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Sequence and structural features of carbohydrate binding in proteins and assessment of predictability using a neural network Adeel Malik and Shandar Ahmad*

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Abstract

Background: Protein-Carbohydrate interactions are crucial in many biological processes with implications to drug targeting and gene expression. Nature of protein-carbohydrate interactions may be studied at individual residue level by analyzing local sequence and structure environments in binding regions in comparison to non-binding regions, which provide an inherent control for such analyses. With an ultimate aim of predicting binding sites from sequence and structure, overall statistics of binding regions needs to be compiled. Sequence-based predictions of binding sites have been successfully applied to DNA-binding proteins in our earlier works. We aim to apply similar analysis to carbohydrate binding proteins. However, due to a relatively much smaller region of proteins taking part in such interactions, the methodology and results are significantly different. A comparison of protein-carbohydrate complexes has also been made with other protein-ligand complexes.

Results: We have compiled statistics of amino acid compositions in binding versus non-binding regions-general as well as in each different secondary structure conformation. Binding propensities of each of the 20 residue types and their structure features such as solvent accessibility, packing density and secondary structure have been calculated to assess their predisposition to carbohydrate interactions. Finally, evolutionary profiles of amino acid sequences have been used to predict binding sites using a neural network. Another set of neural networks was trained using information from single sequences and the prediction performance from the evolutionary profiles and single sequences were compared. Best of the neural network based prediction could achieve an 87% sensitivity of prediction at 23% specificity for all carbohydrate-binding sites, using evolutionary information. Single sequences gave 68% sensitivity and 55% specificity for the same data set. Sensitivity and specificity for a limited galactose binding data set were obtained as 63% and 79% respectively for evolutionary information and 62% and 68% sensitivity and specificity for single sequences. Propensity and other sequence and structural features of carbohydrate binding sites have also been compared with our similar extensive studies on DNA-binding proteins and also with protein-ligand complexes.

Conclusion: Carbohydrates typically show a preference to bind aromatic residues and most prominently tryptophan. Higher exposed surface area of binding sites indicates a role of hydrophobic interactions. Neural networks give a moderate success of prediction, which is expected to improve when structures of more protein-carbohydrate complexes become available in future.

Background

Carbohydrates are often referred to as the third molecular chain of life, after DNA and proteins [1]. These interactions are responsible for important biological functions such as inter cellular communication particularly in the immune system [2]. Living cells in all organisms are usually covered with one or another type of carbohydrate [2]. Some viruses like influenza, use sugars on the outside of human cells to gain entry. Sometimes the carbohydratebinding proteins and their sugar-ligands are expressed on the same cell, and the sugar is a part of the regulation machinery of the cell [3]. The functional roles of carbohydrates and their interactions with proteins are drawing more attention than before, since it has been recognized that carbohydrates are used as information carriers, rather than simple storage material [1]. Protein-carbohydrate interactions are involved in a broad range of biological processes. These processes include, among others, infection by invading microorganisms and the subsequent immune response, leukocytic trafficking and infiltration, and tumor metastasis [4-12]. Carbohydrates are uniquely suited for this role in molecular recognition, as they possess the capacity to generate an array of structurally diverse moieties from a relatively small number of monosaccharide units [13]. This could be attributed to the fact, that unlike the components of nucleic acids, carbohydrates can link together in multiple, nonlinear ways because each building block has about four functional groups for linkage. They can even form branched chains. Hence, the number of possible polysaccharides is enormous (Figure 1). Since carbohydrates assume a large variety of configurations, many carbohydrate-binding proteins are being considered as targets for new medicines.

In view of the above, accurate in silico identification of carbohydrate-binding sites is a key issue in genome annotation and drug targeting. The information about the factors, which prevent or support carbohydrate binding of an amino acid, is expected to be present in the evolutionary profile of the sequence as well as the identity and structure of amino acid residues in the neighborhood of potential carbohydrate binding sites. A number of reviews have been published on protein- carbohydrate interactions [14-18]. Different aspects of protein carbohydrate recognition have also been extensively studied [19-26]. However, bioinformatics approaches with a predictive goal are relatively rare [1,27]. Compared with the abundance of methodologies developed for protein-nucleic acid [28-32] or protein-protein interactions [33-38], there are still very few methods for predicting carbohydrate-protein interactions. Shionyu-Mitsuyama [1] has developed a program that uses the empirical rules of the spatial distribution of protein atoms at known carbohydrate-binding sites for prediction. In that work an analysis of the characteristic properties of sugar binding sites was performed on

a set of 19 sugar binding proteins. For each site six parameters were evaluated viz. solvation potential, residue propensity, hydrophobicity, planarity, protrusion and relative accessible surface area. Three of the parameters were found to distinguish the observed sugar binding sites from the other surface patches. These parameters were then used to calculate the probability for a surface patch to be a carbohydrate-binding site [27]. These prediction methods are based on local structural descriptors of proteins and cannot be used if complete 3 dimensional structures are not available. On the other hand, neural network based predictions of post-translational modification (Oglycosylation and phosphorylation) sites have been reported by two groups [39,40]. However, these studies are restricted to only one type of protein-carbohydrate interactions and therefore do not capture all protein-carbohydrate interactions, as sought out in this work.

In this work we explore the exact contribution from different sequence and evolutionary attributes of proteins in determining their carbohydrate binding regions. Propensity of each of the 20 amino acid residues in binding regions has been calculated and compared with non-binding regions. Solvent accessibility, secondary structure and packing density of binding sites have been analyzed in a similar way. We go on to design a neural network to model sequence and evolutionary information (obtained by Position Specific Scoring Matrices) and determine their role in the predictability of carbohydrate binding sites.

We also study the binding sites of other protein-ligand and protein-DNA complexes and compare the propensity scores of all residues and their secondary structures with protein-carbohydrate complexes.

Results and Discussion Residue-wise propensity scores

We started with a non-redundant set of all carbohydrate binding proteins (Procarb40) collected from PDB as described in the Methods section. Residue-wise propensities of carbohydrate binding sites in Procarb40 dataset were calculated and compared with the propensities of protein-ligand interaction database PLD116 and protein-DNA interaction database PDNA62 complexes, former of which is used as control data sets and the later for additional comparison. These datasets are described in Methods section. Results obtained from this analysis are summarized in Figure 2 and Table 2. It may be observed that certain residues (e.g. TRP, GLN, and ASN) are overrepresented within the binding sites of these 40 proteincarbohydrate complexes, which signifies their importance in protein-carbohydrate interactions. These results may be understood in the light of reported experimental and theoretical studies on carbohydrate interactions. For example, it has been argued that the side chain residues with

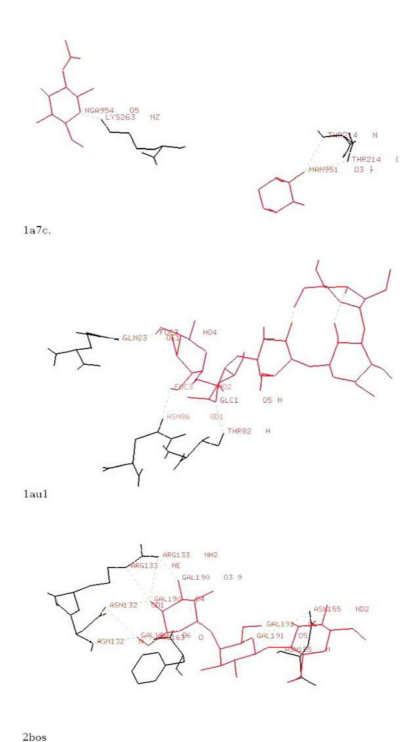


Figure 1 Some typical protein-carbohydrate interactions. All the atoms making hydrogen bonded contacts between sugars and amino acids are labelled.

polar planar groups- ASN, ASP, GLU, GLN, ARG, and HISare the only ones participating in all three forms of hydrogen bonding with sugars and are abundant in sugar-binding sites, which explains why their propensities in the binding sites is higher [14]. Our analyses show that aromatic amino acid residues are often present in carbohydrate-binding sites of proteins. These binding sites are characterized by a placement of a carbohydrate moiety in a stacking orientation to an aromatic ring. This arrangement is an example of CH/pi interactions, which have been shown to play an important role in carbohydrate recognition by glycosidases and carbohydrate-binding proteins [41]. Apart from confirming some of the widely accepted ideas on residue preference for carbohydrate binding, our study determines exact role and contribution of each residue to carbohydrate binding.

Highest propensity score (331% over representation) in the carbohydrate binding sites is observed from Figure 2 and Table 2 for tryptophan (TRP), which is in accordance with many reported mutational studies [42]. This and other studies have provided experimental and theoretical evidence that the presence of TRP residues in mutation sites is crucial for their binding to carbohydrates [43-45].

Additionally, the conservation of aromatic residues, such as tyrosine and phenylalanine, on an exposed surface is common in carbohydrate-binding modules (CBMs) from families 1, 3, 5 and 10, highlighting the role of aromatic residues in carbohydrate binding [46-50]. (It may be noted that CBMs have been previously classified into different families in which groupings of Carbohydrate binding domains or CBDs were called "Types" and numbered with roman numerals (e.g. Type I or Type II CBDs) [51]).

The modification of tryptophan residues has also been shown to cause a compete loss of hemagglutinating activity [52]. Involvement of two tryptophan residues in carbohydrate-binding site was also shown to be essential in the same study. Similarly, Lafora disease-related mutation of TRP32 to glycine (W32G) has also been shown to disrupt the polysaccharide-binding pocket and also potentially unfold the region immediately adjacent to the binding pocket [53]. All these experimental results are well reflected in the high propensity of TRP in carbohydrate binding sites presented in this work (Figure 2).

If the propensity scores of carbohydrate binding are compared with other ligand-binding residues identified by

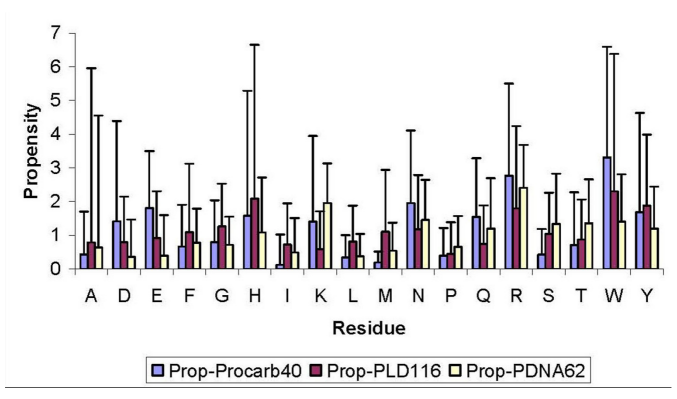


Figure 2
Comparison of binding site propensity of each residue in Procarb40, PDNA62 & PLD116 (residue was marked as binding if any of its atom fell within 3.5 Å of any atom of the ligand/DNA/carbohydrate. Propensity values were obtained by pooling all residues of the same type in all proteins to a single database of binding and non-binding sites. To compute the error bars, propensity values were calculated for each protein separately and standard deviations in propensity values was used as an error bar.

http://www.biomedcentral.com/1472-6807/7/1

Table I: Showing Procarb40 dataset. (Some cells are left empty as no Pfam ID could be found for them).

PDB ID	Pfam ID	Ligand name	Ligand formula	Ligand ID	
<u>IA7C</u>	Serpins	ALPHA-D-MANNOSE	C6 H12 O6	MAN	
		N-ACETYL-D-GALACTOSAMINE	3(C8 HI5 NI O6)	NGA	
		N-METHYLCARBONYLTHREONINE	2(C6 H11 N1 O4)	THC	
<u>IAUI</u>	Interferon	ZINC ION	ZNI 2+	ZN	
		FUCOSE	2(C6 H12 O6)	FUC	
		GLUCOSE	4(C6 H12 O6)	GLC	
		D-GALACTOSE	C6 H12 O6	GAL	
		ALPHA-D-MANNOSE	2(CI HI2 O6)	MAN	
<u>IAXM</u>	Fibroblast Growth Factors	SELENOMETHIONINE	6(C5 H11 N1 O2 SE1)	MSE	
		O2-SULFO-GLUCURONIC ACID	7(C6 HI0 OI0 SI)	IDS	
LCVA.	L et L D	N,O6-DISULFO-GLUCOSAMINE	8(C6 HI3 NI OII S2)	SGN	
<u>ICVN</u>	Lectin legB	CALCIUM ION	4(CAI 2+)	CA	
		MANGANESE (II) ION	4(MNI 2+)	MN	
LEZNI	CDM F 12	ALPHA-D-MANNOSE	12(C1 H12 O6)	MAN	
<u>1E6N</u>	CBM_5_12	GLYCEROL SULFATE ION	C3 H8 O3	GOL SO4	
	Glycol_hydro_18	N-ACETYL-D-GLUCOSAMINE	12(O4 S1 2-) 10(C8 H15 N1 O6)	NAG	
15/2	Tarria D bind C	GLUCOSE	,	GLC	
<u>IFV3</u>	Toxin_R_bind_C Toxin R bind N	PHOSPHATE ION	2(C6 H12 O6) O4 P1 3-	PO4	
	Toxin_N_bind_IN Toxin trans	D-GALACTOSE	4(C6 H12 O6)	GAL	
	TOXIII_ti aris	ETHYL-TRIMETHYL-SILANE	2(C5 H14 SII)	CEQ	
		N-ACETYL-D-GALACTOSAMINE	2(C8 HI5 NI O6)	NGA	
		5-N-ACETYL-BETA-D-NEURAMINIC ACID	2(C11 H19 N1 O9)	SLB	
		5-N-ACETYL-ALPHA-D-NEURAMINIC ACID	4(CII HI9 NI O9)	NAN	
IFWU	Ricin_B_lectin	FUCOSE	C6 H12 O5	FUC	
		O3-SULFONYLGALACTOSE	C6 H12 O9 S1	SGA	
		ALPHA-METHYL-N-ACETYL-D-GLUCOSAMINE	C9 H17 N1 O6	MAG	
<u>IGIT</u>	EGF	FUCOSE	C6 H12 O5	FUC	
	Lectin C	CALCIUM ION	CAI 2+	CA	
		D-GALACTOSE	C6 H12 O6	GAL	
		O-SIALIC ACID	CII HI9 NI O9	SIA	
		N-ACETYL-O-METHYL-D-GLUCOSAMINE	C9 H17 N1 O6	INA	
<u> 1G5N</u>	Annexin	CALCIUM ION	9(CAI 2+)	CA	
		N,O6-DISULFO-GLUCOSAMINE	4(C6 H13 N1 O11 S2)	SGN	
		1,4-DIDEOXY-O2-SULFO-GLUCURONIC ACID	2(C6 H10 O8 S1)	IDU	
		1,4-DIDEOXY-5-DEHYDRO-O2-SULFO-GLUCURONIC ACID	2(C6 H8 O8 S1)	UAP	
<u>IGMN</u>	Kringle	O2-SULFO-GLUCURONIC ACID	3(C6 H10 O10 S1)	IDS	
	PAN	N,O6-DISULFO-GLUCOSAMINE	2(C6 H13 N1 O11 S2)	SGN	
		4-(2-HYDROXYETHYL)-1-PIPERAZINE ETHANESULFONIC ACID	2(C8 H18 N2 O4 S1)	EPE	
<u>IGUI</u>	CBM4/9	CALCIUM ION	CAI 2+	CA	
		GLYCEROL	5(C3 H8 O3)	GOL	
		BETA-D-GLUCOSE	6(C6 H12 O6)	BGC	
<u>IGWM</u>	Family 29 carbohydrate binding module	GLUCOSE	C6 H14 O6	GLC	
		COBALT (II) ION	COI 2+	CO	
		I,2-ETHANEDIOL	8(C2 H6 O2)	EDO	
111/4//	Dan aka t	BETA-D-GLUCOSE	5(C6 HI2 O6)	BGC	
<u>11W6</u>	Bac_rhodopsin	GLUCOSE PETINAL	C6 H12 O6	GLC	
		RETINAL D-GALACTOSE	C20 H28 O1 C6 H12 O6	RET GAL	
		ALPHA-D-MANNOSE	C6 H12 O6 C6 H12 O6	MAN	
		2.3-DI-PHYTANYL-GLYCEROL	C43 H88 O3	L2P	
		2,3-DI-O-PHYTANLY-3-SN-GLYCERO-I-PHOSPHORYL-3'-SN-GLYCEROL-I'-	4(C46 H94 O11 P2 2-)	L3P	

Table I: Showing Procarb40 dataset. (Some cells are left empty as no Pfam ID could be found for them). (Continued)

<u>IJ8R</u>	PapG _N	GLUCOSE	C6 H12 O6	GLC
		D-GALACTOSE	2(C6 H12 O6)	GAL
		N-ACETYL-D-GLUCOSAMINE	C8 H15 N1 O6	NAG
		SELENOMETHIONINE	3(C5 H11 N1 O2 SE1)	MSE
<u>IJPC</u>	B_lectin D-mannose binding lectin	ALPHA-D-MANNOSE	8(C1 H12 O6)	MAN
<u>ILGB</u>	Lectin_legB	FUCOSE	C6 H12 O6	FUC
	Transferrin	CALCIUM ION	CAI 2+	CA
		D-GALACTOSE	C6 H12 O6	GAL
		MANGANESE (II) ION	MNI 2+	MN
		ALPHA-D-MANNOSE	3(CI HI2 O6)	MAN
		N-ACETYL-D-GLUCOSAMINE	4(C8 HI5 NI O6)	NAG
<u>1M5J</u>		ALPHA-D-MANNOSE	8(C6 H12 O6)	MAN
		O1-PENTYL-MANNOSE	C11 H22 O6	OPM
		2- [N-CYCLOHEXYLAMINO]ETHANE SULFONIC ACID	C8 H17 N1 O3 S1	NHE
<u>10H4</u>		CALCIUM ION	CAI 2+	CA
		GLYCEROL	2(C3 H8 O3)	GOL
		SULFATE ION	O4 S1 2-	SO4
		BETA-D-MANNOSE	5(C6 H12 O6)	BMA
		ALPHA D-GALACTOSE	2(C6 H12 O6)	GLA
<u>108V</u>	Lectin_legB	CALCIUM ION	2(CA1 2+)	CA
		MANGANESE (II) ION	2(MNI 2+)	MN
		ALPHA-D-MANNOSE	5(C6 H12 O6)	MAN
		PYROGLUTAMIC ACID	2(C5 H7 N1 O3)	PCA
<u>IQFO</u>	V-set	GLUCOSE	2(C6 H12 O6)	GLC
		D-GALACTOSE	2(C6 H12 O6)	GAL
		O-SIALIC ACID	3(C11 H19 N1 O9)	SIA
<u>IRID</u>	Sushi	O2-SULFO-GLUCURONIC ACID	8(C6 H10 O10 S1)	IDS
		N,O6-DISULFO-GLUCOSAMINE	8(C6 H13 N1 O11 S2)	SGN
<u> 1SE3</u>	Stap_Strp_tox_C	GLUCOSE	C6 H12 O6	GLC
	Stap_Strp_toxin	D-GALACTOSE	C6 H12 O6	GAL
		O-SIALIC ACID	CII HI9 NI O9	SIA
<u> ISL4</u>	Lectin_C	CALCIUM ION	3(CAI 2+)	CA
		ALPHA-D-MANNOSE	4(C6 H12 O6)	MAN
<u>ISLC</u>	Gal-bind_lectin	D-GALACTOSE	4(C6 H12 O6)	GAL
		ALPHA-D-MANNOSE	6(C1 H12 O6)	MAN
		N-ACETYL-D-GLUCOSAMINE	6(C8 H15 N1 O6)	NAG
<u>1T0W</u>	Chitin_bind_I	AMINO GROUP	H2 N1	NH2
		N-ACETYL-D-GLUCOSAMINE	3(C8 HI5 NI O6)	NAG
<u>1T8U</u>	Sulfotransfer_I	SODIUM ION	2(NAT T+)	NA
		SULFATE ION	O4 S1 2-	SO4
		O2-SULFO-GLUCURONIC ACID	C6 H10 O10 S1	IDS
		N,O6-DISULFO-GLUCOSAMINE	2(C6 H13 N1 O11 S2)	SGN
		ADENOSINE-3'-5'-DIPHOSPHATE	2(C10 H15 N5 O10 P2)	A3P
		1,4-DIDEOXY-5-DEHYDRO-O2-SULFO-GLUCURONIC ACID	C6 H8 O8 S1	UAP
<u>IULE</u>		D-GALACTOSE	4(C6 HI2 O6)	GAL
		N-ACETYL-D-GLUCOSAMINE	2(C8 HI5 NI O6)	NAG
<u>IUX7</u>	CBM_6	CALCIUM ION	2(CAI 2+)	CA
		SULFATE ION	O4 S1 2-	SO4
		BETA-D-XYLOPYRANOSE	3(C5 H10 O5)	XYP

Table I: Showing Procarb40 dataset. (Some cells are left empty as no Pfam ID could be found for them). (Continued)

IUY4		SODIUM ION	NAI I+	NA
		CALCIUM ION	CAI 2+	CA
		GLYCEROL	C3 H8 O3	GO
		BETA-D-XYLOPYRANOSE	4(C5 HI0 O5)	XYF
IUYY	CBM 6	CALCIUM ION	4(CAI 2+)	CA
		BETA-D-GLUCOSE	7(C6 H12 O6)	BGC
IVBO		ALPHA-D-MANNOSE	20(C6 H12 O6)	1AM
1.00		N-ACETYLALANINE	8(C5 H9 N1 O3)	AY
IVPS	Polyoma Coat	D-GALACTOSE	5(C6 H12 O6)	GA
1115	1 olyonia Coac	O-SIALIC ACID	10(C11 H19 N1 O9)	SIA
		N-ACETYL-D-GLUCOSAMINE	5(C8 HI5 NI O6)	NA:
IW9T	CBM 6	SODIUM ION	6(NALI+)	NA NA
17771	CBI 1_0	XYLOPYRANOSE	2(C5 H10 O5)	XY
		BETA-D-XYLOPYRANOSE	8(C5 H10 O5)	XY
JXT3	Toxin I	CITRIC ACID	C6 H8 O7	CI
1713	TOXIII_T	N,O6-DISULFO-GLUCOSAMINE	3(C6 H13 N1 O11 S2)	SGI
		1,4-DIDEOXY-O2-SULFO-GLUCURONIC ACID	3(C6 HI0 O8 SI)	IDI
2BOS	SLT beta	BUTYL GROUP	3(C4 H9)	BU
<u> 2603</u>	SLI beta	GLUCOSE	5(C4 H7) 5(C6 H12 O6)	GL
		D-GALACTOSE	14(C6 H12 O6)	GA GA
arcn	DI	GLUCOSE	C6 H12 O6)	GL
2FCP	Plug	PHOSPHATE ION	O4 P1 3-	PO
	TonB_dep_Rec	D-GALACTOSE		GA
		NICKEL (II) ION	2(C6 H12 O6) 2(NI1 2+)	GA N
		3-OXO-BUTYRIC ACID	C4 H6 O3	LIN
		3-OXO-PENTADECANOIC ACID	C15 H28 O3	LIN
		GLUCOSAMINE I-PHOSPHATE	C6 H14 N1 O8 P1	GP
		GLUCOSAMINE 4-PHOSPHATE	C6 H14 N1 O8 P1	GP GP
		ETHANOL AMINE PYROPHOSPHATE	C2 H9 N1 O7 P2	EA
		L-GLYCERO-D-MANNO-HEPTOPYRANOSE	2(C7 H14 O7)	GM
		3-DEOXY-D-MANNO-OCT-2-ULOSONIC ACID	2(C8 H14 O8)	KD
		2-TRIDECANOYLOXY-PENTADECANOIC ACID	2(C28 H54 O4)	LII
2MPR	LamB	GLUCOSE	C ₆ H ₁₂ O ₆	GL
ZITIFIX	Lailib	CALCIUM ION	C ₆ H ₁₂ O ₆ Ca ²⁺	C/
2CUD	Entanatavin h	GLUCOSE		GL
3CHB	Enterotoxin b	D-GALACTOSE	5(C6 H12 O6)	GA
		D-GALACTOSE O-SIALIC ACID	10(C6 H12 O6)	GA SIA
			5(CII HI9 NI O9)	NG.
		N-ACETYL-D-GALACTOSAMINE	5(C8 HI5 NI O6)	
254.651	C 11 1	N-(EHTYLSULFITE)MORPHOLINE	2(C6 HI4 NI O4 SI)	ME
3MAN	Cellulase	ALPHA-D-MANNOSE	3(C6 H12 O6)	MAI
3MBP		GLUCOSE	3(C6 H12 O6)	GLO

Table 2: Propensities of Procarb40, PDNA62 & PLD116 along with their binding and non-binding data

	PI	ROCARB4	0		PDNA62			PLD116	
Residue	Propensity	BS	NBS	Propensity	BS	NBS	Propensity	BS	NBS
Α	0.43	9	494	0.64	42	389	0.79	109	2684
С	0.00	0	29	0.34	7	143	1.07	24	436
D	1.41	27	433	0.36	18	292	0.79	84	2009
E	1.81	29	356	0.39	32	510	0.92	92	1952
F	0.66	9	318	0.77	33	245	1.09	70	1346
G	0.80	20	581	0.71	46	372	1.26	176	2633
Н	1.58	8	114	1.08	39	194	2.09	81	712
ı	0.12	2	392	0.48	30	373	0.72	70	1837
K	1.40	26	419	1.95	180	423	0.59	65	2053
L	0.34	8	561	0.38	39	624	18.0	120	2872
M	0.19	I	124	0.54	14	149	1.11	42	716
N	1.96	38	429	1.45	74	260	1.17	92	1485
P	0.40	5	297	0.66	35	307	0.45	38	1597
Q	1.54	18	263	1.19	61	272	0.74	46	1123
R	2.77	32	246	2.41	208	360	1.80	139	1450
S	0.43	9	499	1.33	91	355	1.03	112	2049
Т	0.70	15	499	1.36	85	325	0.87	90	2030
V	0.00	0	472	0.59	40	399	0.73	92	2315
W	3.31	23	144	1.40	22	81	2.30	67	518
Υ	1.68	25	333	1.19	43	189	1.88	125	1189

PLD116 database (see Methods), TRP remains the most prominent high propensity residue. However, high propensity score of HIS residues is also shown by PLD binding sites, indicating that the role of HIS in ligand-binding sites does not have specific preference for carbohydrates, but HIS in general being an active site shows high propensity of binding to any ligand, including carbohydrates. Next important residue is ARG whose propensity for carbohydrate binding is less than TRP within Procarb40, yet the propensity (277% over representation) is even higher than what is observed for ARG in DNA-binding proteins database PDNA62 (= 241% over representation) (see Table 2 and Methods section). These results are supported by some published results of transmutagenesis experiments reporting crucial role of ARG residues in some protein-carbohydrate interactions [54]. Lower propensity scores for the other basic residue LYS indicate that the interaction between ARG and sugar is not purely electrostatic in nature. Dahms et al. in 1993, [54] also report that the substitution of ARG residues by LYS in Insulin-like growth factor also caused loss of binding despite similar electropositive property of these residues and also despite overall conservation of structure upon this mutation. These results were interpreted that the proteins utilize residues with planar side chains (ARG, ASN, ASP, GLU) for their interaction with sugars. Higher propensity scores of ASP and GLU, which are also negatively charged residues, also support this argument. These propensity scores are higher than what is observed for other ligands (PLD116 database), thus highlighting a preference of these residues to interact carbohydrates in contrast to other types of ligands. In comparison to DNA-binding propensity scores of ASP and GLU are much higher, obviously because negatively charged bases in DNA repel negative charged residues.

Solvent accessibility of binding sites compared with the rest of the protein

We next attempted to establish a residue-wise relationship between solvent accessibility and carbohydrate binding. Figure 3 shows the mean solvent accessibility (ASA) values for the binding and non-binding regions in Procarb40 database. We observe that the most frequent carbohydrate binder TRP has a significantly higher ASA in binding locations compared with non-binding ones. Similar higher ASA for binding regions are also observed for other aliphatic residues ALA, GLY, ILE and LEU. Thus, the hydrophobic residues, which are usually in the buried states, do not apparently participate in sugar binding. In order to bind sugars they are expected to be on the surface, thus facilitating their hydrophobic interactions with carbohydrate atoms of protein-carbohydrate complexes and reveals that polar uncharged and certain hydrophobic residues (e.g. TYR, TRP, ALA, LEU and ILE) seem to have higher mean ASA-values in the binding regions. This result contrasts with similar binding sites analysis on DNA-binding proteins, where ASA of charged residues showed a better discrimination between binding and non-binding regions [28]. Most charged and polar residues do not show any difference in their ASA for binding and non-binding regions, presumably because their probability to be on the surface is higher irrespective of their role in binding. For a quick comparison of role of ASA in binding regions of PLD116 and PDNA62 databases with Procarb40, ratio of mean ASA in binding to non-binding regions of the three databases have been plotted in Figure 4 (see Additional file 1). As discussed above, aliphatic residues ILE, LEU and GLY show the highest ratio for Procarb40, in addition to the most frequent binder TRP. Very low values of CYS and VAL residues are not significant as there are very few binding residue of this type (see Table 2).

Role of Secondary structure

We tried to explore if certain residues prefer any secondary structure for binding to carbohydrates. Results of these statistics are p

resented in Additional File 1 as Tables 5-10 and Figures 5a-g. If the number of binding sites is resolved into their secondary structure types, very few binding sites are assigned to each category. This leaves the resulting data to be insufficient for any statistical conclusions. These results are therefore not discussed here, but only provided in Additional File 1 for reference.

Packing Density

We also tried to find out the difference between the packing density of the binding and non-binding residues and observed that there is no statistically significant difference of packing density between binding and non-binding residues.

Prediction results

Looking at clear preferences of residues for binding carbohydrates (Figure 1), we sought to develop a prediction method, which could take the predisposition of residues and their sequence environments as an input and thereby identify binding residues from the information of proteinsequence. To do so, sequence environment at each residue level could be represented either as binary 20 bit vectors or by the rows of the matrices depicting evolutionary profiles of residues at each location. Sequence neighbor environment could be added as the corresponding rows of this matrix (called position-specific substitution matrix or PSSM) on either side. Schemes of these representations have been extensively developed for the problem of solvent accessibility and other residue-wise features of proteins [55]. Table 3 summarized the results of predictions obtained in this way, using a leave-one-out method. This method also allows us to compute the standard errors in the prediction scores. Further, prediction performance of sequence-only predictors has been compared with those

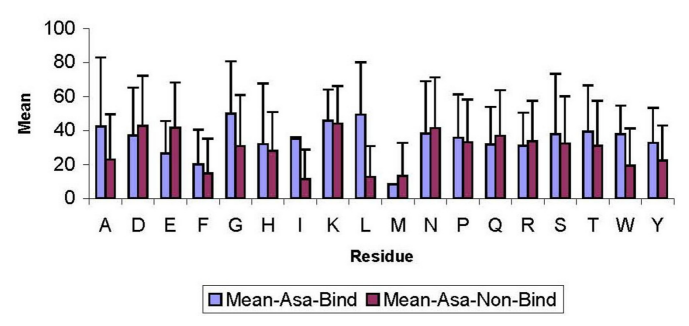


Figure 3

Comparison between mean ASA values of residues in binding and non-binding sites for Procarb40. Error bars are taken from their standard deviation in each protein. The graph does not contain cystein and valine data as none of these residues were found to be in the binding regions.

using PSSMs. The best performance for Procarb40 data set was found to be a modest 61%, indicating that the sequence and evolutionary information do not decisively determine a binding site. This not-so-good prediction performance for Procarb40 is is apparently because carbohydrates are diverse and finding overall general rules for their binding sites in proteins may not be possible with the amount of data we have. We need to have large data with sufficient representation of all types of sugars. To ensure that the low performance is caused by the diversity of sugars, we tried to develop a prediction model for only one type of sugar. We tried many differently classified carbohydrates, but due to further small size of data, could only use galactose binding proteins (GalBind18) data set used by Sujatha et al. (2004) [57] to have a sufficient number of binding sites to model. As expected prediction performance for proteins binding to only one type of sugars, was very much higher than all carbohydrates taken together. Table 3 shows that in GalBind18 carbohydrate binding sites could be predicted with as much as 79% specificity and 63% sensitivity. We speculate that much better prediction methods will be developed when a large number of proteins binding to each type of carbohydrates become available.

Single sequence versus evolutionary information

It may be a little surprising to note that PSSM based predictions (55%) were somewhat poorer than single sequences (61%) in Procarb40. However in the case of GalBind18 the situation in reversed. Lower values of prediction in PSSM based methods could be due to two reasons. First of all the number of sequences which gave significant alignments with Procarb40 was roughly 400, which is small and hence the evolutionary information transferred to PSSM may not be enough to improve performance. Secondly, the diversity of Procarb40 may lead to higher conservation scores to some residues and hence there would be many false positive predictions by this (that is why the specificity of PSSM based method was as low as 23%). In the case of GalBind18, the situation is reversed because the carbohydrates are more similar and

hence conservation of a residue within them does convey positive information about its binding behaviour. Thus PSSMs do not carry false information to the neural network.

Comparison with other studies

Although some of the results presented in this work may be obvious to some experienced biologists, yet this work is the first attempt to summarize the sequence and structure features of carbohydrate binding proteins in such a comprehensive way. Previous studies have either focused on a small set of proteins aiming to analyze one or a few types of residues [43-45], or tried to focus either on the structural aspect [e.g. [16,17,26]] or just the sequence aspect of these interactions. This is also the first attempt to use sequence and evolutionary information to predict carbohydrate-binding sites using neural network based approach, which has been proved to be successful in making other sequence-based predictions. Earlier, structurebased methods have been employed to develop empirical rules on patches and other structure descriptors with a somewhat better (65%) accuracy. However, sequencebased methods, employing only sequence information presented in this work are new and will have a much wider application as no structure information will be required for prediction. We expect that this will trigger interest in the prediction of carbohydrate binding sites using machine learning methods and the performance will improve with the availability of more data.

Conclusion

This analysis of protein-carbohydrate interactions in terms of proteins sequence and solvent accessibility reveals that TRP and ARG residues have the highest overall binding propensity for all types of carbohydrates. Planar side chains of polar residues are also confirmed to have overall high propensity of binding. Mean solvent accessibility of hydrophobic residues has been found to be higher for binding regions, whereas charged residues have almost the same solvent accessibility in binding and non-binding regions. A neural network, trained to use evolu-

Table 3: Comparison of Binary and PSSM prediction results using jackknife leave-one-out method (binding sites were labeled at 3.5 Å cut-off distance between carbohydrate and protein atoms).

Data type	Validation type	Average-sensitivity	Average-specificity	Average -net Prediction	P-value	
GalBind18	Leave I out (Using PSSM)	0.63 (0.19)	0.79 (0.09)	0.71 (0.09)	0.08859	
GalBind18	Leave I out (Using single sequences)	0.62 (0.26)	0.68 (0.12)	0.65 (0.11)		
Procarb40	Leave I out (Using PSSM)	0.87 (0.12)	0.23 (0.08)	0.55 (0.06)	0.00209	
Procarb40	Leave I out (Using single sequences)	0.68 (0.22)	0.55 (0.16)	0.61 (0.12)		

Due to a large number of iterations required in a leave-one-out method, the prediction performance has a significant standard deviation, which has been shown in brackets. P-values are for two-tailed t-test conducted to distinguish between the predictions performances of single sequences versus evolutionary information coded by PSSM. In Procarb40, evolutionary profiles give a significantly poorer result than single sequences, due to a high false positive rate (low specificity).

tionary information of residues and their neighbors could correctly make prediction of binding or non-binding residues with 69–72% specificity and 55–57% sensitivity.

Methods

Definition of a binding site

A binding residue is defined as any amino acid in the protein such that any of its atoms is within a cut-off distance from any atom from the sugar in the protein-carbohydrate complex. We tried to determine the best cut-off distance and found that 3.5 Å distance could best separate the binding residues from non-binding ones in the propensity graphs and also gives the best accuracy figures in neural network based predictors. Thus, all the reported results are based on this distance cut-off unless otherwise stated.

Data Sets

Procarb40

PDB search was performed for protein-carbohydrate complexes with a pair-wise similarity of 50% or less. Only one structure was taken in case there were more than one representative from the same family. For polypeptides, only one chain was selected on the basis of maximum number of binding sites present. FASTA formatted sequences were subsequently formatted using *formatdb* program of the BLAST package. BLASTCLUST program [56] at 30% threshold refined our search to 40 structures (Table 1). We call this database Procarb40.

GalBind I 8

This is a data set of 18 Galactose specific proteins selected for another analysis by Sujatha *et al.* [57].

PDNA62

This is the (non redundant) data set of 62 DNA-binding proteins [28].

PLD116

This is a non-redundant data set of ligand-binding proteins developed for the current study. To begin with, all the 485 protein-ligand complexes were downloaded from Protein-Ligand Database [58] (v1.3 as on 25/01/06). Redundancy among sequences was first removed by using CD-HIT program from [59] with a threshold of 40% sequence identity. This resulted in 178 clusters. FASTA formatted sequences were subsequently formatted using *formatdb* program of the BLAST package. The redundancy was further removed with a threshold of 30% sequence identity using BLASTCLUST program [56]. A data set was thus created, by retaining only the representative ones such that no two sequences in the resulting data set have more than 30% sequence identity. We call this database PLD116.

Other data sets

PDB-ALL (47,189 sequences) is a data set of all protein sequences obtained from NCBI. PIR is the sequence data set (283,177 sequences) of Protein Information Resource at Georgetown University [60]. SWISSPROT is another well-known database of sequences [61]. NCBI-NR is a non-redundant data set of all protein sequences compiled from GeneBank, PIR, SwissProt, PDB and other resources by NCBI [62] were also used in the current work.

Generation of PSSMs

Target sequences are scanned against the reference data sets to compile a set of alignment profiles or position specific scoring matrices (PSSMs) using Position Specific Iterative BLAST (PSI BLAST) program [63]. Three cycles of PSI-BLAST were run for each protein and the scores were saved as profile matrices (PSSMs). NR database of NCBI, PDBAA (database of all amino acid sequences of proteins in PDB), SWISSPROT and PIR were used for building the profiles. Profiles from NR database of NCBI were used for most of the calculations presented in this work unless otherwise specified.

Calculation of amino acid composition, solvent accessibility and secondary structure at binding sites

We collected statistics on amino acid residues, which were involved in carbohydrate binding. An attempt was then made to determine whether there was a preference for any particular amino acid residue. Frequency of occurrence for each residue type is calculated and corresponds to the relative number of residues of that type out of all the residues that were found in the carbohydrate-binding proteins.

Solvent accessibility or accessible surface area (ASA) values of Procarb40, PDNA62 and PLD116 complexes were obtained from our earlier database of (relative) solvent accessibility of proteins ASAVIEW [64], whereas the secondary structure was obtained using DSSP program [65].

Propensity scores

Propensity of a residue in the binding site was calculated by the formula: -

$$\frac{NB_i}{N_i} \frac{NB_{all}}{N_{all}}$$

where NB_i is the number of residues of type i, which bind to carbohydrate, N_i is the total number of residues of type i, NB_{all} is the total number of all binding residues, N_{all} is the total number of all residues. To compute the propensity score of each residue, the data of binding and non-binding residues were pooled together and a single propensity score was obtained for the entire data of proteins.

Also, propensity scores for each protein were calculated separately and standard deviation in all propensity scores for the same residue type was used as the error bar.

Neural network

Neural network inputs

Conservation scores in 20 amino acid positions for every residues form 20 columns (column 3 onwards) of corresponding row in a PSI-BLAST PSSM. For every residue, we make a binary (1 for binding and 0 for non-binding) prediction of that residue being a binding site or not. Input for every prediction is the PSSM score on the row corresponding to this target residue and one more rows on either side ($20 \times 3 = 60$ inputs) as well as two more rows on either side ($20 \times 5 = 100$ inputs).

Network architecture and transfer function

A fully connected, feed-forward neural network was constructed using Stuttgart Neural Network Simulator (SNNS) version 4.2, developed at University of Stuttgart [66]. After varying the number of units, and hidden layers, it was found that a network with two units in the hidden layer and a single output unit performed slightly better than other choices.

Training and validation

Different datasets and their cross validation were tried. Out of these results are presented for which prediction performance was better than others. We use a leave-one-out approach for training and validation. In this approach, data corresponding to one protein is removed from the data set and the remaining proteins are trained using a neural network. The performance on the left out protein is than measured. The process is systematically repeated for all proteins, leaving them out one by one and measuring their prediction accuracy. Finally reported accuracy scores correspond to the averages of the left out proteins.

Most other procedures for training and assessment of prediction accuracy were the same as in our earlier work [67].

Assessment of prediction performance

Three scores were used for the measure of prediction performance viz. Sensitivity (S1), Specificity (S2) and their average Net Prediction (NP). They are defined as follows:

Sensitivity (S1) = TP/(TP+FN)

Specificity (S2) = TN/(TN+FP)

Where *TP* stands for correctly identified binding sites, *TN* stands for correctly identified non-binding residues, *FP* stands for number of non-binding residues wrongly iden-

tified as binding by predictor, and *FN* is the number of binding residues predicted as non-binding.

Authors' contributions

This work is part of the doctoral research of AM. SA conceived the problem and designed the protocols. All calculations were performed by AM under SA's guidance and supervision.

Additional material

Additional file 1

Supplementary Material. The data provides tables and figures with additional information on topics presented in the main text. Some of these results may not be statistically significant due to small size of data. Click here for file

[http://www.biomedcentral.com/content/supplementary/1472-6807-7-1-\$1.doc]

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References

- Shionyu-Mitsuyama C, Shirai T, Ishida H, Yamane T: An empirical approach for structure-based prediction of carbohydrate binding sites on proteins. Protein Eng 2003, 16:467-78.
- 2. Scripps [http://www.scripps.edu/news/press/120704.html]
- Scripps [http://www.Scripps.edu/newsandviews/e 20011008/paulson1.html]
- Sharon N, Lis H: Lectins as cell recognition molecules. Science 1989, 246:227-234.
- Lasky LA: Selectins: interpreters of cell-specific carbohydrate information during inflammation. Science 1992, 258:964-969.
- Sastry K, Ezekowitz RA: Collectins: pattern recognition molecules involved in first line host defense. Curr Opin Immunol 1993, 5:59-66.
- Barondes SH, Cooper DN, Gitt MA, Leffler H: Galectins: Structure and function of a large family of animal lectins. J Biol Chem 1994, 269:20807-10.
- Hoppe HJ, Reid KB: Collectins soluble proteins containing collagenous regions and lectin domains - and their roles in innate immunity. Protein Sci 1994, 3:1143-58.
- Rosen SD, Bertozzi CR: The selectins and their ligands. Curr Opin Cell Biol 1994, 6:663-73.
- Sharon N, Lis H: Lectins-proteins with a sweet tooth: function in cell recognition. Essays Biochem 1995, 30:59-75.
- Sharon NH, Lis H: Lectins: Carbohydrate-Specific Proteins That Mediate Cellular Recognition. Chem Rev 1998, 98:637-674.
- Karlsson KA: Meaning and therapeutic potential of microbial recognition of host glycoconjugates. Mol Microbio 1998, 29:1-11.
- Audette GF, Delbaere LT, Xiang J: Mapping protein: carbohydrate interactions. Curr Protein Pept Sci 2003, 4:11-20.
- Quiocho FA: Protein-carbohydrate interactions: basic molecular features. Pure & Appl Chem 1989, 61:1293-1306.
- Vyas NK: Atomic features of protein-carbohydrate interactions. Curr Opin Struct Biol 1991, 1:732-740.
- Spurlino JC, Rodseth LE, Quiocho FA: Atomic interactions in protein-carbohydrate complexes. Tryptophan residues in the periplasmic maltodextrin receptor for active transport and chemotaxis. J Mol Biol 1992, 226:15-22.

- 17. Toone El: Structure and energetics of protein-carbohydrate complexes. Curr Opin Struct Biol 1994, 4:719-728.
- Meyer JE, Schulz GE: Energy profile of maltooligosaccharide permeation through maltoporin as derived from the structure and from a statistical analysis of saccharide-protein interactions. Protein Sci 1997, 6:1084-91.
- Rini JM: Lectin structure. Annu Rev Biophys Biomol Struct 1995, **24:**51-77
- 20. Weis WI, Drickamer K: Structural basis of lectin-carbohydrate recognition. Annu Rev Biochem 1996, 65:441-73.
- 21. Elgavish S, Shaanan B: Lectin-carbohydrate interactions: different folds, common recognition principles. Trends Biochem Sci 1997, **22:**462-7.
- Rao VS, Lam K, Qasba PK: Architecture of the sugar binding sites in carbohydrate binding proteins – a computer modeling study. Int J Biol Macromol 1998, 23:295-307.
- 23. Garcia-Hernandez E, Hernandez-Arana A: Structural bases of lectin-carbohydrate affinities: comparison with protein-folding energetics. Protein Sci 1999, 8:1075-86.
- 24. Garcia-Hernandez E, Zubillaga RA, Rodriguez-Romero A, Hernandez-Arana A: Stereochemical metrics of lectin-carbohydrate interactions: comparison with protein-protein interfaces. Glycobiology 2000, 10:993-1000.
- Clarke C, Woods RJ, Gluska J, Cooper A, Nutley MA, Boons GJ: Involvement of water in carbohydrate-protein binding. J Am Chem Soc 2001, 123:12238-47.
- 26. Neumann D, Kohlbacher O, Lenhof HP, Lehr CM: Lectin-sugar interaction. Calculated versus experimental binding energies. Eur J Biochem 2002, 269:1518-24.
 Taroni C, Jones S, Thornton JM (2): Analysis and prediction of
- carbohydrate binding sites. Protein Eng 2000, 13:89-98.
- Ahmad S, Gromiha MM, Sarai A: Analysis and prediction of DNAbinding proteins and their binding residues based on composition, sequence and structural information. Bioinformatics 2004, **20:**477-86.
- Ahmad S, Sarai A: PSSM-based prediction of DNA binding sites in proteins. BMC Bioinformatics 2005, 6:33.
- Wang L, Brown SJ: BindN: a web-based tool for efficient prediction of DNA and RNA binding sites in amino acid sequences. Nucleic Acids Res 2006, 34:W243-248.
- Kuznetsov IB, Gou Z, Li R, Hwang S: Using evolutionary and structural information to predict DNA-binding sites on DNA-binding proteins. Proteins 2006, 64:19-27.
- Yan C, Terribilini M, Wu F, Jernigan RL, Dobbs D, Honavar V: Predicting DNA-binding sites of proteins from amino acid sequence. BMC Bioinformatics 2006, 7:262.
- 33. Fariselli P, Pazos F, Valencia A, Casadio R: Prediction of proteinprotein interaction sites in heterocomplexes with neural networks. Eur J Biochem 2002, 269:1356-1361.
- Jones S, Thornton JM: Prediction of protein-protein interaction sites using patch analysis. | Mol Biol 1997, 272:133-143
- Lu L, Lu H, Skolnick J: MULTIPROSPECTOR: an algorithm for the prediction of protein-protein interactions by multimeric threading. Proteins 2002, 49:350-64.
- Keskin O, Ma B, Rogale K, Gunasekaran K, Nussinov R: Proteinprotein interactions: organization, cooperativity and mapping in a bottom-up Systems Biology approach. Phys Biol 2005, 2:S24-S35
- 37. Raih MF, Ahmad S, Zheng R, Mohamed R: Solvent accessibility in native and isolated domain environments: general features and implications to interface predictability. Biophys Chem 2005, 114:63-9
- 38. Hoskins J, Lovell S, Blundell TL: An algorithm for predicting protein-protein interaction sites: Abnormally exposed amino acid residues and secondary structure elements. Protein Sci 2006, 15:1017-29.
- 39. Julenius K, Molgaard A, Gupta R, Brunak S: Prediction, conservation analysis, and structural characterization of mammalian mucin-type O-glycosylation sites. Glycobiology 2005, 15:153-64.
- Blom N, Gammeltoft S, Brunak S: Sequence and structure-based prediction of eukaryotic protein phosphorylation sites. J Mol Biol 1999, 94:1351-62
- Spiwok V, Lipovova P, Skalova T, Vondrackova E, Dohnalek J, Hasek J, Kralova B: Modelling of carbohydrate-aromatic interactions: ab initio energetics and force field performance. J Comput Aided Mol Des 2005, 19:887-901.

- 42. McLean BW, Bray MR, Boraston AB, Gilkes NR, Haynes CA, Kilburn DG: Analysis of binding of the family 2a carbohydrate-binding module from Cellulomonas fimi xylanase IOA to cellulose: specificity and identification of functionally important amino acid residues. Protein Eng 2000, 13:801-9.
- Poole DM, Hazlewood GP, Huskisson NS, Virden R, Gilbert HJ: The role of conserved tryptophan residues in the interaction of a bacterial cellulose-binding domain with its ligand. FEMS Microbiol Lett 1993, 106:77-83.
- Din N, Forsythe IJ, Burtnick LD, Gilkes NR, Miller RC Jr, Warren RA, Kilburn DG: The cellulose-binding domain of endoglucanase A (CenA) from Cellulomonas fimi: evidence for the involvement of tryptophan residues in binding. Mol Microbiol 1994, 11:747-55.
- Bray MR, Johnson PE, Gilkes NR, McIntosh LP, Kilburn DG, Warren RA: Probing the role of tryptophan residues in a cellulosebinding domain by chemical modification. Protein Sci 1996,
- Reinikainen T, Ruohonen L, Nevanen T, Laaksonen L, Kraulis P, Jones TA, Knowles JK, Teeri TT: Investigation of the function of mutated cellulose-binding domains of Trichoderma reesei cellobiohydrolase I. Proteins 1992, 14:475-82.
- Nagy T, Simpson P, Williamson MP, Hazlewood GP, Gilbert HJ, Orosz L: All three surface tryptophans in Type IIa cellulose binding domains play a pivotal role in binding both soluble and insoluble ligands. FEBS Lett 1998, 429:312-6.
- Simpson HD, Barras F: Functional analysis of the carbohydratebinding domains of Erwinia chrysanthemi Cel5 (Endoglucanase Z) and an Escherichia coli putative chitinase. J Bacteriol 1999, 18:4611-6.
- Ponyi T, Szabo L, Nagy T, Orosz L, Simpson PJ, Williamson MP, Gilbert HJ: Trp22, Trp24, and Tyr8 play a pivotal role in the binding of the family 10 cellulose-binding module from Pseudomonas xylanase A to insoluble ligands. Biochemistry 2000, 39:985-91.
- Raghothama S, Simpson PJ, Szabo L, Nagy T, Gilbert HJ, Williamson MP: Solution structure of the CBM10 cellulose binding module from Pseudomonas xylanase A. Biochemistry 2000, 39:978-84
- CBM [http://afmb.cnrs-mrs.fr/CAZY/fam/acc CBM.html]
- Gao S, An J, Wu CF, Gu Y, Chen F, Yu Y, Wu QQ, Bao JK: Effect of amino acid residue and oligosaccharide chain chemical modifications on spectral and hemagglutinating activity of Millettia dielsiana Harms. ex Diels. lectin. Acta Biochim Biophys Sin (Shanghai) 2005, 37:47-54.
- Wang J, Stuckey JA, Wishart MJ, Dixon JE: A unique carbohydratebinding domain targets the lafora disease phosphatase to glycogen. J Biol Chem 2002, 277:2377-80.
- Dahms NM, Rose PA, Molkentin JD, Zhang Y, Brzycki MA: The bovine mannose 6-phosphate/insulin-like growth factor II receptor. The role of arginine residues in mannose 6-phosphate binding. J Biol Chem 1993, 268:5457-63. Arauzo-Bravo M, Ahmad S, Sarai A: " Dimensionality of amino
- acid space and solvent accessibility prediction with neural networks". Comput Bio Chem 2006, 30:160-168.
- Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ: Basic local alignment search tool. J Mol Biol 1990, 215:403-410.
- Sujatha MS, Balaji PV: Identification of common structural features of binding sites in galactose-specific proteins. Proteins 2004, 5:44-65
- Puvanendrampillai D, Mitchell JB: L/D Protein Ligand Database (PLD): additional understanding of the nature and specificity of protein-ligand complexes. Bioinformatics 2003, 19:1856-7.
- Li W, Jaroszewski L, Godzik A: Clustering of highly homologous sequences to reduce the size of large protein database. Bioinformatics 2001, 17:282-283.
- Apweiler R, Bairoch A, Wu CH: Protein sequence databases. Curr Opin in Chem Biol 2004, 8:76-80.
- 61. Bairoch A, Boeckmann B: The SWISS-PROT protein sequence data bank. Nucleic Acids Res 1991, 19:2247-2249.
- NCBI BLAST database download site ftp.ncbi.nlm.nih.gov/blast/db/]
- Altschul SF, Madden TL, Schäffer AA, Zhang J, Zhang Z, Miller W, Lipman DJ: Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res 1997, **25:**3389-3402.

- Ahmad S, Gromiha M, Fawareh H, Sarai A: ASAView: database and tool for solvent accessibility representation in proteins. BMC Bioinformatics 2004, 5:51.
- Kabsch W, Sander C: Dictionary of protein secondary structure: pattern recognition of hydrogen-bonded and geometrical features. Biopolymers 1983, 22:2577-637.
- 66. SNNS [http://www-ra.informatik.uni-tuebingen.de/SNNS/]
- Ahmad S, Gromiha MM: Design and training of a neural network for predicting the solvent accessibility of proteins. J Comput Chem 2003, 24:1313-1320.

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