponding 10 versions of the predictive model are defined in [Supplementary Tables S3](#) and [S4](#), available online at <http://bib.oxfordjournals.org/>, i.e. modifications of the sequence profile and modifications of the topology, respectively.

We summarize the results of the ablation analysis on the test dataset in [Table 2](#). The top portion of the [Table 2](#) focuses on the sequence profile and reveals that all major parts of this profile that we employ provide useful information for the predictive model. More specifically, removal of the sequence, putative disorder or binding propensities (versions v1–v3) leads to a substantial drop in predictive performance from 0.75 to between 0.72 and 0.73 in the average AUC and from 0.56 to between 0.47 and 0.50 in the average sensitivity; we average over the three partner types. Removal of two or more parts of the profile (versions v4–v7) further deteriorates the performance, with the average AUC dropping to between 0.70 and 0.71. Interestingly, the v7 model that relies solely on the amino acid level propensities for binding (5-dimensional RAAP input) is comparable to the v6 model that uses the protein sequence [23-dimensional amino acid sequence (AAS) input], where both models secure the average AUC of 0.7. This shows that the RAAP scores provide a high-quality reduced representation of the sequence for the purpose of the prediction of the protein and nucleic acids interactions. [Supplementary Figure S1A](#), available online at <http://bib.oxfordjournals.org/>, provides the corresponding ROC curves. The curves demonstrate that DeepDISOBind offers particularly strong improvements over the models that exclude certain types of inputs for the low values of FPR (false positive rate)  $< 0.3$  ([Supplementary Figure S1B](#), available online at <http://bib.oxfordjournals.org/>). The increase in the sensitivity at the same FPR can be as high as 7%

when compared to the best input-reduced version. We argue that predictions with the low FPRs are more practical than the predictions with higher FPRs, given our imbalanced dataset where only about 20% of residues are interacting. In other words, FPRs of  $> 0.3$  would correspond to substantial overprediction of interactions. Altogether, these results indicate that all elements of the sequence profile contribute to the quality of predictions produced by the DeepDISOBind model.

We also study benefits of the application of the multi-task architecture by comparing it with the implementation that combines three single-task networks that use corresponding subsets of the layers from the original network and the same complete sequence profile ([Supplementary Table S4](#), available online at <http://bib.oxfordjournals.org/>). We summarize these results in the bottom section of [Table 2](#) (versions v8–v10). Each of the three single-task models underperforms when compared with DeepDISOBind. More specifically, the AUC for the prediction of the disordered protein interactions drops from 0.77 (DeepDISOBind) to 0.75 (single-task deep network), the AUC for the RNA interactions drops from 0.75 to 0.72 and the AUC for the DNA interactions decreases from 0.74 to 0.70. Moreover, average (over the three types of interactions) F1 and sensitivity (measured as the same specificity = 0.8) are reduced from 0.30 and 0.56 to 0.25 and 0.46, respectively, when comparing the multi-task and the single-task networks. This suggests that the use of the multi-task design leads to substantial improvements in the predictive performance across the three types of the interactions. This conclusion is in agreement with literature that similarly demonstrates that the multi-task learning improves over the single-task learning in a generic machine learning setting [61, 83] as well as when applied to bioinformatics problems [62, 63, 84]. We note that the multi-task learning was not previously used for the prediction of the disordered protein–protein and protein–nucleic acids interactions.

### Comparative assessment of predictive performance between DeepDISOBind and related methods

We compare results produced by DeepDISOBind with other relevant and representative methods that predict protein, DNA and RNA interactions from protein sequences. These methods include the only other tool that predicts disordered protein, DNA and RNA interactions, DisoRDPbind [29] and two popular and accurate predictors of the disordered protein interactions, ANCHOR2 [31] and MoRFChiBiLight [25]. These methods secured the top three spots in the assessment of the prediction of interacting IDRs in the recent CAID experiment [38]. We also include a comprehensive selection of the structure-trained predictors, including SCRIBER [49], which predicts protein-binding residues and which was recently shown to outperform other structure-trained predictors of protein-interacting residues [59];

**Table 2.** Ablation analysis for the DeepDISOBind predictor on the test dataset

Ablation design	Model	Protein interactions			RNA interactions			DNA interactions			Average			
		AUC	Sensitivity	F1	AUC	Sensitivity	F1	AUC	Sensitivity	F1	AUC	Sensitivity	F1	
Exclusion of inputs from the profile	DeepDISOBind	0.77	0.60		0.31	0.75	0.61	0.32	0.74	0.47	0.26	<b>0.75</b>	<b>0.56</b>	<b>0.30</b>
	v1 (excludes AAS)	0.75	0.55		0.29	0.74	0.52	0.28	0.70	0.44	0.24	<b>0.73</b>	<b>0.50</b>	<b>0.27</b>
	v2 (excludes PID)	0.74	0.53		0.28	0.69	0.43	0.24	0.72	0.46	0.25	<b>0.72</b>	<b>0.47</b>	<b>0.26</b>
	v3 (excludes RAAP)	0.77	0.55		0.29	0.67	0.46	0.25	0.73	0.40	0.22	<b>0.72</b>	<b>0.47</b>	<b>0.25</b>
	v4 (excludes AAS and RAAP)	0.76	0.56		0.30	0.68	0.40	0.22	0.70	0.46	0.25	<b>0.71</b>	<b>0.47</b>	<b>0.26</b>
	v5 (excludes PSS and PID)	0.72	0.53		0.28	0.68	0.45	0.25	0.70	0.42	0.23	<b>0.70</b>	<b>0.47</b>	<b>0.25</b>
	v6 (excludes RAAP, PSS and PID)	0.71	0.45		0.25	0.67	0.38	0.21	0.72	0.43	0.24	<b>0.70</b>	<b>0.42</b>	<b>0.23</b>
Single-task prediction	v7 (excludes AAS, PSS and PID)	0.69	0.47		0.25	0.72	0.51	0.28	0.68	0.48	0.26	<b>0.70</b>	<b>0.49</b>	<b>0.26</b>
	v8 (single-task prediction of protein-binding)	0.75	0.51		0.27	N/A	N/A	N/A	N/A	N/A	N/A			
	v9 (single-task prediction of RNA-binding)	N/A	N/A		N/A	0.72	0.44	0.24	N/A	N/A	N/A	<b>0.72</b>	<b>0.46</b>	<b>0.25</b>
	v10 (single-task prediction of DNA-binding)	N/A	N/A		N/A	N/A	N/A	N/A	0.70	0.44	0.24			

Note. We compare the complete DeepDISOBind model against 10 versions where we remove specific parts of the sequence profile (v1–v7) and where we implement the model as the combination of three single-task networks (versions v8–v10). [Supplementary Tables S3 and S4](#), available online at <http://bib.oxfordjournals.org/>, define further details. The profile includes AAS, RAAP for binding, putative secondary structure (PSS) and putative intrinsic disorder (PID). Sensitivity and F1 are calibrated to the same specificity=0.8. The last set of columns shown in bold font shows the average values over the three types of the partner molecules.

RNABindRPlus [50] that was ranked as the best tool in the recent assessment of the structure-trained predictors of the RNA interactions [45]; TargetDNA [52], one of the most accurate and popular predictors of the DNA interactions in the structured regions [85]; two representative structure-trained methods that predict DNA- and RNA-binding regions, popular BindN+ [56] that was shown to provide strong predictive performance in comparative surveys [39, 43] and one of the most recent methods, NCBRPred [55]; and two structure-trained methods which target prediction of protein, DNA and RNA interactions, ProNA2020 that was released in 2020 [57] and MTDsites that was published in 2021 [58]. The latter two methods offer the same scope of predictions as DeepDISOBind and DisoRDPbind, but they address predictions for structured rather than disordered regions. We use the author-provided webserver or implementations to make the predictions for these ten tools: DisoRDPbind, ANCHOR2, MoRFChibiLight, SCRIBER, RNABindRPlus, TargetDNA, BindN+, NCBRPred, ProNA2020 and MTDsites.

We compare results produced by DeepDISOBind with the 10 representative tools and our implementation that is based on the single-task networks on the test dataset, as shown in [Table 3](#). We empirically assess whether DeepDISOBind offers statistically significant improvements over the other solutions that are robust across different datasets. We bootstrap 50% of the test proteins for 50 times and compare the corresponding results with the t-test (for normal measurements) or with the Wilcoxon test (otherwise). We test normality with the Kolmogorov–Smirnov test at the *P*-value of 0.05. Similar tests were done in related comparative studies [45, 64, 86]. [Table 3](#) reveals that DeepDISOBind consistently secures the best predictive performance across the three binding partner types and the three metrics of performance. Moreover, the improvements in AUC, sensitivity and F1 are statistically significant

compared to each of the 10 other methods for the predictions of protein, DNA and RNA interactions (*P*-value < 0.05).

The average AUC, sensitivity and F1 (computed over the three interactions) for DeepDISOBind are 0.75, 0.56 and 0.30 compared the other three tools that provide the same scope of predictions that covers protein, DNA and RNA interactions: DisoRDPbind (0.66, 0.42 and 0.23), ProNA2020 (0.47, 0.28 and 0.13) and MTDsites (0.64, 0.35 and 0.22). The corresponding ROC curves for these four predictors are separated by a relatively wide margin ([Supplementary Figure S2](#), available online at <http://bib.oxfordjournals.org/>). We also assess multi-label predictions for these four methods using the macro-average and micro-average metrics ([Table 3](#)). Consistent with the single-label assessment, DeepDISOBind outperforms the other three predictors by securing macro-F1 of 0.30, macro-sensitivity of 0.56, micro-F1 of 0.30 and micro-sensitivity of 0.58. These results are statistically better than the results of the three other methods (*P*-value < 0.05), with the second-best DisoRDPbind that obtains macro-F1 of 0.23, macro-sensitivity of 0.43, micro-F1 of 0.25 and micro-sensitivity of 0.46. The predictive performance of MTDsites and ProNA2020 is worse than DeepDISOBind and DisoRDPbind since the two former methods are trained using structured proteins. The lower predictive quality of these tools for the prediction of interactions in the IDR is in agreement with similar observations obtained in a recent comparative survey of the disorder-trained and structure-trained predictors of protein-binding residues [59].

For the disordered protein interactions' prediction, the sensitivity of DeepDISOBind is better by (0.595–0.456)/0.456 = 30.5%, 190.2%, 95.7%, 40.5%, 18.8% and 18.5% when compared with DisoRDPbind, ProNA2020, MTDsites, SCRIBER, ANCHOR2 and MoRFChibiLight, respectively. This means that DisoRDPbind correctly

**Table 3.** Comparative assessment on the test dataset

Predictive target	Method	Protein-binding			RNA-binding			DNA-binding			Multi-label macro-average		Multi-label micro-average	
		AUC	Sensitivity	F1	AUC	Sensitivity	F1	AUC	Sensitivity	F1	Sensitivity	F1	Sensitivity	F1
Protein-, DNA- and RNA-binding residues	DeepDISOBind	<b>0.771</b>	<b>0.595</b>	<b>0.313</b>	<b>0.746</b>	<b>0.611</b>	<b>0.320</b>	<b>0.736</b>	<b>0.472</b>	<b>0.255</b>	<b>0.559</b>	<b>0.297</b>	<b>0.580</b>	<b>0.305</b>
	Single-task predictor (combination of v8-v10)	0.746+	0.516+	0.277+	0.725+	0.446+	0.243+	0.697+	0.443+	0.242+	0.468+	0.254+	0.503+	0.271+
	DisoRDPbind	0.727+	0.456+	0.249+	0.594+	0.364+	0.202+	0.671+	0.452+	0.246+	0.426+	0.234+	0.457+	0.248+
	MTDsites	0.576+	0.304+	0.173+	0.677+	0.479+	0.258+	0.675+	0.253+	0.242+	0.406+	0.225+	0.322+	0.182+
	ProNA2020	0.398+	0.205+	0.120+	0.468+	0.193+	0.08+	0.551+	0.441+	0.187+	0.215+	0.132+	0.204+	0.120+
Protein-binding residues	ANCHOR2	0.719+	0.501+	0.270+										
	MoRFChibiLight	0.735+	0.502+	0.271+										
	SCRIBER	0.684+	0.423+	0.232+										
DNA and RNA-binding residues	BindN+				0.685+	0.473+	0.257+	0.615+	0.331+	0.187+				
	NCBRPred				0.662+	0.455+	0.243+	0.617+	0.367+	0.205+				
DNA-binding residues	TargetDNA						0.580+	0.274+	0.157+					
RNA-binding residues	RNABindRPlus				0.576+	0.336+	0.186+							

Note. The binary predictions use thresholds that equalize specificity to 0.8 across the methods to allow for direct comparisons (details in the Evaluation criteria section). + means that DeepDISOBind is statistically significantly better ( $P$ -value  $< 0.05$ ). = means that the difference between DeepDISOBind and another predictor is not significant ( $P$ -value  $\geq 0.05$ ). The best results for each column are shown in bold font.

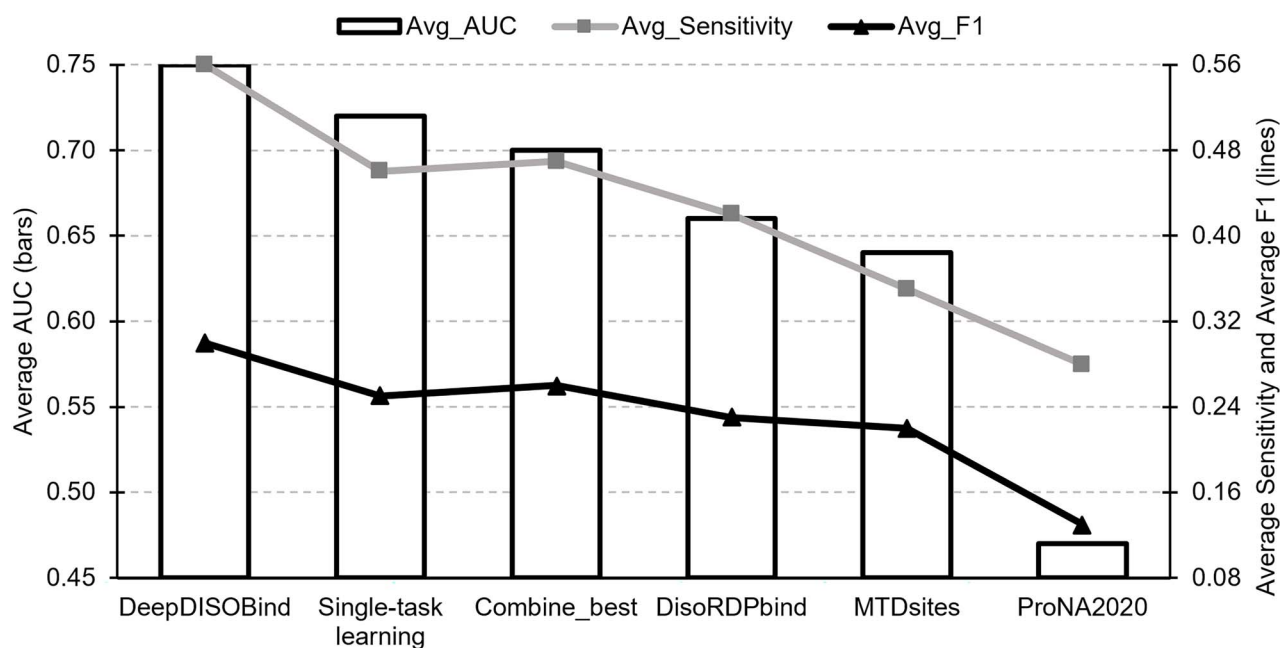
identifies at least 18.5% more interacting residues at the same false positive rate, i.e. we fix specificity at 0.8 for all methods, which corresponds to 0.2 false positive rate. Similarly, for the RNA interactions, DeepDISOBind's sensitivity is better by 67.9%, 216.5%, 27.5%, 34.3%, 29.2% and 81.8% when contrasted with DisoRDPbind, ProNA2020, MTDsites, NCBRPred, BindN+ and RNABindRPlus, respectively. The improvements in the sensitivity for the DNA interaction predictions are at 4.4%, 7.0%, 86.5%, 28.6%, 42.6% and 72.3% when compared against DisoRDPbind, ProNA2020, MTDsites, NCBRPred, BindN+ and TargetDNA, respectively. Similar observations are true when using the F1 and AUC metrics.

Figure 2 offers a more direct approach to compare DeepDISOBind with the state of the art. We compare the average values of the AUC (bars), sensitivity (gray line) and F1 (black line) computed over the three interaction types. The comparison includes DeepDISOBind, the single-task network (combination of the v8-v10 networks), the 'combine best' approach which uses the best method (i.e. having highest AUC) for each interaction type selected across the 10 predictors (i.e. MoRFChibiLight for the protein interactions; BindN+ for the RNA interactions and MTDsites for the DNA interactions); DisoRDPbind which is the only other disorder-trained predictor with the same scope as DeepDISOBind; and MTDsites and ProNA2020 which are the two recently published structure-trained methods that predict protein-, DNA- and RNA-interacting residues. First, we note a substantial and statistically significant (Table 3) improvement when contrasting the multi-task (DeepDISOBind) versus single-task solutions across the three metrics ( $P$ -value  $< 0.05$ ). Second, DeepDISOBind improves against the combination of the best current methods by a large and statistically significant margin (0.75 versus 0.70 in AUC, 0.56 versus 0.41 in sensitivity and 0.30 versus 0.26

in F1). Third, DeepDISOBind and the single-task networks outperform DisoRDPbind, primarily because the latter relies on simpler logistic regression models that are applied utilizing the single-task architecture. Lastly, DeepDISOBind improves over ProNA2020 and MTDsites because the latter two are trained on the structured proteins.

Finally, we investigate impact of similarity between the test proteins and the proteins that were used to train PSIPRED and SPOT-Disorder-Single methods, which we utilize to derive inputs for DeepDISOBind (Figure 1). We collect and combine the training datasets of these two predictors. Next, we align each test protein to every training protein with BLASTp [87] to annotate regions in the test proteins that share similarity  $> 30\%$ . Finally, we retest the predictive performance of DeepDISOBind and the other predictors of protein-, DNA- and RNA-binding residues on the test proteins when excluding the similar regions. We summarize these results in Supplementary Table S5, available online at <http://bib.oxfordjournals.org/>. DeepDISOBind secures results that are on average very similar to the results on the complete test dataset, with the average AUC (over the protein, DNA and RNA predictions) of 0.752 versus 0.751 and the average F1 of 0.295 versus 0.296. Moreover, DeepDISOBind's predictions consistently maintain statistically significant advantage over the results of the other 10 predictors ( $P$ -value  $< 0.05$ ). Altogether, the results on the complete test dataset and the sequence regions that share low similarity to the training data of PSIPRED and SPOT-Disorder-Single are similar. This could be explained by the fact that we use the single-sequence version of PSIPRED and the inherently single-sequence SPOT-Disorder-Single. Both methods do not use sequence alignment, thus minimizing the likelihood of overfitting training datasets [72, 73]. To sum up, the empirical analysis demonstrates that DeepDISOBind





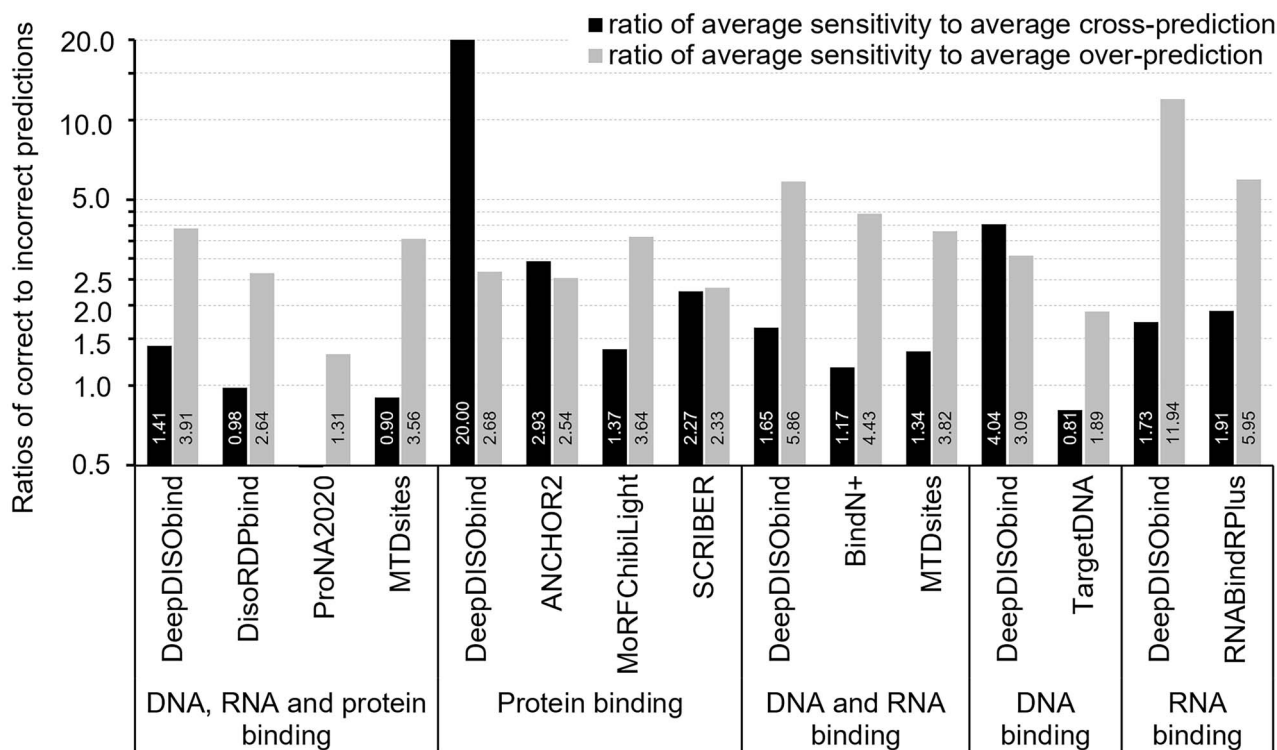
**Figure 2.** Comparison of the predictive performance on the test dataset between DeepDISOBind, MTDsites, ProNA2020, the single-task network (combination of the v8–v10 networks), the Combine\_best approach which uses the best method for each interaction type selected across the six predictors (i.e. MoRFChibiLight for the protein interactions, BindN+ for the RNA interactions and MTDsites for the DNA interactions) and DisoRDPbind. We quantify the predictive performance with the average (over the three interaction types) values of AUC (bars and vertical axis on the left), F1 and sensitivity (lines and vertical axis on the right).

provides accurate predictions of the disordered protein, DNA and RNA interactions.

### Assessment of cross-predictions and over-predictions

The binding residues share certain characteristics, such as high levels of evolutionary conservation and high solvent accessibility, irrespective of the type of their binding partners. This may lead to a substantial amount of cross-predictions, which are measured as the fraction of residues that bind a given partner type, which are predicted to interact with another ligand type, e.g. protein-binding residues predicted as DNA- or RNA-binding residues. Recent studies have found that majority of methods that predict interacting residues for the structured regions generate substantial amounts of cross-predictions, which in some cases, are as high as their sensitivity that quantifies the rate of correct predictions [42, 43, 59, 88]. Correspondingly, we assess the cross-predictions and over-predictions (fraction of non-binding residues predicted to interact with a given partner type) for DeepDISOBind and the other 10 considered to be predictors. Figure 3 quantifies the average (over the different partner types) ratios of sensitivity (rate of correct predictions) to the cross-prediction and over-prediction rates (rates of incorrect predictions) on the test dataset; ratios  $>1$  denote methods for which the rate of the correct predictions is higher than the rate of over- or cross-predictions. We normalize rate of predictions of binding residues across predictors to allow for side-by-side comparisons

of the ratios across methods, i.e. we equalize the number of the predicted protein-/DNA-/RNA-binding residues to the number of the native protein-/DNA-/RNA-binding residues. We provide the complete set of results including cross-prediction rates, over-prediction rates and sensitivity values for each partner type (protein, DNA and RNA) in Supplementary Table S6, available online at <http://bib.oxfordjournals.org/>. We compare DeepDISOBind to the other methods using the same set of predictions, e.g. we compare DeepDISOBind's predictions of protein-binding residues to the SCRIBER's, ANCHOR2's and MoRFChibiLight's results which also predict only the protein-binding residues. The ratios to the over-predictions are relatively high across all methods, ranging between 1.89 for TargetDNA and 11.94 for DeepDISOBind's prediction of the RNA-binding residues (gray bars in Figure 3). This means that relatively few non-binding residues are predicted to bind. We also observe that DeepDISOBind generates the highest/best ratios to the cross-predictions across all scenarios, except when compared for the RNA-binding prediction with RNABindRPlus where both methods achieve good results, 1.73 and 1.91 (black bars in Figure 3). Moreover, the DeepDISOBind's ratios are always  $>1$ , which means that that its rates of correct prediction of binding residues outperform the rates of the cross-predictions. We observe that relatively few protein-binding residues are incorrectly predicted as RNA-binding (7%) or DNA-binding (11%) compared to the corresponding average sensitivity (26%). Overall, when making predictions of the protein, DNA- and RNA-binding, the DeepDISOBind's



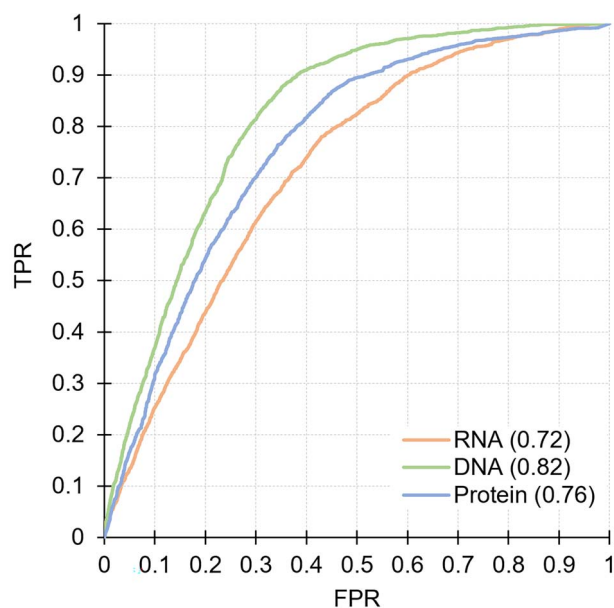
**Figure 3.** Comparison of the ratios of the average sensitivity (over the three interaction types) to the average cross-prediction and over-prediction rates on the test dataset. Larger ratios indicate higher quality predictions. The predictions rely on thresholds that equalize the number of the predicted binding residues with the number of native binding residues for each predictor. Predictors are grouped by the scope of their predictions, as described on the x-axis, where DeepDISOBind's predictions are limited to the predictions of the other methods in the same group.

ratio to cross-predictions equals 1.41. This means that its average rate of correct predictions is 40% higher than the rate of the cross-predictions, which is substantially better than the 0.98, 0.46 and 0.90 ratios secured by DisoRDPbind, ProNA2020 and MTDsites.

### Assessment of predictions in the human proteome

We assess DeepDISOBind's predictions on the proteome scale. While the coverage of the residue-/region-level annotations is limited at this scale, we can obtain a comprehensive set of experimental annotations at the protein level. We evaluate DeepDISOBind's predictions of the disordered DNA- and RNA-binding proteins in one of the most comprehensively annotated proteomes, the human proteome. To do that, we collect disordered human proteins that are annotated to interact with DNA and with RNA as well as the human proteins that are unlikely to interact with the nucleic acids. First, we collect the human proteome from UniProt version 2019\_09 [89] and remove partial sequences that we identify based on the "Sequence status" term "Fragment". This produces 43 789 protein sequences. Second, we annotate the DNA-interacting proteins by combining data from a comprehensive collection of relevant databases including 3D-footprint [90], CIS-BP [91], JASPAR [92], HumanTF2 [93], SMiLE-seq [94], animalTFDB [95] and the gene ontology (GO) terms [96] in UniProt. We also annotate the RNA-binding

proteins based on the data from ATTRACT [97], RBPDB [98] and the GO terms in UniProt. We map proteins in these diverse resources into the human set based on the UniProt's accession numbers. This results in 2379 DNA-binding and 2371 RNA-binding proteins, which is in line with related studies [99]. We identify the disordered subset of these proteins using the popular VSL2B predictor [100]. This method is different than the SPOT-Disorder-Single predictor used in DeepDISOBind and offers high-quality predictions of the disordered proteins [38, 64]. We annotate a given DNA-/RNA-binding protein as disordered if its putative disorder content is  $>0.2$ . Consequently, we identify 1739 and 1711 disordered DNA- and RNA-interacting proteins, respectively. Third, we derive proteins that are unlikely to interact with the nucleic acids. We select the human proteins that share  $<30\%$  sequence similarity with the annotated DNA- and RNA-binding proteins, which we quantify with BLASTp [87, 101]. This results in the set of 24 435 proteins. Finally, we convert the residue-/region-level propensities produced by DeepDISOBind into protein-level propensities of the disordered RNA and DNA interactions. Since typically only a small portion of the amino acids interact with the nucleic acids, we compute average of the highest 5% of the residue-level propensities produced by DeepDISOBind for a given protein to quantify the protein-level propensities. We emphasize that this approach does not validate correctness of the positions of the predicted binding residues in the protein



**Figure 4.** ROC curves for the DeepDISOBind's prediction of the DNA-interacting proteins (green line) and the RNA-interacting proteins (orange line) in the human proteome. The blue ROC curve is for the scenario where DeepDISOBind classifies disordered human hub proteins (proteins that interact with many proteins) versus human proteins that interact with a few protein partners.

sequence (which we assess in Comparative assessment of predictive performance between DeepDISOBind and related methods and Assessment of cross-predictions and over-predictions sections) but rather the ability to quantify propensity for binding at the whole protein level. We assess these protein-level predictions of the DNA- and RNA-interacting proteins in the human proteome with the ROC curves and the corresponding AUC scores (Figure 4). DeepDISOBind secures AUCs of 0.72 and 0.82 for the prediction of the human RNA- and DNA-interacting proteins, respectively, which are consistent with the results on the test dataset.

Moreover, motivated by the discussion in Assessment of cross-predictions and over-predictions section, we evaluate the potential for cross-predictions of the protein-level scores. We group the considered human proteins into four sets: (i) proteins that bind DNA and do not bind RNA, (ii) proteins that bind RNA and do not bind DNA, (iii) proteins that bind both RNA and DNA and (iv) proteins that do not bind neither DNA nor RNA. Next, we compare the protein-level scores for DNA and RNA interactions that we extract from DeepDISOBind's predictions (i.e. average of the highest 5% of the residue-level propensities) inside the sets 1–3 to study the cross-prediction. We utilize the pairwise t-test (for normal measurements) or the Wilcoxon test (otherwise), where we test normality with the Kolmogorov–Smirnov test at the 0.05  $P$ -value. The protein-level DNA-binding propensities are higher than the protein-level RNA propensities within the protein set 1 and the difference is statistically significant ( $P$ -value < 0.01). Similarly, the protein-level RNA-binding propensities are significantly higher than

the corresponding DNA propensities for the set 2 ( $P$ -value < 0.01). Interestingly, the protein-level RNA- and DNA-binding propensities are not significantly different for the protein set 3 ( $P$ -value = 0.66). These results suggest that the predictions of DeepDISOBind that we aggregate at the protein-level successfully differentiate between DNA- and RNA-binding proteins. Finally, we further examine the accuracy of the prediction of the DNA- and RNA-binding proteins by comparing the protein-level DNA-binding propensities between sets 1 and 4 and the protein-level RNA-binding propensities between sets 2 and 4. In both cases, the protein-level propensities for DNA and RNA interactions are higher in the sets 1 and 2, respectively, when compared with the set 4 and these differences are statistically significant ( $P$ -value < 0.01).

We also assess whether DeepDISOBind accurately predicts the disordered protein interactions. Since majority of human proteins interact with proteins, and thus, it would be virtually impossible to reliably identify non-protein-binding proteins; we use DeepDISOBind's predictions to differentiate between disordered hub proteins [102], which interact with many protein partners, and proteins that interact with relatively few proteins. This is motivated by the finding that the human hub proteins are enriched in the intrinsic disorder [103, 104]. First, we collected a comprehensive set of PPI annotations in the human proteome from the mentha resource, which combines data from several relevant source databases [105]. Second, we process the corresponding set of 17 598 protein-interacting proteins to extract the highly promiscuous hub proteins (25% of proteins with the highest PPI counts) and proteins that interact with a few protein partners (25% that interact with the smallest number of proteins). The same as for the assessment of the DNA/RNA interactions, we use VSL2B to identify a subset of the disordered hub proteins. Finally, we convert the residue-/region-level protein-binding propensities produced by DeepDISOBind into the protein-level propensities of the disordered proteins interactions using the same approach as for the assessment of the nucleic acid-binding proteins. Blue ROC curve in Figure 4 quantifies the predictive quality of DeepDISOBind applied to differentiate between the disordered hubs and the proteins that interact with few proteins. DeepDISOBind obtains AUC of 0.76, which is similar to the results on the test dataset. Altogether, these results suggest that the outputs produced by DeepDISOBind can be converted into protein-level scores that correctly predict disordered RNA-, DNA- and protein-interacting proteins. The DeepDISOBind's predictions for the human proteins are available at <https://www.csuligroup.com/DeepDISOBind/>.

### Case study

We illustrate DeepDISOBind's predictions on one of the test proteins, the silent information regulator Sir3p from budding yeast (DisProt: DP00533; UniProt: P06701). Sir3p is involved in the initiation, propagation and compaction of the silenced chromatin [106, 107].

Sir3p has a long IDR (positions 216–549) that interacts with RAP1p [108], RAD7p [109] and Sir4p coiled-coil domain [110]. This DNA- and protein-interacting IDR is flanked by structured regions that extend to the termini.

The case study aims to visualize the putative propensities and binary predictions produced by DeepDISOBind and the other methods that we cover in Comparative assessment of predictive performance between DeepDISOBind and related methods section and Table 3. This example is not intended to quantify or compare the predictive performance. Supplementary Figure S3A, available online at <http://bib.oxfordjournals.org/>, reveals that the disorder-trained predictors (ANCHOR2, MoRFChiBiLight, DisoRDBbind and DeepDISOBind) correctly identify this IDR as interacting with proteins. MoRFChiBiLight slightly overpredicts protein-interacting regions in the structured domain at the N-terminus. On the other hand, the structure-trained predictors of the protein-binding residues, such as SCRIBER, ProNA2020 and MTDsites, miss this binding region. This can be explained by the fact that they target prediction of protein-binding in structured regions. Supplementary Figure S3B, available online at <http://bib.oxfordjournals.org/>, shows that, while most methods (except for ProNA2020) correctly predict DNA interactions in this IDR, DisoRDPbind, TargetDNA, BindN+, MTDsites and NCBRPred overpredict DNA interactions outside this region. TargetDNA, BindN+, MTDsites and NCBRPred were designed to identify the interactions in the structured regions and this is likely why they make more predictions at both structured termini. Moreover, predictions from DeepDISOBind, DisoRDPbind, ProNA2020 and RNABindRPlus suggest that this protein is unlikely to interact with RNA, while BindN+, NCBRPred and MTDsites predict multiple RNA-binding regions (Supplementary Figure S3C, available online at <http://bib.oxfordjournals.org/>). This can be again attributed to the fact that BindN+, NCBRPred and MTDsites aim to predict RNA interactions in the structured regions.

### DeepDISOBind webserver

DeepDISOBind is available as a user-friendly webserver at <https://www.csuligroup.com/DeepDISOBind/>. With the user's convenience in mind, we make predictions on the server side and process up to 20 proteins in a single request. The only required inputs are the FASTA-formatted protein sequences. Users can opt to provide an email address where we send links to the results when predictions are completed. Predictions take about 30 seconds for an average-size sequence. The server outputs numeric propensities for the protein, RNA and DNA interactions and the three corresponding binary predictions for each residue in the input chain(s). We also provide putative propensities and binary annotations of disorder generated by SPOT-Disorder-Single. The results are available in three convenient formats: (i) parseable text file that can be downloaded from a request-specific

URL, (ii) color-coded (to ease identification of interacting residues) table in the browser window and (iii) an interactive graphical format in the browser window. We will store these predictions for at least 1 month. The graphical format allows users to select predictions of specific interactions, identify propensity scores, amino acid type and position on mouse hover and zoom on a specific protein segment. Users should employ the putative propensities as a measure of confidence, i.e. residues predicted with higher values of propensity are more likely to interact with the corresponding partner. Moreover, the binary predictions can be used to identify the putative protein-, RNA- and DNA-binding residues when assuming low false positive rate at 0.2; we use the same calibration in Tables 2 and 3 and in Supplementary Tables S5 and S6, available online at <http://bib.oxfordjournals.org/>.

Importantly, DeepDISOBind targets prediction of IDRs that interact with proteins, DNA and RNA and, by design, is not going to produce reliable predictions for the structured regions. Thus, predictions of the interacting residues for the structured regions, which can be identified with the help of the SPOT-Disorder-Single predictions, should be pursued with the structure-trained methods. Recent surveys of the structure-trained predictors can be used to identify suitable methods [39–45].

### Discussion

IDRs interact with a variety of partner molecules including nucleic acids and proteins. The availability of experimental data for hundreds of interacting IDRs gave rise to the development of machine learning models that learn from these data to predict these interactions for the millions of unannotated protein chains. However, only one such tool is available for the prediction of disordered interactions with the nucleic acids, and the recent CAID experiment concludes that new and more accurate predictors of the interacting regions are needed [38]. To this end, we develop DeepDISOBind, a novel multi-task deep learner, which provides accurate predictions of the DNA-, RNA- and protein-binding IDRs. We empirically demonstrate that our selection of the predictive inputs and the multi-task design of DeepDISOBind's model contribute to its predictive performance. Side-by-side evaluation on an independent (low similarity) test dataset reveals that DeepDISOBind offers statistically significant improvements over the single-task topology and a representative collection of 10 existing tools that cover both disorder-trained and structure-trained methods. These improvements are consistent across the three interactions types. Evaluation on the human proteome shows that DeepDISOBind accurately identifies hubs and DNA- and RNA-binding proteins. We provide a convenient webserver at <https://www.csuligroup.com/DeepDISOBind/>. This webserver allows for batch predictions, performs calculations on the server side and provides results



in multiple formats, including an interactive graphical visualization.

### Key Points

- CAID experiment shows that current predictors of disordered regions interacting with nucleic acids and proteins offer modest levels of predictive accuracy.
- DeepDISOBind uses an innovative deep multi-task architecture to accurately predict DNA-, RNA- and protein-binding disordered regions from protein sequences.
- DeepDISOBind's predictions outperform results of current disorder- and structure-trained methods across the interactions with DNA, RNA and protein partners.
- DeepDISOBind accurately identifies protein hubs and DNA- and RNA-binding proteins in the human proteome.
- DeepDISOBind's webserver is available at <https://www.csuligroup.com/DeepDISOBind/>.

### Supplementary data

Supplementary data are available online at <https://academic.oup.com/bib>.

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