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# COMPARATIVE MODELLING AND DOCKING STUDIES OF CYTOCHROME C OXIDASE SUBUNIT 1 PROTEIN

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Keywords: Cytochrome c oxidase, Homology modeling, Modeller9.21, Docking, Natural compounds Correspondence to Author: Shravan Kumar Gunda Faculty, Bioinformatics Division, PGRRCDE, Osmania University, Hyderabad - 500007, Telangana, India. E-mail: gunda14@gmail.com **ABSTRACT:** Cytochrome c oxidase, a mitochondrial metalloenzyme acting as the terminal enzyme of the mitochondrial respiratory chain. It provides a critical function in cellular respiration in both eukaryotes and prokaryotes. The recent development in technology has enabled researchers to understand the genetic, molecular, structural, and functional properties of proteins to identify appropriate targets against diseases, and as a result, many antiinflammatory and anti-proliferative drugs have been developed. In the absence of X-ray and NMR protein crystal structure, homology modeling provides a useful 3D model for a protein that is related to at least one known protein structure. In the present study, the 3D molecular structures of nine Cytochrome c oxidase subunit 1 protein from different species such as Myxine glutinosa, Struthio camelus, Sus scrofa, Homo sapiens, Xenopus laevis, Halichoerus grypus, Zygogeomys trichopus, Buteo buteo, and Plasmodium falciparum were predicted using homology modeling software MODDLLER. The modeled structures were docked using Autodock4.2 software with ten different natural compounds and three drugs as a control to study the molecular interactions of these compounds with the coat protein. The results show that all the compounds exhibited good interactions with modeled proteins.

**INTRODUCTION:** Cytochrome c oxidase (EC 1.9.3.1) is a complex metalloprotein. Cytochrome c oxidase provides a critical function in cellular respiration in both eukaryotes and prokaryotes <sup>1</sup>. The gene COI plays a central role in metabolism, and it is present in almost all eukaryotes <sup>2</sup>. Cytochrome c oxidase I as a standard for molecular barcoding of animals, was proposed by Hebert <sup>3</sup>.



Cox1 has a phylogenetic signal system than the other mitochondrial genes <sup>4</sup>. The enzyme Cytochrome c oxidase is a terminal enzyme of mitochondria and aerobic bacteria respiratory chains. It is crucial that the complexes of the respiratory chains are combined into the membranes of bacteria and mitochondria <sup>5</sup>. Cytochrome c oxidase is a transmembrane protein, it looks like Y shaped, and is located in the inner mitochondrial membrane <sup>6</sup>.

Three subunits I, II, and III are mitochondrially encoded and are present in all eukaryotes. The enzyme catalyzes 4-electron reduction of  $O_2$  to  $H_2O^{-7}$ . In eukaryotes, the enzyme Cytochrome c oxidase is encoded in two genomes.

These three subunits I, II, and III have been isolated and sequenced from fungus, yeast and mammalian sources <sup>8</sup> and for plants <sup>9-12</sup>. Hence, the aim of the present study is to build a three-dimensional structure of nine different important Cytochrome c oxidase subunit 1 protein from different species by using homology modeling.

In addition, this study also focuses on performing a molecular docking for the identification of the effective inhibitory activity of known broadspectrum compounds using molecular docking studies.

**MATERIALS AND METHODS:** The amino acid sequences of the Cytochrome c oxidase (COX) proteins from different species Myxine glutinosa isolate (Uniprot ID: O21079), Struthio camelus (Uniport ID: O21399), Sus scrofa (Uniprot ID: O79876), Homo sapiens (Uniprot ID: P00395), Xenopus laevis (Uniprot ID: P00398), Halichoerus (Uniprot ID: P38595), Zygogeomys grypus trichopus (Uniprot ID: Q6EGH7), Buteo buteo (Uniprot ID: Q94WR7), Plasmodium falciparum (Uniprot ID: Q02766) were retrieved from the database protein sequence **UniprotKB** (www.uniprot.org).<sup>13</sup>

To find related protein templates in order to build models for these primary sequences, a sequence similarity search has been carried out separately by using Protein BLAST <sup>14</sup> tool against solved protein structures deposited in Protein Data Bank (PDB). ClustalX and ClustalW2 are used for the correction of alignment <sup>15</sup>. Modeller9.21 <sup>16</sup> was used to gain satisfactory models <sup>17</sup>. The modeller is an implementation of an automated approach to comparative modeling by satisfaction of spatial restraints, which employs position-dependent gap penalties based on structural information of the template for generating alignments <sup>18</sup>.

After manually modifying the alignment input file in MODELLER 9.21 to match the template and query sequence, 20 models were generated and selected the PDB file on the basis of the Modeller Objective Function. The selected model was subjected to a series of tests for its internal consistency and reliability. Backbone conformation was evaluated by the inspection of the psi/phi Ramachandran plot obtained from PROCHECK analysis<sup>19</sup>. Active Site Prediction: The active site for docking was predicted using the Tripos Sybyl6.7 SiteID<sup>20</sup> module after adding hydrogen atoms to the modeled proteins. The amino acids fall under active site for all the modeled proteins are shown in **Table** 2. The potential binding sites were predicted and identified by the SiteID module by correlating and combining key criteria such as depth, exposure, temperature factor. hydrophobicity, solvent accessible surface, and hydrogen-bonding capability. While detecting the potential binding sites, the SiteID module generates necessary files required for performing Autodock4.2 docking studies.

Docking Studies: Molecular docking studies were performed to elucidate the binding mode of COX proteins and the ligands. A total of ten molecules were docked to the protein to locate the appropriate binding orientations and conformations of various inhibitors in the binding pocket using the Graphical User Interface program "Autodock4.2".<sup>21</sup> Auto Dock 4.2 is an automatic docking tool designed to predict how small molecules bind to a receptor of 3D structures, which generates grids and calculates the docking score to evaluate the conformers. Gasteiger - Huckel united atom charges, polar hydrogen's and solvation parameters were added to the receptor for the preparation of protein in docking simulation, Gasteiger - Huckel charges assigned and then non-polar hydrogens were merged as the docking ligands were not peptides. All torsions were allowed to rotate during docking. The generation of PDBQT files for protein and ligands preparation, grid box creation, was carried out with Auto Dock Tools 1.5.6 (ADT) <sup>22</sup>. AutoDock saved the prepared ligand file and the PDB file in PDBQT format. Generated AutoGrid was used for the preparation of the grid map using a grid box. The grid size was set to  $60 \times 60 \times 60$ XYZ points, and the grid center was designated at dimensions (x, y and z). In Autodock, both the protein and ligands are considered as rigid. The binding poses with the lowest docked energy belonging to the top-ranked cluster were selected as the final model for post-docking analysis.

### **RESULTS AND DISCUSSION:**

**Homology Modelling of Proteins:** After protein BLAST of the primary sequences of COX proteins with the predetermined structures deposited in PDB bank, sequences that showed the greatest similarity were considered as template sequences. Twenty models were generated using the MODELLER 9.21 program based on the sequence alignment files generated by the ClustalX program. The alignment file was tweaked manually to best fit the sequences. Among the generated models for all the primary sequences, the model with the least object function was selected for further evaluation for protein stereochemistry (phi and psi angles) with PROCHECK software. The final models were validated using PROCHECK. The PROCHECK software generates a number of files that list complete residue by residue data and the assessment of the overall quality of generated structure compared to well-refined structures of the same resolution.

Fig. 1 shows the predicted homology models of all the Cytochrome c oxidase subunit 1 proteins belonging to different species. The Ramachandran plots for all the generated models are shown in Fig. 2. Table 1 shows the residues falling in different regions of the Ramachandran plot. The results show that there are no amino acids falling in the disallowed region of the plot and a minor percentage of amino acids in the generously allowed regions, which indicate that the models are the better conformational structures. The template protein showed 87.7% of amino acid residues in the most favored region, 11.3% of amino acid residues in the additionally allowed region, 0.6% of amino acids in the generously allowed region, and 0.4% of amino acids in the disallowed region. Loop building was performed by using SPDBV.

TABLE 1: % OF RESIDUES FALLING IN DIFFERENT REGIONS OF RAMACHANDRAN PLOT OFCYTOCHROME C OXIDASE SUBUNIT 1 PROTEINS FROM DIFFERENT SPECIES

S.	Name	Core re	gion	Allowed	region	Generously all	owed region	Disallowed	region
no.	of species	No. of	%	No. of	%	No. of	%	No. of	%
		residues		residues		residues		residues	
1	O21079	416	94.3	24	5.4	1	0.2	0	0
	Myxine glutinosa								
2	O21399	416	95.4	18	4.1	2	0.5	0	0
	Struthio camelus								
3	O79876	419	95.9	15	3.4	3	0.7	0	0
	Sus scrofa								
4	P00395	416	95.4	16	3.7	4	0.9	0	0
	Homo sapiens								
5	P00398	414	93.0	28	6.3	3	0.7	0	0
	Xenopus laevis								
6	P38595	409	93.6	24	5.5	4	0.9	0	0
	Halichoerus grypus								
7	Q6EGH7	416	95.0	19	4.3	4	0.7	0	0
	Zygogeomys trichopus								
8	Q94WR7	413	94.7	18	4.1	5	1.1	0	0
	Buteo buteo								
9	Q02766	393	94.7	17	4.1	5	1.2	0	0
	Plasmodium falciparum								

**Docking Studies of Modeled Proteins:** The active site for docking was predicted using the Tripos Sybyl 6.7 SiteID module, and the active site is shown in **Table 2**. During the docking procedure, selected only the best fit, active site pocket with respect to the ligands in order to dock them. AutoDock 4.2 was used for molecular docking studies. Results obtained from AutoDock 4.2 provided information on the binding orientation of ligand-receptor interactions. Free energies of binding ( $\Delta$ Gb) and dissociation constants (Ki) as calculated by AutoDock4.2. The modeled protein was docked with natural compounds consisting of Plant compounds like flavonoids, phenolic compounds, alkaloids, *etc.* The docking programs place both the ligand and protein molecule in various orientations, conformational positions, and the lowest energy confirmations, which are energetically favorable are evaluated and analyzed for interactions. All the ten molecules that were docked showed good interactions.

From the docking simulation, we observed the free energy charge of binding for the protein-ligand complex. A possible explanation may be that the radio-graphical structure of the protein from crystals differs from that of the aqueous system. The total energy was calculated for all the compounds against different targets are summarized in **Table 3-11**. Therefore, it can be concluded that the interaction of all the compounds with the corresponding targets is hydrophobic interactions in nature. Molecular docking indicated that the distance of hydrogen bonding between the ligand and the protein, respectively.

Molecular interaction between Cycloartocarpin A with different targets calculated the docking score. The modeled protein was used for molecular docking analysis. It was found that Cycloartocarpin A made a hydrogen bond with Gly355 with binding energy of -10.27 kcal/mol a P00395-Homo sapiens Cytochrome c oxidase subunit 1 protein. The compound also showed best interactions with Trp127 and Arg439 with binding energy of -10.49 kcal/mol an O21399-*Struthio camelus* protein.

It also shows Hydrogen bond interaction with the amino acid residue Met178 of modeled protein (Uniprot ID: O21079-Myxine glutinosa), a COX target with a binding energy of -6.26 kcal/mol. Four proteins showed the highest binding energy, and interactions with Cycloartocarpin A and other proteins showed the highest binding energy and interactions with other proteins. Similarly, we docked with the following targets O79876\_Sus scrofa, P00395-Homo sapiens, P00398-Xenopus laevis, P38595-Halichoerus grypus, Q6EGH7-Zygogeomys trichopus, Q94WR7-Buteo buteo, Q02766-Plasmodium falciparum. Total ten natural compounds were docked with all the nine modeled proteins, and the results of the highest binding energy conformations and interacting amino acids are shown in Fig. 3. All the natural compounds exhibited good binding energies and interactions than standard drugs.



FIG. 1: HOMOLOGY MODELS OF CYTOCHROME C OXIDASE SUBUNIT 1 PROTEINS [A] MYXINE GLUTINOSA (UNIPROT ID: O2179), [B]STRUTHIO CAMELUS (UNIPORT ID: O21399), [C] SUS SCROFA (UNIPROT ID: O79876), [D] HOMO SAPIENS (UNIPROT ID: P00395), [E] XENOPUS LAEVIS (UNIPROT ID: P00398), [F] HALICHOERUS GRYPUS (UNIPROT ID: P38595), [G]ZYGOGEOMYS TRICHOPUS (UNIPROT ID: Q6EGH7), [H]BUTEO BUTEO(UNIPROT ID: Q94WR7), [I]PLASMODIUM FALCIPARUM(UNIPROT ID: Q02766).







FIG. 2: RAMACHANDRAN PLOTS OF CYTOCHROME C OXIDASE SUBUNIT 1 PROTEINS FROM DIFFERENT SPECIES

#### TABLE 2: LIGAND BINDING POCKETS AS DETECTED BY USING SITEID MODULE OF TRIPOS SYBYL6.7 SOFTWARE

S. no.	Name of Species	Amino acids fall under receptor site
1	O21079_Myxine	His151, Ala153, Gly154, Ser156, Gly160, Ala161, Phe164, Thr167, Ile168, Ile188, Leu199, Leu202,
	glutinosa	Ala203, Phe238, Glu242, Phe67, Ile75, Asn80, Val83, Met92, Asn98, Ser101, Leu105
2	O21399_Struthio	Asn136, Val144, Ala147, Ile148, Leu207, Leu210, Tyr232, Thr60, Phe64, Phe68, Ser116, Ser117,
	camelus	Ala123, Thr125, Thr128, Val129
3	O79876_Sus scrofa	Gly239, Glu242, Val243, Leu246, His291, Thr31, Ser34, Tyr371, Val374, Ile37, Leu381, Arg38,
		Val421, Thr424, Phe425, Arg438, Arg439, Ser458, Tyr54, Asn55, Ile57, Val58, Thr59, His61, Ala62,
		Met65, Ile66, Val70, Gly123, Gly125, Trp126
4	P00395_Homo sapiens	Trp236, Gly239, Glu242, Val243, Leu246, His291, Thr31, Ser34, Tyr371, Ile37, Val374, Phe377,
		Leu381, Val386, Met390, Leu41, Met417, Thr424, Phe425, Gln428, Arg438, Arg439, Tyr54, Ile57,
		Val58, His61, Ala62, Met65, Ile66, Val70, Trp126
5	P00398_Xenopus	Phe184, His240, Val243, Tyr244, Leu246, Ile247, Leu248, Pro249, Phe251, Met277, Ile280, His290,
	laevis	His291, Thr309, Ile312, Thr316, Val320, Phe344, Ile345, Phe348, Gly352, Gly355, Leu358, Ala359,
		Asp364, His368, Val373, His376, Val380, Leu381, Ala385, Val70, Met74, Phe78
6	P38595_Halichoerus	Tyr261, Pro336, Met390, Phe393, Val394, His395, Pro398, Leu399, Tyr403, Leu405, Ala410, His413,
	grypus	Leu467, Met468, Met471, Glu481, Ala484, Trp494, Cys498.
7	Q6EGH7_Zygogeomys	Thr17, Met20, Trp340, Met390, Phe393, Val394, His395, Phe397, Leu399, Phe400, Leu405, Ala410,
	trichopus	His413, Met468, Met471, Ile472, Ser484, Trp494, His496, Cys498
8	Q94WR7_Buteo buteo	Gly135, Asn136, Val144, Ala147, Ile148, Leu151, Ile207, Leu210, Ala27, Val30, Thr60, Ala61, Val65,
		Met66, Phe69, Phe110, Leu113, Leu114, Ser116, Ser117, Ala123, Val129
9	Q02766_Plasmodium	Gly126, Gly127, Leu133, Pro146, Val147, Ile152, Gly155, Leu156, Leu208, Gly211, Val212, Tyr237,
	falciparum	Leu240, Phe241, Tyr31, Met61, Ile62, Ile65, Ile68, Ile69, Phe72, Phe73, Phe114, Val117, Thr121

# TABLE 3: BINDING ENERGY AND PROTEIN LIGAND INTERACTIONS OF NATURAL COMPOUNDS AGAINST MODELLED O21079\_MYXINE GLUTINOSA PROTEIN

S. no.	Ligand	Interactions	Binding Energy ΔG (Kcal/Mol)	Disassociation Constant (µM)
1	Artocarpin	Asn170 (2)	-4.95	234.15
2	Avenanthramide A	Phe505, Lys172 (2)	-6.17	300.0
3	Avenanthramide B	Asn170, Met176, Lys172	-5.56	83.90
4	Cycloartocarpin A	Asn170	-5.99	40.95
5	Cyclokievitone	Met178	-6.26	25.92
6	Eriosemaone A	Gly88, Lys172	-5.94	44.43
7	Khonklonginol A	Met171, Thr173	-5.77	58.69
8	Khonklonginol F	-	-5.98	41.33
9	Khonklonginol H	Asn170	-5.86	51.00
10	Racemosol	Lys172	-5.85	51.91
11	Celecoxib*	Gly88, Met171	-5.94	44.51
12	Valdecoxib*	Lys172	-5.80	56.52
13	Rofecoxib*	Gly88, Met 171	-6.45	18.58

# TABLE 4: BINDING ENERGY AND PROTEIN LIGAND INTERACTIONS OF NATURAL COMPOUNDS AGAINST MODELLED 021399\_STRUTHIO CAMELUS PROTEIN

S. no.	Ligand	Interactions	Binding Energy ΔG (Kcal/Mol)	Disassociation Constant (µM)
1	Artocarpin	Trp127, Arg439	-9.24	160.86 μM
2	Avenanthramide A	Trp127, His369, His291,	-8.08	1.20 μM
		Arg439		
3	Avenanthramide B	Trp127, Arg439	-7.52	3.06 µM
4	Cycloartocarpin A	Trp127, Arg439	-10.49	20.40 nm
5	Cyclokievitone	Arg439, His369	-7.83	1.83 μM
6	Eriosemaone A	Trp127, His292, Arg439	-9.92	53.15 nm
7	Khonklonginol A	Arg439	-9.27	161.59 nm
8	Khonklonginol F	Trp127	-10.80	12.00 nm
9	Khonklonginol H	Trp127, His292	-9.44	120.29 nm
10	Racemosol	Trp127, His292	-9.22	175.01 nm
11	Celecoxib*		-8.26	885.50 nM
12	Valdecoxib*	Arg439	-8.35	752.19 nM
13	Rofecoxib*	Trp127, Arg439, Asp365,	-9.51	106.40 nM
		His291		

### TABLE 5: BINDING ENERGY AND PROTEIN LIGAND INTERACTIONS OF NATURAL COMPOUNDS AGAINST MODELLED 079876 SUS SCROFA PROTEIN

S. no.	Ligand	Interactions	Binding Energy ΔG (Kcal/Mol)	Disassociation Constant (µM)
1	Artocarpin	Tyr54, Arg439	-9.53	103.93 nM
2	Avenanthramide A	Tyr371, Arg438, Arg439,	-7.62	2.61 μM
		Trp126, Glu242		
3	Avenanthramide B	Arg38, Arg439	-6.31	23.56 μM
4	Cycloartocarpin A		-8.23	924.96 nM
5	Cyclokievitone	His378, Arg38	-7.78	1.97 μM
6	Eriosemaone A	His61, His378, Glu242	-9.04	263.63 nM
7	Khonklonginol A	Tyr371	-9.19	182.46 nM
8	Khonklonginol F	Tyr371	-8.95	277.22 nM
9	Khonklonginol H	Arg438	-9.47	114.98 nM
10	Racemosol	Arg439, Tyr371	-9.32	148.25 nM
11	Celecoxib*	Gln428	-6.84	9.63 μM
12	Valdecoxib*	Arg438, Ala62	-7.91	1.6 µM
13	Rofecoxib*	Arg38, Arg439	-8.40	700.42 nM

# TABLE 6: BINDING ENERGY AND PROTEIN LIGAND INTERACTIONS OF NATURAL COMPOUNDS AGAINST MODELLED P00395\_HOMO SAPIENS PROTEIN

S. no.	Ligand	Interactions	Binding Energy ∆G (Kcal/Mol)	Disassociation Constant (µM)
1	Artocarpin	Tyr126, His368	-6.96	7.85 μM
2	Avenanthramide A	Gly355	-9.71	76.58 nM
3	Avenanthramide B	Tyr126, His291, Arg438	-5.85	51.4 μM
4	Cycloartocarpin A	Gly355	-10.27	29.61 nM
5	Cyclokievitone	Asp364, His368	-8.68	431.36 nM
6	Eriosemaone A	His368, Gly355	-9.65	84.33 nM
7	Khonklonginol A	Asp364, Ala359	-9.99	47.26 nM
8	Khonklonginol F	His376	-9.99	47.81 nM
9	Khonklonginol H	His368, Asp 364	-10.09	39.96 nM

10	Racemosol	His368, Asp364	-9.67	81.62 nM
11	Celecoxib*	His368, His290, Asp364	-8.94	277.51nM
12	Valdecoxib*	His240, His291, Thr316	-8.65	453.7nM
13	Rofecoxib*	His368, His 376, Asp 364,	-9.33	145.94nM
		His290		

# TABLE 7: BINDING ENERGY AND PROTEIN LIGAND INTERACTIONS OF NATURAL COMPOUNDS AGAINST MODELLED P00398\_XENOPUS LAEVIS PROTEIN

S. no.	Ligand	Interactions	Binding Energy ∆G (Kcal/Mol)	Disassociation Constant (µM)
1	Artocarpin	Gly355	-8.82	341.34 nM
2	Avenanthramide A	Asp364, Thr316, Gly355	-8.25	896.97 nM
3	Avenanthramide B	-	-5.64	72.95 μM
4	Cycloartocarpin A	His376, Gly355	-10.72	13.82 nM
5	Cyclokievitone	His240, His290, His368,	-9.02	243.68 nM
		Leu358		
6	Eriosemaone A	His368, Asp364	-9.31	150.29 nM
7	Khonklonginol A	-	-9.09	217.61 nM
8	Khonklonginol F	-	-9.56	98.23 nM
9	Khonklonginol H	-	-9.24	168.06 nM
10	Racemosol	His376	-9.14	199.08 nM
11	Celecoxib*	-	-8.23	930.24 nM
12	Valdecoxib*	His240, His290	-8.11	1.14 μM
13	Rofecoxib*	His368, Asp364	-9.21	177.51 nM

### TABLE 8: BINDING ENERGY AND PROTEIN LIGAND INTERACTIONS OF NATURAL COMPOUNDS AGAINST MODELLED P38595\_HALICHOERUS GRYPUS PROTEIN

S. no.	Ligand	Interactions	Binding Energy ∆G (Kcal/Mol)	Disassociation Constant (µM)
1	Artocarpin	Lys13, Phe400, Tyr502	-6.90	8.81 μM
2	Avenanthramide A	Thr17, Thr502, Thr17	-7.55	2.91 μM
3	Avenanthramide B	Lys13, Phe400, Tyr17	-6.33	22.9 μM
4	Cycloartocarpin A	Lys13	-9.04	236.84 nM
5	Cyclokievitone	Lys13, Pro500, Ser401	-7.92	1.55 μM
6	Eriosemaone A	Thr17	-8.34	771.67 nM
7	Khonklonginol A	Lys13, Thr17, Pro500	-8.35	762.44 nM
8	Khonklonginol F	Lys13, Pro500	-8.12	1.12 μM
9	Khonklonginol H	Ser401	-7.85	1.77 μM
10	Racemosol	Ser401, Pro500	-8.91	292.46 nM
11	Celecoxib*	Lys13	-7.98	1.14 μM
12	Valdecoxib*	Lys13, ASP14	-8.26	877.06nM
13	Rofecoxib*	Lys13	-8.58	650.82nM

# TABLE 9: BINDING ENERGY AND PROTEIN LIGAND INTERACTIONS OF NATURAL COMPOUNDS AGAINST MODELLED Q6EGH7\_ZYGOGEOMYS TRICHOPUS PROTEIN

S. no.	Ligand	Interactions	Binding Energy ΔG (Kcal/Mol)	Disassociation Constant (µM)
1	Artocarpin	Arg5, Lys13, Asp14	-9.78	67.63 nM
2	Avenanthramide A	Lys13, Pro500, Asp14	-8.03	1.29 μM
3	Avenanthramide B	Lys13, Thr401	-8.04	1.28 μM
4	Cycloartocarpin A	-	-6.66	13.21 μM
5	Cyclokievitone	Lys13	-8.15	1.07 μM
6	Eriosemaone A	Arg5	-8.17	1.03 μM
7	Khonklonginol A	Arg5, Asp14, Pro500	-8.59	504.7 nM
8	Khonklonginol F	Arg480, Thr401	-9.10	213.32 nM
9	Khonklonginol H	-	-9.13	204.0 nM
10	Racemosol	Thr401	-8.74	392.28 nM
11	Celecoxib*	Lys13	-8.08	1.19 μM
12	Valdecoxib*	Lys13, Pro500	-8.60	500.75Nm
13	Rofecoxib*	Asn11, Lys13, Asp14	-8.41	650.82nM

# TABLE 10: BINDING ENERGY AND PROTEIN LIGAND INTERACTIONS OF NATURAL COMPOUNDS AGAINST MODELLED Q94WR7\_*BUTEO BUTEO* PROTEIN

S. no.	Ligand	Interactions	Binding Energy ΔG (Kcal/Mol)	Disassociation Constant (µM)
1	Artocarpin	Ala123, Ser143	-6.83	9.43 μM
2	Avenanthramide A	Ala123, Ser143	-7.72	2.19 μM
3	Avenanthramide B	-	-5.68	68.95 μM
4	Cycloartocarpin A	Val119	-6.96	7.88 µM
5	Cyclokievitone	Ser117	-7.47	3.35 µM

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6	Eriosemaone A	Leu113	-7.21	5.21 μM
7	Khonklonginol A	-	-7.43	3.59 µM
8	Khonklonginol F	Ala123	-7.47	3.37 µM
9	Khonklonginol H	-	-6.85	9.47 μM
10	Racemosol	Ala123	-7.05	6.80 µM
11	Celecoxib*	-	-7.46	3.43 µM
12	Valdecoxib*	-	-7.72	2.17 µM
13	Rofecoxib*	Ala123	-8.94	279.83nM

TABLE 11: BINDING ENERGY AND PROTEIN LIGAND INTERACTIONS OF NATURAL COMPOUNDS AGAINST MODELLED Q02766 PLASMODIUM FALCIPARUM PROTEIN

S. no.	Ligand	Interactions	<b>Binding Energy ΔG (Kcal/Mol)</b>	Disassociation Constant (µM)
1	Artocarpin	-	-4.52	488.63 μM
2	Avenanthramide A	-	-4.01	1.16 μM
3	Avenanthramide B	Thr209	-3.43	3.08 mM
4	Cycloartocarpin A	-	-5.21	152.82 μM
5	Cyclokievitone	Phe154	-4.68	371.81 μM
6	Eriosemaone A	-	-5.33	124.09 µM
7	Khonklonginol A	-	-5.22	148.48 µM
8	Khonklonginol F	-	-5.15	167.27 μM
9	Khonklonginol H	-	-5.18	159.22 μM
10	Racemosol	-	-5.29	132.15 μM
11	Celecoxib*	-	-6.64	13.48 µM
12	Valdecoxib*	Thr209	-4.99	221.22 μM
13	Rofecoxib*	Thr209	-5 32	126.61 uM



FIG. 3: INTERACTIONS OF CYCLOKIEVITONE INTERACTIONS WITH CYTOCHROME C OXIDASE SUBUNIT 1 FROM *MYXINE GLUTINOSA* [B] CYCLOARTOCARPIN A INTERACTIONS WITH CYTOCHROME C OXIDASE SUBUNIT 1 FROM *STRUTHIO CAMELUS* [C] ARTOCARPIN INTERACTIONS WITH CYTOCHROME C OXIDASE SUBUNIT 1 FROM *SUS SCROFA*,[D] CYCLOARTOCARPIN A INTERACTIONS WITH CYTOCHROME C OXIDASE SUBUNIT 1 FROM *HOMO SAPIENS* [E] CYCLOARTOCARPIN A INTERACTIONS WITH CYTOCHROME C OXIDASE SUBUNIT 1 FROM *HOMO SAPIENS* [E] CYCLOARTOCARPIN A INTERACTIONS WITH CYTOCHROME C OXIDASE SUBUNIT 1 FROM *XENOPUS LAEVIS*, [F] CYCLOARTOCARPIN A INTERACTIONS WITH CYTOCHROME C OXIDASE SUBUNIT 1 FROM *HALICHOERUS GRYPUS*, [G] ARTOCARPIN INTERACTIONS WITH CYTOCHROME C OXIDASE SUBUNIT 1 FROM *ZYGOGEOMYS TRICHOPUS*, [H] AVENANTHRAMIDE A INTERACTIONS WITH CYTOCHROME C OXIDASE SUBUNIT 1 FROM *BUTEO BUTEO*, [I] RACEMOSOL INTERACTIONS WITH CYTOCHROME C OXIDASE SUBUNIT 1 FROM *PLASMODIUM FALCIPARUM* 

**CONCLUSION:** In the present study, the stable three-dimensional models were constructed for Cytochrome c oxidase protein from nine different species and further used for molecular docking with the natural inhibitors. Molecular docking results revealed some important amino-acid residues in the active sites of Cytochrome c oxidase protein, which play a key role in the maintenance of their conformation and are directly associated with substrate binding. The interactions between the ligands and the binding sites of Cytochrome c oxidase protein shown in the present study are useful for the understanding of the binding mechanisms of inhibitors and active site information of these proteins. The use of combinatorial approaches may result in the rapid development of better anti-inflammatory and anti-proliferative in future. Successful docking methods search highdimensional spaces effectively and use a scoring function that correctly ranks candidate dockings. Molecular docking studies of the above compounds showed favorable interactions in the binding site of the modeled COX proteins. Therefore, it is concluded that these molecules are the potential candidates for cytochrome c oxidase inhibitors.

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