## **Description of features used for training**

## Glossary:

kDa - kilodaltons

 $Å^2$  - Angstrom squared

Å<sup>3</sup> - Angstrom cubed

DSSP – Dictionary of Secondary Structure of Proteins

 $O(I)>H-N(I\pm n)$  H bonds' type descriptors explained:

Alpha helices are part of a protein's secondary structure. They form when the backbone of the amino acid chain arranges into a helical structure with hydrogen bonds stabilising it.

The H-bond happens between the carbonyl oxygen from amino acid i and the N-H group of amino acid  $i\pm n$ . These bonds can happen between amino acids up to 5 positions away from each other. So 'O(I)>H-N(I+2)\_H\_bonds' for example represents the number of H-bonds that happen between the i-th amino acid oxygen and the i+2 amino acid N-H group.

## Notes on some known mistakes and issues:

- 1. The '%polar' descriptor in the training data is labelled wrong, it should be named '%apolar', and it contains the percentage of apolar, not polar, amino acids in the protein.
- 2. If a description is marked as '-', it is because a description was not found for all the molecular descriptors calculated by Mordred and external information could not be found.

<b>Feature name</b>	Description
prot_MW	Protein molecular weight (from sequence). In
	kDa
tot_aa_SASA	Total amino acid solvent accessible surface area
	(SASA), calculated by adding up individual
	amino acid SASA. In Å <sup>2</sup>
tot_VWvol	Total Van der Waals volume (VWV), calculated
	by adding up amino acid VWV. In Å <sup>3</sup>
%A	Percentage of alanine in the protein
%R	Percentage of arginine in the protein
%N	Percentage of aspargine in the protein

Feature name	Description
%D	Percentage of aspartic acid in the protein
%C	Percentage of cysteine in the protein
%E	Percentage of glutamic acid in the protein
<b>%</b> Q	Percentage of glutamine in the protein
<b>%</b> G	Percentage of glycine in the protein
%Н	Percentage of histidine in the protein
%I	Percentage of isoleucine in the protein
%L	Percentage of leucine in the protein
%K	Percentage of lysine in the protein
%M	Percentage of methionine in the protein
%F	Percentage of phenylalanine in the protein
%P	Percentage of proline in the protein
%S	Percentage of serine in the protein
%T	Percentage of threonine in the protein
%W	Percentage of tryptophan in the protein
%Y	Percentage of tyrosine in the protein
%V	Percentage of valine in the protein
%polar	Percentage of apolar amino acids in the protein
%pol_unch	Percentage of polar uncharged amino acids in the
	protein
%pol_ch	Percentage of polar charged amino acids in the
	protein
%pol_ch_basic	Percentage of polar charged basic amino acids in
	the protein
%pol_ch_acid	Percentage of polar charged acidic amino acids in
	the protein
%aromatic	Percentage of aromatic amino acids in the protein
%aliphatic	Percentage of aliphatic amino acids in the protein
%hydrox	Percentage of hydroxylic amino acids in the
	protein
%amidic	Percentage of amidic amino acids in the protein
%w_S	Percentage of amino acids with a Sulfur atom in
	the protein
tot_aa_TPSA	Total topological polar surface area (TPSA) of all

Feature name	Description
	amino acids in the protein. In Å <sup>2</sup>
apolar_SA	Apolar surface area of amino acids in the protein.
	In Å <sup>2</sup>
pol_unch_SA	Polar uncharged surface area of amino acids in
	the protein. In Å <sup>2</sup>
pol_ch_SA	Polar charged surface area of amino acids in the
	protein. In Å <sup>2</sup>
pol_ch_basic_SA	Polar charged basic surface area of amino acids in
	the protein. In Å <sup>2</sup>
pol_ch_acid_SA	Polar charged acidic surface area of amino acids
	in the protein. In Å <sup>2</sup>
tot_SASA/actual_asa	SASA calculated from amino acids. In Å <sup>2</sup>
	(tot_aa_SASA) divided by the actual SASA of the
	protein (from DSSP). In Å <sup>2</sup>
length	Length of the protein sequence (all chains added)
n_chains	Number of individual chains in the protein
aa/chain	Average amino acids per chain
tot_SS_bri	Total S-S bridges
intrachain_SS_bri	Intrachain S-S bridges
interchain_SS_bri	Interchain S-S bridges
asa	*From here onwards, until stated otherwise, all
	data is from DSSP.*
	Accessible surface area of the protein . In $\text{ Å}^2$
tot_O(I)>H-N(J)_H_bonds	Total H bonds of type $O(I)>H-N(J)$
O(I)>H-N(J)_H_bonds%	Percentage of H bonds of type O(I)>H-N(J)
parallel_bri_H_bonds	Total H bonds in parallel bridges
parallel_bri_H_bonds%	Percentage of H bonds in parallel bridges
antiparallel_bri_H_bonds	Total H bonds in antiparallel bridges
antiparallel_bri_H_bonds%	Percentage of H bonds in antiparallel bridges
O(I)>H-N(I-5)_H_bonds	Total H bonds of type O(I)>H-N(I-5)
O(I)>H-N(I-5)_H_bonds%	Percentage of H bonds of type O(I)>H-N(I-5)
O(I)>H-N(I-4)_H_bonds	Total H bonds of type O(I)>H-N(I-4)
O(I)>H-N(I-4)_H_bonds%	Percentage of H bonds of type O(I)>H-N(I-4)
O(I)>H-N(I-3)_H_bonds	Total H bonds of type $O(I)>H-N(I-3)$

Feature name	Description
O(I)>H-N(I-3)_H_bonds%	Percentage of H bonds of type O(I)>H-N(I-3)
$O(I)>H-N(I-2)_H_bonds$	Total H bonds of type O(I)>H-N(I-2)
$O(I)>H-N(I-2)_H_bonds\%$	Percentage of H bonds of type O(I)>H-N(I-2)
$O(I)>H-N(I-1)_H_bonds$	Total H bonds of type O(I)>H-N(I-1)
$O(I)>H-N(I-1)_H_bonds\%$	Percentage of H bonds of type O(I)>H-N(I-1)
$O(I)>H-N(I+2)_H_bonds$	Total H bonds of type O(I)>H-N(I+2)
$O(I)>H-N(I+2)_H_bonds\%$	Percentage of H bonds of type O(I)>H-N(I+2)
$O(I)>H-N(I+3)_H_bonds$	Total H bonds of type O(I)>H-N(I+3)
$O(I)>H-N(I+3)_H_bonds\%$	Percentage of H bonds of type O(I)>H-N(I+3)
$O(I)>H-N(I+4)_H_bonds$	Total H bonds of type O(I)>H-N(I+4)
$O(I)>H-N(I+4)_H_bonds\%$	Percentage of H bonds of type O(I)>H-N(I+4)
$O(I)>H-N(I+5)_H_bonds$	Total H bonds of type O(I)>H-N(I+5)
$O(I)>H-N(I+5)_H_bonds\%$	Percentage of H bonds of type O(I)>H-N(I+5)
	End of DSSP data.
nAcid	*From here onwards, until stated otherwise, the
	descriptors pertain to the ligand.*
	Number of acidic groups in the ligand
nBase	Number of basic groups
nAromAtom	Number of atoms in an aromatic structure
nAromBond	Number of bonds in an aromatic structure
nAtom	Total number of atoms in the ligand
nHeavyAtom	Number of heavy atoms in the ligand
nSpiro	-
nBridgehead	-
nHetero	-
nH	Number of hydrogen atoms in the ligand
nB	Number of boron atoms in the ligand
nC	Number of carbon atoms in the ligand
nN	Number of nitrogen atoms in the ligand
nO	Number of oxygen atoms in the ligand
nS	Number of sulphur atoms in the ligand
nP	Number of phosphorus atoms in the ligand
nF	Number of fluorine atoms in the ligand
nCl	Number of chlorine atoms in the ligand

Feature name	Description
nBr	Number of bromine atoms in the ligand
nI	Number of iodine atoms in the ligand
nX	Number of other atoms in the ligand (?)
BalabanJ	*topological indexes*
	Balaban J connectivity topological index
BertzCT	Bertz topological index meant to quantify
	"complexity"
nBonds	*bonds descriptors*
	Total number of bonds in the ligand
nBondsO	-
nBondsS	-
nBondsD	-
nBondsT	-
nBondsA	-
nBondsM	-
nBondsKS	-
nBondsKD	-
PNSA1	*Charged Partial Surface Area (CPSA)
	descriptors*
	Partial negative surface areas
PNSA2	
PNSA3	
PNSA4	
PNSA5	
PPSA1	Partial positive surface areas
PPSA2	
PPSA3	
PPSA4	
PPSA5	
DPSA1	Differences in charged partial surface areas
DPSA2	
DPSA3	
DPSA4	
DPSA5	

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Feature name	Description
FNSA1	Fractional charged partial negative surface areas
FNSA2	
FNSA3	
FNSA4	
FNSA5	
FPSA1	Fractional charged partial negative surface areas
FPSA2	
FPSA3	
FPSA4	
FPSA5	
WNSA1	Surface weighted charged partial negative surface
	areas
WNSA2	
WNSA3	
WNSA4	
WNSA5	
WPSA1	Surface weighted charged partial negative surface
	areas
WPSA2	
WPSA3	
WPSA4	
WPSA5	
RNCG	Relative negative charge
RPCG	Relative positive charge
RNCS	Relative negative charge surface area
RPCS	Relative positive charge surface area
TASA	Total hydrophobic surface area
TPSA	Total polar surface area
RASA	Relative hydrophobic surface area
RPSA	Relative polar surface area
	*end of CPSA descriptors*
ECIndex	<del>-</del>
fragCpx	Fragment complexity score
GeomDiameter	-

Feature name	Description
GeomRadius	-
GeomShapeIndex	-
GeomPetitjeanIndex	-
GRAV	-
GRAVH	-
GRAVp	-
GRAVHp	-
nHBAcc	Number of hydrogen bond acceptors
nHBDon	Number of hydrogen bond donors
Lipinski	Lipinski rule of 5. Checks whether ligand is drug-
	like or not
GhoseFilter	Ghose Filter. Checks whether ligand is drug-like
VMcGowan	McGowan volume
apol	Sum of the atomic polarizabilities (including
	implicit hydrogens) with polarizabilities
bpol	Sum of the absolute value of the difference
	between atomic polarizabilities of all bonded
	atoms in the molecule (including implicit
	hydrogens) with polarizabilities
nRot	Number of rotatable bonds
RotRatio	-
SLogP	Log of the octanol/water partition coefficient
	(including implicit hydrogens)
SMR	Molecular refractivity. This property is an atomic
	contribution model [Crippen 1999] that assumes
	the correct protonation state (washed structures)
TopoPSA(NO)	-
TopoPSA	Topological polar surface area
Diameter	Largest value in the distance matrix of the ligand
Radius	Diameter/2 (?)
TopoShapeIndex	Topological shape index
PetitjeanIndex	Petit jean index
Vabc	-
MW	Molecular weight of the ligand

Feature name	Description
AMW	-
WPath	Wiener path number
WPol	Wiener polarity number