

6lu7_pdb-Eucalyptol

Docking Scores

No.	Affinity	rmsd l.b	rmsd u.b
1	-3.8	0	0
2	-3.7	1.137	3.354
3	-3.7	1.221	2.592
4	-3.5	1.395	3.126
5	-3.4	1.288	2.357
6	-3.4	1.2	1.968
7	-3.4	1.199	3.187
8	-3.2	8.075	9.504
9	-3.2	8.041	9.662

Visualisation

Natural Ligand 1

Hydrophobic Interactions

No.	Res.	AA	Dist	Ligand Atom	Proton Atom
1	25	THR	3.73	2416	178
2	26	THR	3.81	2416	185

Hydrogen Bonds

No.	Res.	AA	Dist H-A	Dist D-A	Don Angle	Protisdon?	Sidechain?	D. Atom	A. Atom
1	143	GLY	1.93	2.8	145.29	True	False	1105[Nam]	2412[O3]

Natural Ligand 2

Hydrophobic Interactions

No.	Res.	AA	Dist	Ligand Atom	Proton Atom
1	168	PRO	3.53	2372	1303

No Hydrogen Bonds Found

Natural Ligand 3

No Hydrophobic Interactions Found

Hydrogen Bonds

No.	Res.	AA	Dist H-A	Dist D-A	Don Angle	Protisdon?	Sidechain?	D. Atom	A. Atom
1	164	HIS	2.16	3.07	153.73	False	False	2398[N3]	1266[O2]
2	166	GLU	2.61	3.38	135.52	False	True	2399[Nam]	1289[O-]

Docked Ligand Interactions

Hydrophobic Interactions Found

Hydrophobic Interactions Found

No Hydrogen Bonds Found

No Water Bridges Found

No Salt Bridges Found

No Pi Stacks Found

No Pi Cation Interactions Found

No Halogen Bonds Found

No Metal Complexes Found

Hydrophobic Interactions

No.	Res.	AA	Dist	Ligand Atom	Proton Atom
1	142	ASN	3.56	2425	1101

Figures

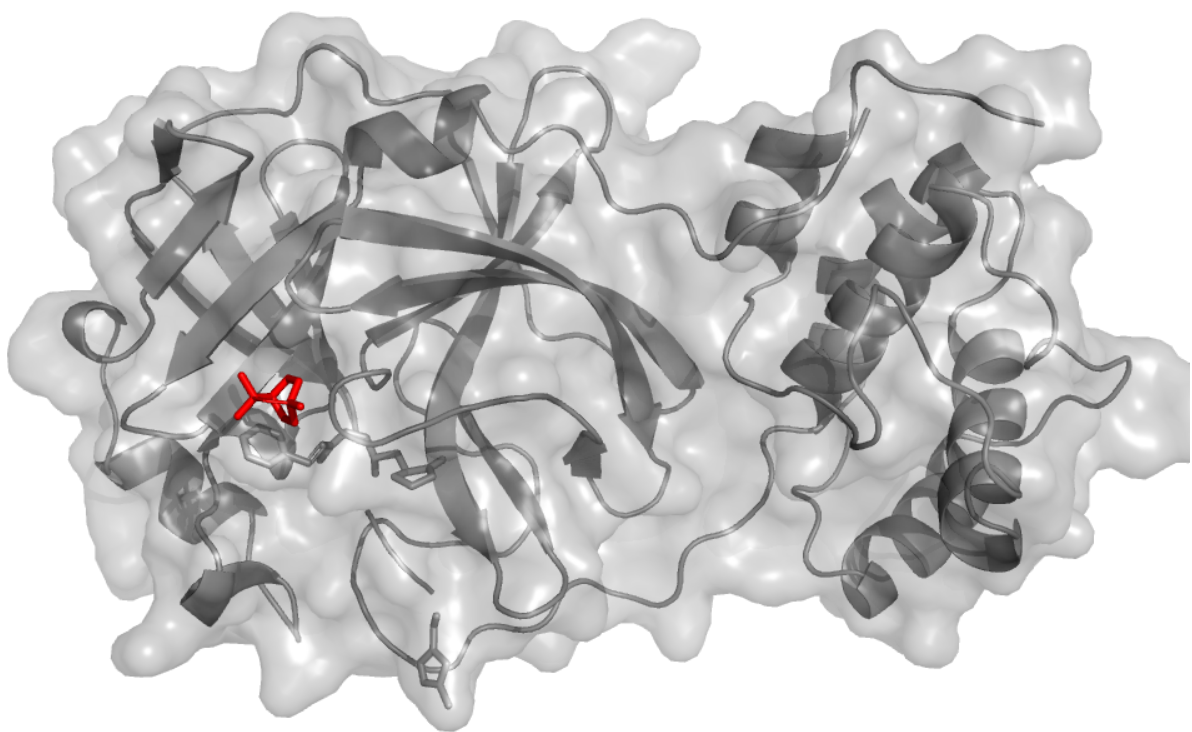


Figure 1: Back View

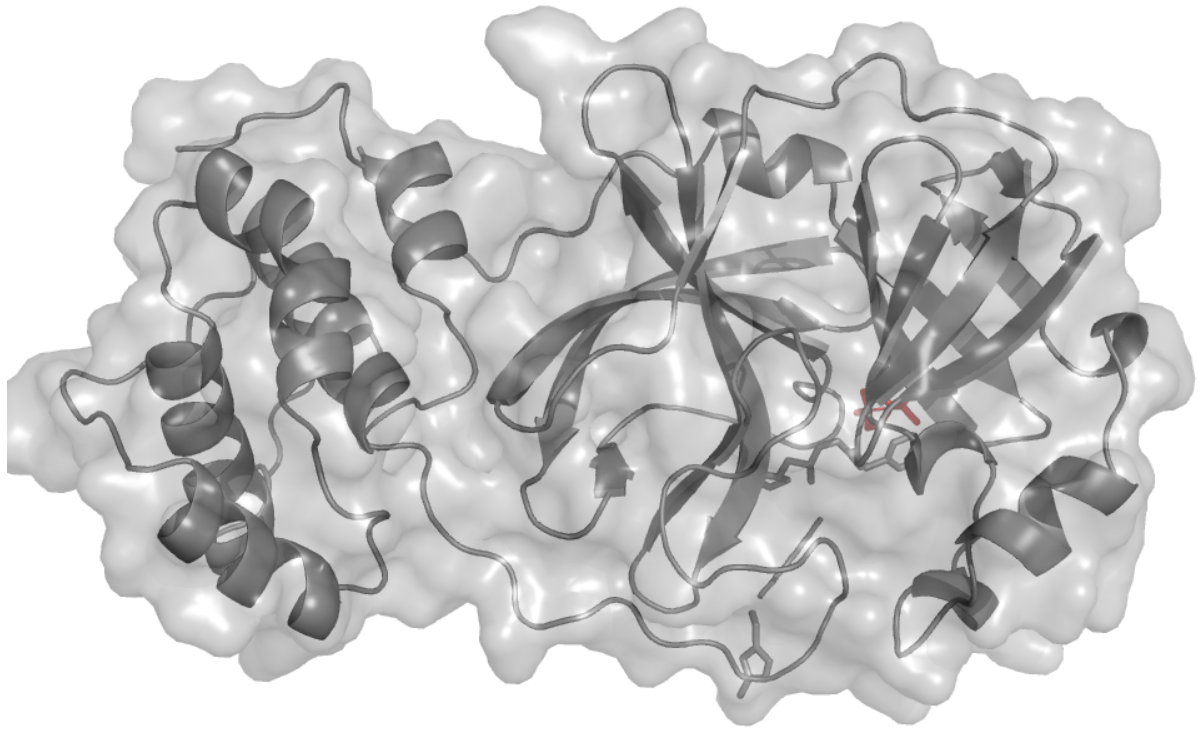


Figure 2: Front View

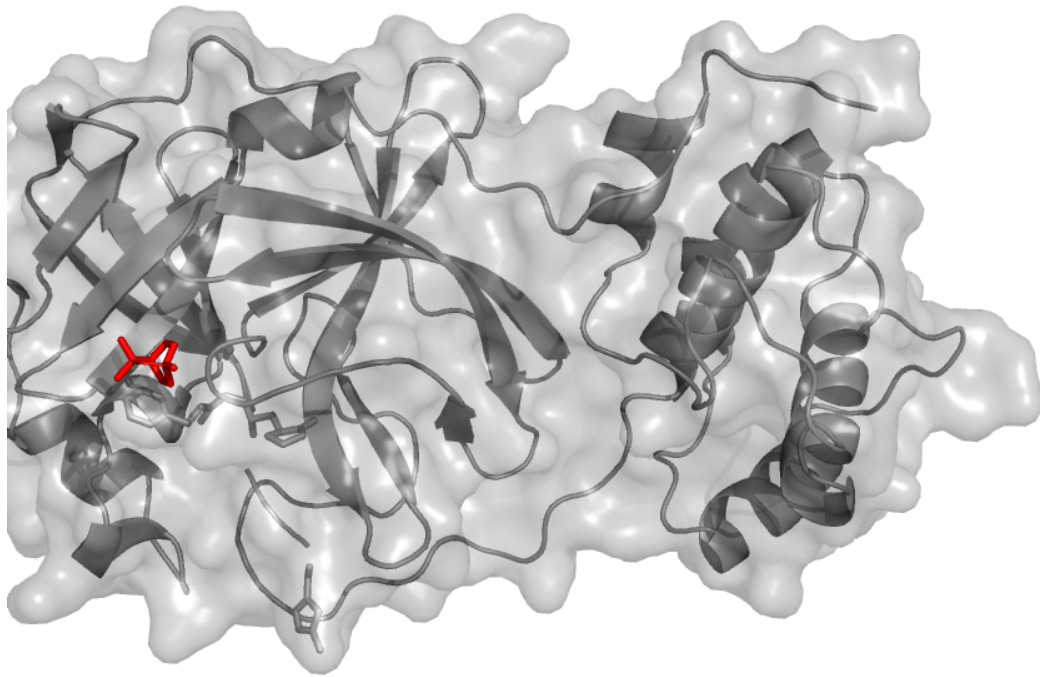


Figure 3: Close Up View of the Back

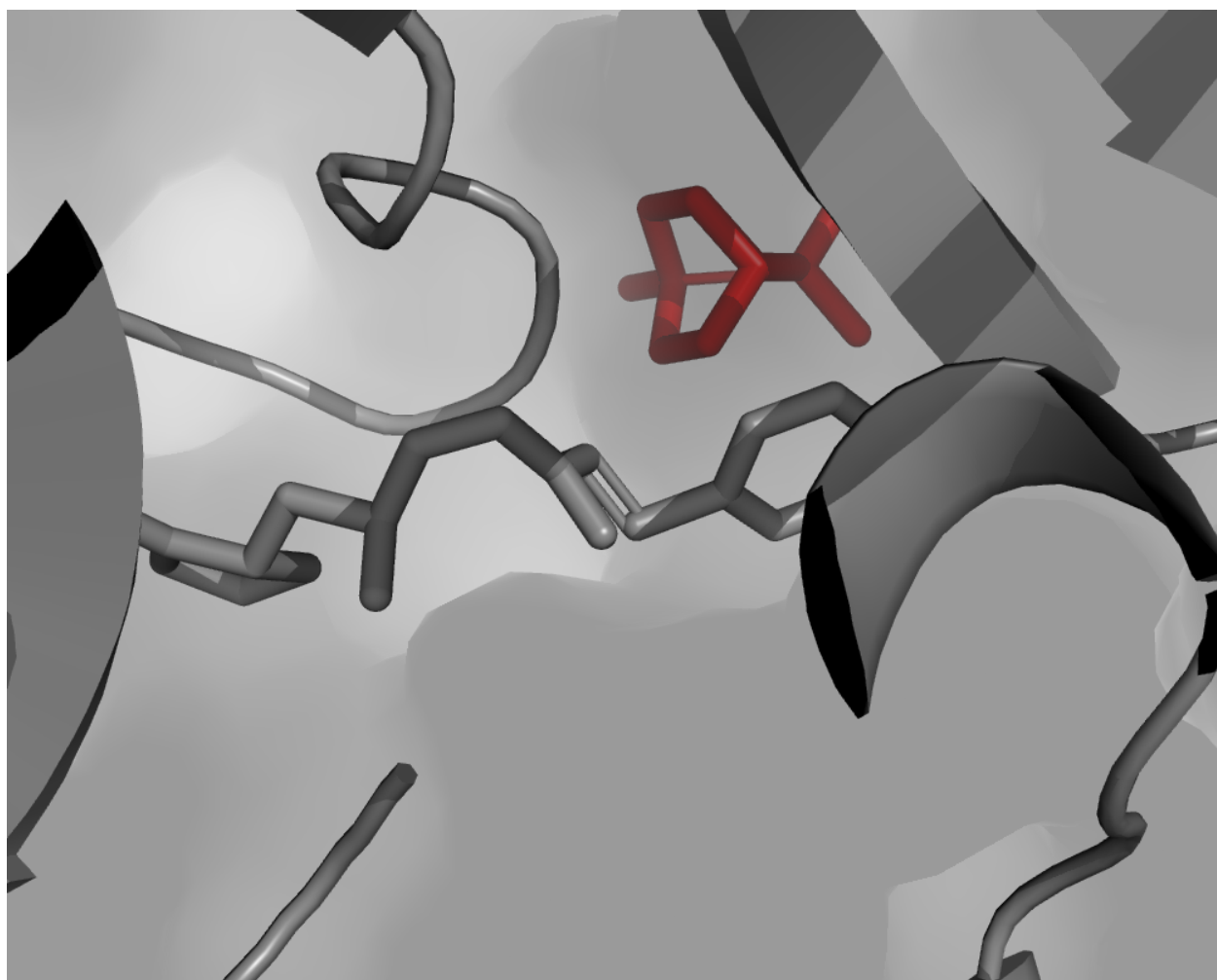


Figure 4: Close Up View of the Front