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**Computational analysis of COX-1 & COX-2 and finding out  
their potent inhibitors**

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**Abstract**

The various NSAID's known to the scientists till date, reduces fever and inflammation when the body gets overzealous in its defenses against infection and damage but it may slows blood flow and blood clotting, reducing the chance of stroke and heart attack in susceptible individuals. Three-dimensional structures of pharmacologically important macromolecules offer a route to the discovery of new drugs. Understanding the macromolecule-ligand interactions and validation of method used for docking and virtual screening of chemical databases is crucial step in structure-based design. We therefore carried out molecular docking for structurally diverse COX-1/COX-2 inhibitors including traditional NSAIDs and Autodock 4.1.2. The complete computational analysis has revealed the best possible ligands combinations for the selective inhibition of COX-2 and COX-1. 3-D Structure of COX-2 has been predicted using the homology modeling tools. Results of docking of bound ligands like Tenoxicam and Valdecoxib have given the best binding scores Autodock.4.1.2. Molecular docking of structurally diverse selective COX-2 and COX-1 inhibitors has been successfully carried out.

Key-Words: Cyclooxygenase (COX-1, COX-2), Classic NSAIDs, Selective COX-2 Inhibitors, Inflammation, Docking, Ligplot, Inhibition

**Introduction**

Non-steroidal anti-inflammatory drugs (NSAIDs) are amongst the most widely used therapeutics, primarily for the curing of pain and inflammation, especially arthritis. From a historical point of view, the first NSAID with therapeutic reimbursement was aspirin, which has now been applied for more than 100 years as an NSAID. The generally worldwide production of about 50 000 tons a year reflects the importance of this substance even today [1]. In the 1970s, a scientific breakthrough occurred with the elucidation of the molecular mechanism of aspirin and other NSAIDs. Vane, Samuelson and Bergstrom succeeded in illustrate that these anti-inflammatory matter block the biosynthesis of prostaglandins (PGs) which contribute to a range of physiological and pathophysiological functions. *Figure 1* recapitulates the biosynthesis of PGs: the preliminary step in the biosynthesis of prostanoids is the emancipation of arachidonic acid (AA) from the phospholipids of the cell film catalyzed by phospholipase A2.

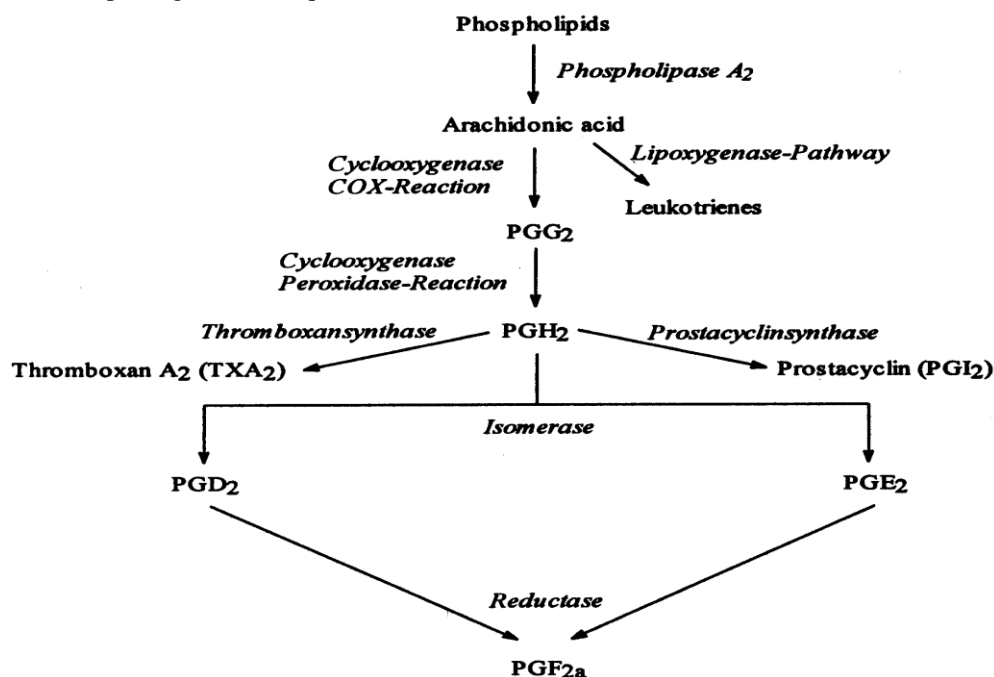
The following important step is the biotransformation of AA by cyclooxygenase. In a bifunctional action, this first produces the unsteady PGG<sub>2</sub>, the cyclooxygenase response itself, which is then instantly converted into PGH<sub>2</sub> by the same enzyme in a peroxidase reaction. As shown in *figure 1*, the ending products of the AA metabolism are PGs, thromboxanes and prostacyclin [2–5]. PGs are generated by most cells and are also current in tissues, which clarify their lane spectrum of biological responses. PGs reconcile a number of characteristic features of the body's reaction to tissue injury or inflammation. The outstanding effects of the PGs include their cytoprotective properties in the gastrointestinal (GI) tract and arrange of renal tasks in the kidney. PGE<sub>2</sub> is the most main PG which mediates the characteristic symptoms of inflammation: rubor, calor, tumor, and dolor. Dilatation of small blood vessels initiates the progress of redness and heat; the increase in vascular permeability causes the characteristic inflammation of tissues. Moreover, PGs sensitize peripheral nerve finish and nociceptors to spread pain signals to the brain and the spinal cord. In adding to the well-accepted proinflammatory role of PGs, there are also details of anti-inflammatory action in certain COX-2-derived PGs in vivo, an experiment lately reported by Gilroy et al. [6]. Like aspirin, all other NSAIDs such as ibuprofen, ketoprofen and naproxen extend their mode of action by blocking

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cyclooxygenase. Therefore, group of NSAIDs, for example to luxury inflammatory diseases such as osteoarthritis or rheumatoid arthritis, unavoidably leads to a lack of the prostaglandins requisite for the

physiological functions revealed above. Therapeutic effects and side-effects of this class of anti-inflammatory drugs are narrowly related to their biochemical mechanism of action.



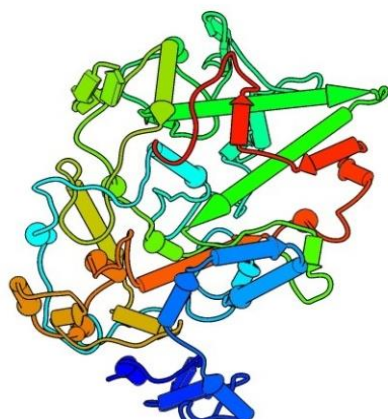
As a outcome, long-term NSAID users endure from a high incidence of GI irritation or, in the worst case, from the progress of life threatening GI ulcers and bleeding. These lesions can lead to improved morbidity in patients [7–9]. Administration of NSAIDs may also lead to renal confusions and have hypertensive effects. Due to a compressed production of PGs, such as PGI<sub>2</sub>, PGE<sub>2</sub> and PDG<sub>2</sub>, in the ruling of renal blood circulation, the rate of glomerularic filtration is condensed. Especially in patients with decreased renal function, this leads to maintenance of water, hypertension and, in some cases, to renal failure [10–12]. The reticence of cyclooxygenase in thrombocytes results in decreased production of thromboxane A<sub>2</sub>. This phenomenon extends bleeding time and leads to inhibition of platelet aggregation. A severe side-effect of NSAIDs is bronchoconstriction with resulting asthmatic events. The condensed amount of bronchodilating PGE<sub>2</sub> on the one hand and a alter in the metabolic lane from the cyclooxygenase pathway to the 5-lipoxygenase pathway on the other hand, seems to be dependable for the bronchoconstriction cause of NSAIDs [13]. The latter pathway metabolizes ‘overflow’ AA, which cannot be changed by the blocked cyclooxygenase pathway. The resultant leukotrienes act as bronchoconstrictors [14]. Because of these problems, a main target of drug research is the

progress of NSAIDs with anti-inflammatory and analgesic action but with no side effects.

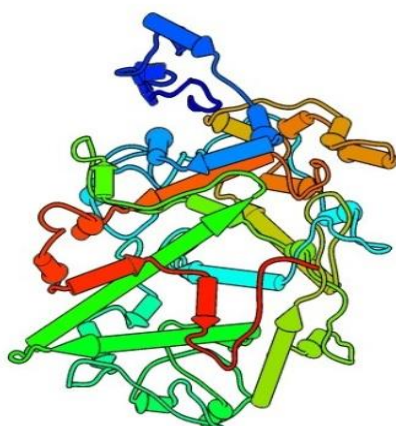
### Material and Methods

Steps involved in carrying out this study are as follows:

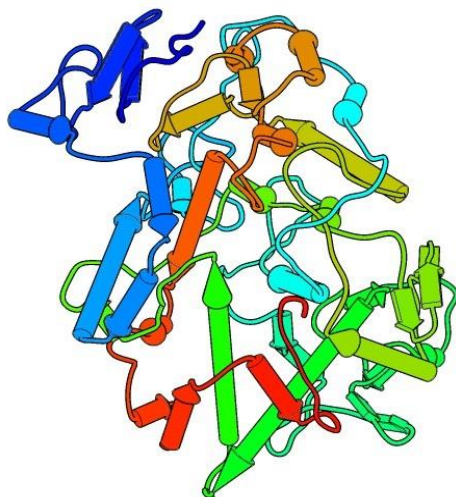
1. Sequence retrieval of COX-1 and COX-2 from GenBank. Protein sequences of COX-1 and COX-2 were retrieved from Genbank that were converted into FASTA format.
2. The sequences were then subjected to BLASTp for identification of local regions and a sequence with maximum similarity. On the basis of the template sequence Homology modeling between the retrieved sequences and the highly similar sequence was done which provides a structure of query sequence (COX-2).
3. After Homology modeling structure refinement was done which is based on energy criteria and other useful parameters for further structure refinement and optimization.
4. The structure are been downloaded from protein data bank (rcsb.org) i.e. 1HT8, 3MQE, 3NTG, 1PGF are given below.



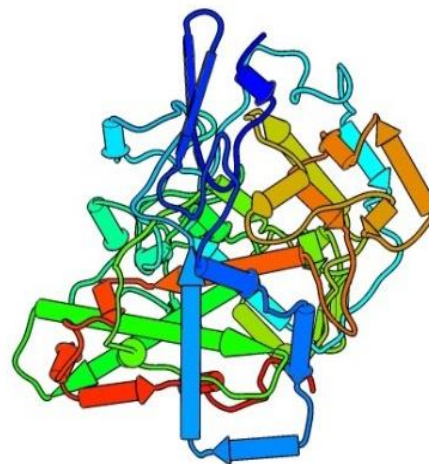
Structure of 1HT8 (Pipes and Planks)



Structure of 3MQE (Pipes and Planks)



Structure of 3NTG (Pipes and Planks)



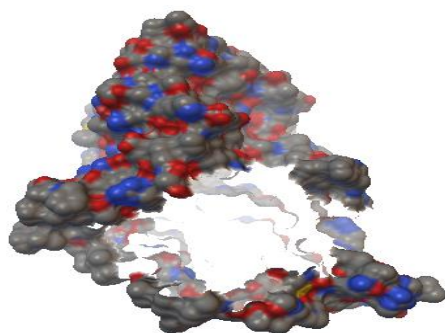
Structure Of 1PGF(Pipes and Planks)

5. Protein cleaning is done with the help of UCSF Chimera ([www.cgl.ucsf.edu/chimera/](http://www.cgl.ucsf.edu/chimera/)) and PNV.
6. Energy is minimized by SPDBV ([www.spdbv.vital-it.ch/](http://www.spdbv.vital-it.ch/)).
7. For docking, ligands were retrieved from drug bank and their physicochemical properties were studied. On the basis of these properties targeted ligand molecules were used for docking. Table No. 1
8. A priority among the ligands was generated.
9. Energy parameters, binding affinity, simulations and Autodock 4.2.1, provide the best possible combinations of COX-2, COX-1 and ligand molecules. Showing in table no. 2,3,4,5 respectively.

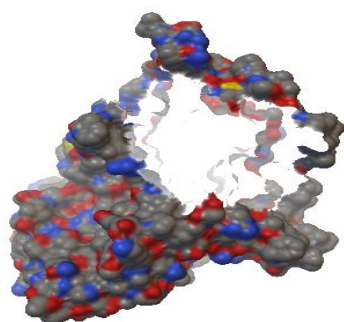
#### Binding site Prediction

Binding sites were characterized by CASTp [15] Q-Site finder and compared by extensive literature search. By comparing prediction of CASTp algorithm and Q-Site Finder, best active sites were selected. CASTp method was used to identify and measure the binding sites, active sites, surface structural pockets (accessible), interior cavities (inaccessible), shape (alpha complex and triangulation), area and volume (solvent and molecular accessible surface) of each pockets and cavities of proteins. CASTp could be used to measure the number, area, circumference of mouth openings of each pocket in solvent and molecular accessible surface [15].

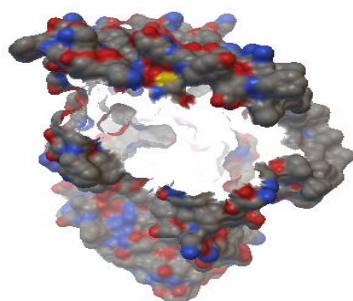




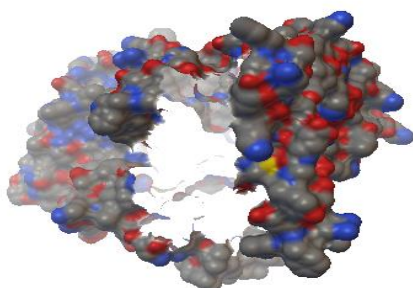
Active site of 1HT8



Active Site of 1PGF



Active Site of 3MQE



Active Site of Valdecoxib

### Analyzing the Docking Results

The search for the best ways is to fit ligand molecules into structure, using Autodock 4.2.1 resulted in docking files that contained detailed records of docking. The obtained log files were read in ADT (Auto Dock Tool) to analyze the results of docking. The similarity of docked structures was measured by computing the root mean square deviation (RMSD) between the coordinates of the atoms and creating clustering of the conformations based on the RMSD values. The lowest binding energy conformation in all cluster were considered as the most favourable docking pose. Binding energies that are reported represent the sum of the total intermolecular energy, total internal energy and torsional free energy minus the energy of the unbound system. The top ligands were selected among the 17 based on the energy score after virtual screening Table, 2,3,4,5 of result section.

**Table 1: List of the Ligands Retrieved from the Drug bank**

Ligand	Chemical formula	Molecular wgt.(avg)
Naproxen.	C <sub>14</sub> H <sub>14</sub> O <sub>3</sub>	230.2592
Etoricoxib.	C <sub>18</sub> H <sub>15</sub> ClN <sub>2</sub> O <sub>2</sub> S	258.842
Flurbiprofen.	C <sub>15</sub> H <sub>13</sub> FO <sub>2</sub>	244.2609
Ibuprofen	C <sub>13</sub> H <sub>18</sub> O <sub>2</sub>	206.2808
Indomethacin.	C <sub>19</sub> H <sub>16</sub> ClNO <sub>4</sub>	357.788
Ketoprofen.	C <sub>16</sub> H <sub>14</sub> O <sub>3</sub>	254.806
Piroxicam.	C <sub>15</sub> H <sub>13</sub> N <sub>3</sub> O <sub>4</sub> S	331.346
Diclofinac.	C <sub>12</sub> H <sub>11</sub> Cl <sub>2</sub> NO <sub>2</sub>	296.149
Ketorolac.	C <sub>15</sub> H <sub>13</sub> NO <sub>3</sub>	255.2686
Tolmetin	C <sub>15</sub> H <sub>15</sub> NO <sub>3</sub>	257.2845
Tenoxicam.	C <sub>13</sub> H <sub>11</sub> N <sub>3</sub> O <sub>4</sub> S <sub>2</sub>	337.374
Valdecoxib.	C <sub>16</sub> H <sub>14</sub> N <sub>2</sub> O <sub>3</sub> S	314.359
Meloxicam.	C <sub>14</sub> H <sub>13</sub> N <sub>3</sub> O <sub>4</sub> S <sub>2</sub>	351.401
Phenylbutazone.	C <sub>19</sub> H <sub>20</sub> N <sub>2</sub> O <sub>2</sub>	308.3743
Rofecoxib.	C <sub>17</sub> H <sub>14</sub> O <sub>4</sub> S	314.356
Sulindac	C <sub>20</sub> H <sub>17</sub> FO <sub>3</sub> S	356.411
Celecoxib.	C <sub>17</sub> H <sub>14</sub> F <sub>3</sub> O <sub>2</sub> S	381.3752

### Results and Discussion

We have successfully carried out docking for 17 structurally diverse COX-2 inhibitors. The obtained ADME score was correlated with the biological activities. Some false positives and false negatives were observed but considering the limitations of the available docking program, the results are encouraging. The detailed analysis of the resulted COX-1&COX-2 - ligand complexes may improve our knowledge in understanding the binding interactions in detail. Thus this study will be useful for the design of novel COX-2 inhibitors based on docking and the resulted bioactive conformations of ligands and the results obtained from

the Autodock of molecular docking and on the basis of binding energy scores we can suggest that tenoxicam and valdecoxib are the best fit ligand combinations which binds selectively with COX-2. This study will provide a platform for the further research and developments of drugs which can selectively suppress COX-2 and would not have any further side effects which were caused earlier due to the inhibition of COX-1. These drugs will surely help a lot in ailing diseases and genetical disorders like colon cancer and various kinds of arthritis. Agents that inhibit COX-2 while sparing COX-1 represent a new attractive therapeutic development and could represent a major advance in the treatment of arthritis and various diseases. The docking model for the substituted tenoxicam and valdecoxib derivatives with the COX-2 receptor has been developed in this project. To the best of literature survey, this is the first report of the

molecular modeling studies of these molecules with the COX-2 receptor. The docking simulation suggested that the modifications in the series that results in better binding potential. The Vander-walls, hydrophobic and charge interactions are responsible for forming the stable compound of the ligands with ligands with receptor. From the **Table.2,3,4,5 (Results)** ligands tenoxicam and valdecoxib do possess minimum dock score i.e. minimum binding energy in kilo joules per mole i.e. these molecule have more affinity for active site of COX-2 enzymes. Clearly, molecules with ester of bulky acids having less affinity for the receptor. Whereas molecules which possesses alcoholic with less bulky function 38-44 are said to have more affinity for COX-2 and can be used as analgesic and anti-inflammatory agents after synthesis.

### Descriptions

#### Sequences producing significant alignments:

Select: All None Selected: 1

Alignments Download Graphics Multiple alignment

Description	Max score	Total score	Query cover	E value	Ident	Accession
3MQE:BIPDBID CHAIN SEQUENCE	787	787	96%	0.0	65%	5059

Download Graphics

3MQE:BIPDBID|CHAIN|SEQUENCE

Sequence ID: lc|5059 Length: 587 Number of Matches: 1

Range 1: 1 to 555 Graphics

Next Match Previous Match

Score	Expect	Method	Identities	Positives	Gaps
787 bits(20333)	0.0	Compositional matrix adjust.	358/555(65%)	449/555(80%)	1/555(0%)
Query 9	VNFCVYPCQHQGICVFRGLRVQCDCTRTGVSQPCNTIPEINTWLRTLRPSSEFIHFL				68
Sbjct 1	NPCC PCQ+G C+ G D+Y+CDCTRTG+ G NCT PE T ++ L+P+P+ +H++				60
Query 69	LHGRWLNDFWAT-FIROTLMLVLTIVSRNLIPSPPTYNIAHDYISNESFSNVSYTRI				127
Sbjct 61	LHFGQVNIWNIIFLASLIMKYLTSRSLIDSPPTNVHYGKSWERFNLSTYTRA				120
Query 128	LPSVPRDCTPMGTGKQKQLPDAEFLSRRLRRKFIDPQGTNLMAFFAQHFTHQFFK				187
Sbjct 121	LPV DCPITMG KG K+LDP++ + LLRR+FIIDPQGN+MFAFFAQHFTHQFFK				180
Query 188	ISGKMGPGFTKALGHGVLDLGHVGNLQVQLRFLKDGKLYQMLNGEVVPPSVEAPV				247
Sbjct 181	T K GPGFT+ LSHGVLDL HIYG+ L+RQ+LRLFKDGKLYQ++ GEVVPF+V++ V				240
Query 248	LMHYRGPQPQSMVQGEVTFGLPGLMLYATINLREHNRVCDLLKAEHPTWDEQLFQT				307
Sbjct 241	EMIVPPIHPENLQFAVQGEVTFGLVPLGMLMYATINLREHNRVCDLLKAEHPTWDEQLFQT				300
Query 308	ARLILIGETIKIVIEYVQQLSGYFLQKDFPELLFGAQFYQYRNRIAMEFNQLYHWHFLM				367
Sbjct 301	+RLILIGETIKIVIE+YVQ LSGY +LKFDELLF QFY+NRIA EFN LYHWHFL+				360
Query 368	PDSFRVGPQDYSEYQFLNLSMLVDYGVLEALVDAFSRQAGRIGGGRNIDHILHVAIVD				427
Sbjct 361	PD+P+ Q+YS++QFL+N S+L+++G+ V++F+RQ AGR+ GGRN+ + VA				420
Query 428	IKESRVLRLQPFNVRKRGWKFYTSFQELTGEKEMAAELIYSDIDALEFYPLGILLEY				487
Sbjct 421	I+SR ++ Q NEYRKRF +FYTSP+ELTGEKEMAAEL+ LY DID +F YP LL+EK				480
Query 488	CHPNSIFGESMIEMGAPFSLKGLLGNFICSPEYWKASTFGGEVGFNLVKTALMLVCLN				547
Sbjct 481	P++IFGE+M+E+GAPFSLKGL+GNFICS+YWK STFGGEVGF ++ IA+++ L+C N				540
Query 548	TKTCFYVSFHVDPDR				562
Sbjct 541	K CP+ SF+V DP+				555

### Results of BLASTp of COX-1(1HT8) AND COX-2(3MQE)

**Descriptions**

Sequences producing significant alignments:

Select: [All](#) [None](#) Selected: 0

[Alignments](#) [Download](#) [Graphics](#) [Multiple alignment](#)

	Description	Max score	Total score	Query cover	E value	Ident	Accession
<a href="#">3NTG.B PDBID CHAIN SEQUENCE</a>		781	781	95%	0.0	65%	40699

**3NTG.B|PDBID|CHAIN|SEQUENCE**  
Sequence ID: ICI40699 Length: 552 Number of Matches: 1

Range 1: 1 to 551 [Graphics](#) [Next Match](#) [Previous Match](#)

Score	Expect	Method	Identities	Positives	Gaps
781 bits(2017)	0.0	Compositional matrix adjust.	356/551(65%)	446/551(80%)	1/551(0%)

Query 9 VNPCCYPCQHQICVRFGLRQDCDCTRTGSGPHCTIPEIWTNLRTLRSPSPFIHFL 68  
Sbjct 1 ANPCCSNPCQHQHGECHSTGDFQYKDCDCTRTGSGPHCTIPEIWTNLRTLRSPSPFIHFL 60

Query 69 LTHGRWLNDFVNIAT-FIRDTLMRLVLTVRNLIPTPTVNIARDVISHESFNSVSYTRI 127  
LTH + +N+ VN F+R +M+ VLT RS LI SPPTYN+ + Y SNE+FSN+SYTR  
Sbjct 61 LTHFGVNNIVNNIPFLASLIMQVLTSSYLDSPTPTVNVHYGKSNFASNLSSYTRA 120

Query 128 LPSVPRDCPTPMGTGKQQLPDAEFLSRRLRRKTFIDPQGNLMFAFFAQHTHQFFK 187  
LP V DCPITMG KG K+LPD++ + + LLRR+FIIDPQGN+MFAFFAQHTHQFFK  
Sbjct 121 LFPVADDCPTPMGTGKQQLPDAEFLSRRLRRKTFIDPQGNLMFAFFAQHTHQFFK 180

Query 188 TSGQMPGFTKALGHGVLDGHIYGNLQVQLRLFKDGKLYQMLNDEVPSVPEAPV 247  
T K GGGFT+ LGHGVLD HIY+ L+RQ+LRLFKDGKLYQ+ GEVPS+V+ V  
Sbjct 181 TSHRGPGFTKALGHGVLDGHIYGNLQVQLRLFKDGKLYQMLNDEVPSVPEAPV 240

Query 248 LMHVFGIPGQMAVGEVFTGLLEGLMLVATINLREHNRVCDLLKAEPTNGDEQLFQT 307  
M VP TP Q AVGEVFTGL+GLM+VATINLREHNRVCDLLK AEPTNGDEQLFQT  
Sbjct 241 EMIVFPHIPEMLQFAVGEVFTGLLEGLMLVATINLREHNRVCDLLKAEPTNGDEQLFQT 300

Query 308 ARLILIGETIKIVIEEYVQQLSGVFLQLWFDPELLFQAFQVNRRIAMEFNQLVHNNPLM 367  
+RLILIGETIKIVIE+YVQ LSGY +LKFDELLF QFQV+NRRI EFN LYHNNPL+  
Sbjct 301 SRLILIGETIKIVIEDYVQQLSGVFLQLWFDPELLFQAFQVNRRIASEFNLYHNNPL 360

Query 368 PDSFRVGPQDYSVEQFLNLSMLVDYGVYALVDFAFRQAGRIAGGRNIDHILRVAVDV 427  
PD+F + Q+VS++QFL+N S+L+++G+ V++F+RQ AGR+ GGRN+ + VA  
Sbjct 361 FDTFNIEDQYSVFWQLYNNILLNGLTQFVESFTRQIAGRVAGGRNVPIAVQAVAKAS 420

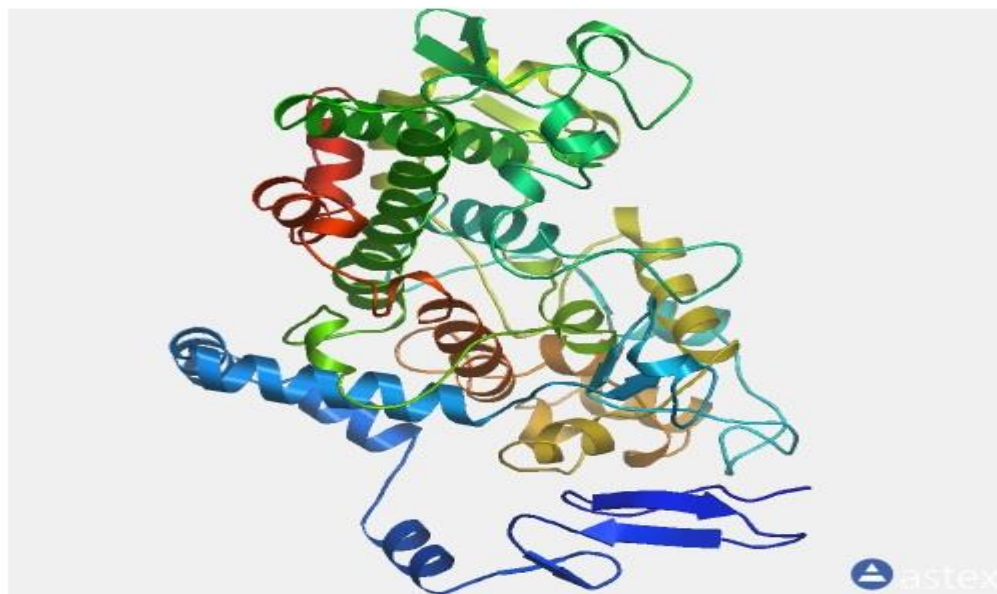
Query 428 IKESRVLRLQPFNEVRRFGMKPFTSFQELTGKEMAAEELLYGDIIDALEFYGLLLEK 487  
I +SR ++ Q NEYRRRF +KFTTSF+ELTGKEMAAEL+ LY DID +E YF LL+EK  
Sbjct 421 IQQSRVMQVSLNIEYRRFSLKFTTSFELTGKEMAAELKALYSDIVMEVYALIVEK 480

Query 488 CHPNSTFGESMIEMGAFFSLKGLLGNFICSPFYKASITFGGEVGNLVKITALKLVCLN 547  
P+IYGS+M+M+GAPFSLKGL+GNFICSP+VWV STFGGEVGF ++ TA+++ L+C N  
Sbjct 481 FRPDATFGEIMVELGAPFSLKGLLGNFICSPFYKWPSTFGGEVGFKINTASIGSLICNN 540

Query 548 TKTCFVVSFHV 558  
K CP+ SF+V  
Sbjct 541 VKGCFTSFHV 551

**Related Information**

# Results of BLASTp of COX-1(1HT8)AND COX-2(3NTG)



Structure of COX-2 from Swiss model server





Workunit: P000002  
Title: cox



#### Model Details: Batch.1

model pic.



Target:  
modelled residue range: 18 to 568  
based on template 3nt1B (1.73 Å)  
Sequence Identity [%]: 88.203  
Evaluate: 0

#### Alignment

TARGET	18	ANPCCSHP	CQNRGVCMSV	GFDQYKCDCT	RIGFYGENCS	TPEFLTRIKL
3nt1B	33	anpccsnp	cqnrgcsmat	gfdqykcdct	rigfygenct	tpelftrikl
TARGET		hh	sssss	sssss	sss	ssssssssss
3nt1B		hh	sssss	sssss	sss	ssssssssss
TARGET	66	FLKPTPNTVH	YILTHFKGFW	NVNNIPFLR	NAIMSYVLTS	RSHLIDSPPT
3nt1B	81	llkptptntvh	yilthfkgvfw	nivnnipflr	alimkyvlts	rsylidsappt
TARGET		hh	ssss	ss	ssss	ssssssssss
3nt1B		hh	ssss	ss	ssss	ssssssssss
TARGET	116	YNADYGYKSW	EAFSNLSYYT	RALPFVPDDC	PTPLGVKGKK	QLPDSNEIVG
3nt1B	130	yvnadygyksw	eafsnlsyvt	ralpvpddc	ptpvgvkgkn	qlpdskeyle
TARGET		h	ssss	sss		ssssss
3nt1B		h	ssss	sss		ssssss
TARGET	166	KLLLRKFIP	DPQGSNMFA	FFAQHFTHQF	EKIDHKRGPA	FTNGLGHGVD
3nt1B	180	kvllrrefip	dpqgsnmfa	ffaghftthgf	fktdhkrgpg	ftnglghgvd
TARGET		h	ss	ss	ssssss	ssss
3nt1B		h	ss	ss	ssssss	ssss

TARGET	216	LNHIYGETLA	RQRKLRLFKD	GKMKYQIIDG	EMYPTVKDT	QAEMIYPPOV
3nt1B	230	lnhiygetld	rqhklrlfkd	gklkyqvigg	exypntvykdt	qvemiypphi

TARGET		hhhh	hh	hhhhh	sss	sss	ssshhh
3nt1B		hhhh	hh	hhhhh	sss	sss	ssshhh

TARGET	266	PEHLRFAVGQ	EVEGLVPGLM	MYATIWLREH	NRVCDVLKQE	HPENGDEQLF
3nt1B	280	penlgfavvg	evfglvpglm	myatiwlreh	nrvcdilke	hpewgdeqlf

TARGET		sss	hhh	hhh	hhhhhhhhh	hhhhhhhhh	hhhhh
3nt1B		sss		hhh	hhhhhhhhh	hhhhhhhhh	hhhhh

TARGET	316	QTSRLILIGE	TIKIVIEDYV	QHLSGYHFKL	KFDPELLFNK	QFQYQNRIAA
3nt1B	330	qtarlilige	tikiviedyv	ghlagyhfkf	kfdpellfng	qfaygnrias

TARGET		hhhhhhhhh	hhhhhhhhh	hhh	hhh	h
3nt1B		hhhhhhhhh	hhhhhhhhh	hhh	hhh	h

TARGET	366	EFNTLYHWHF	LLPDTFQIHD	QKYNYYQFIY	NNSILLEHGI	TQFVESFTRQ
3nt1B	380	efntlyhwhf	llpdtfnied	qevafkqfly	nnaillehgl	tqfvesftrq

TARGET		hhhhh	sss	sss	hhhh	hhhhhhhhh
3nt1B		hhhhh	sss	sss	hhhh	hhhhhhhhh

TARGET	416	IAGRVAGGRN	VPPAVQKVSQ	ASIDQSRQMK	YQSFNEYRKR	FMLKPYESFE
3nt1B	430	iagrvaggrn	vpiavqavak	asidqarem	vgalneyrkr	falkpytsfe



TARGET		aaaa	bbbb	aaaaaaaa	bbbbbb	hh
3nt1B		aaaa	bbbb	aaaaaaaa	bbbbbb	hh
TARGET	466	ELTGEKEMSA	ELEALYGDID	AVELYPALLV	EKPRPDAIFG	ETMVEVGAPF
3nt1B	480	eltgekemaa	elkalyadid	vmelypally	ekprpdaifg	etmvelgapf
TARGET		hhh	bbbb	bbbbbb	bbbbbb	bbbbbbbbbb
3nt1B		hhh	bbbb	bbbbbb	bbbbbb	bbbbbbbbbb
TARGET	516	SLKGLMGNVI	CSPAYWKPST	FGGEVGFQII	NTASIQSLIC	NNVKGCPFTS
3nt1B	530	slkglmgmpi	cspaywkpat	fggevqfkii	ntasigalic	nnvkgcpts
TARGET		bbbbbb		aaaaaa	h	bbbbbb
3nt1B		bbbbbb		aaaaaa	h	bbbbbb
TARGET	566	FSV				
3nt1B	580	fsv				
TARGET		aaa				
3nt1B		aaa				

## Docking and ADME

Table 2: Binding energy and other parameters of the ligands with 1HT8

Ligand	Binding Energy	RMSD	Inhibition Constant	H Bonds
Celecoxib.	-9.28	0	257.36	4
Tenoxicam	-12.29	0	988.24	4

Table 3: Binding energy and other parameters of the ligands with 3MQE

Ligand	Binding Energy	RMSD	Inhibition Constant	H Bonds
Tenoxicam	-12.37	0	856.71	3
Valdecoxib	-12.75	0	452.7	9

Table 4: Binding energy and other parameters of the ligands with 3NTG

Ligand	Binding Energy	RMSD	Inhibition Constant	H Bonds
Ketoprofen	-9.22	0	173.52	3
Telometin	-9.13	0	204.19	3
Valdecoxib	-13.4	0	150.74	7

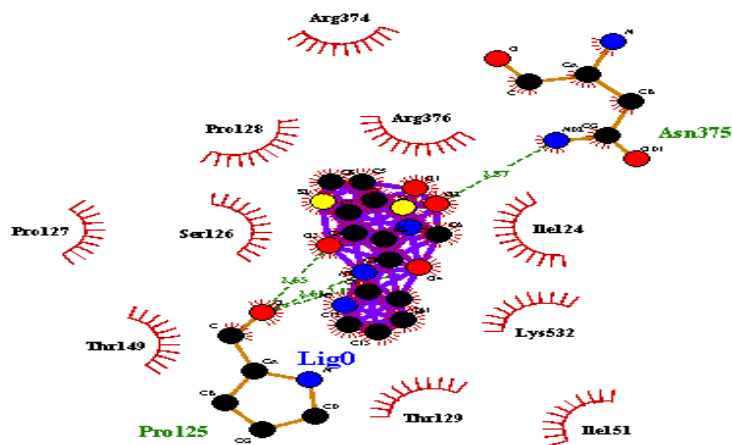
Table 5: Binding energy and other parameters of the ligands with 1PGF

Ligand	Binding Energy	RMSD	Inhibition Constant	H Bonds
Piroxicam	-16.59	0	690.85	4
Tenoxicam	-13.65	0	98.64	3

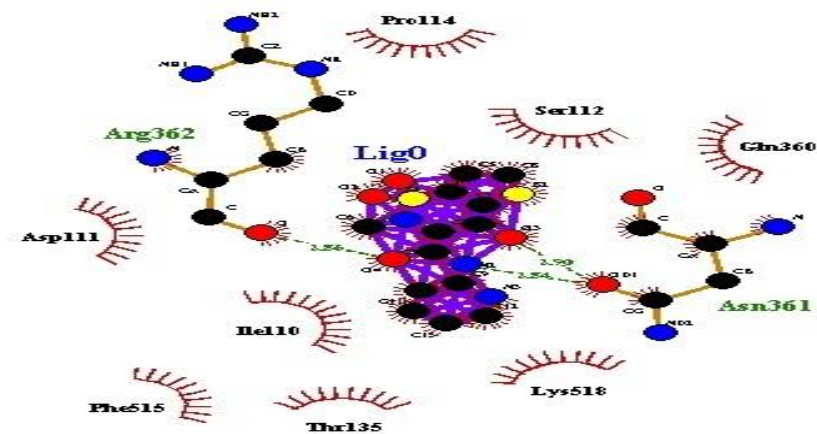
Table 6: Drug Likelihood Prediction (ADME)

Ligand	Intestinal absorption	Blood brain barrier	Caco-2 permeable	Ames Test
Tenoxicam	+0.9955	-0.9455	+0.8867	Negative
Piroxicam	+0.9898	-0.9659	+0.8867	Negative
Valdecoxib	+1	+0.9386	+0.5	Negative

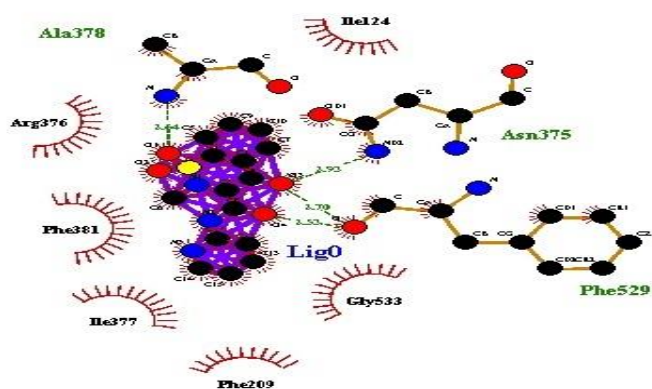
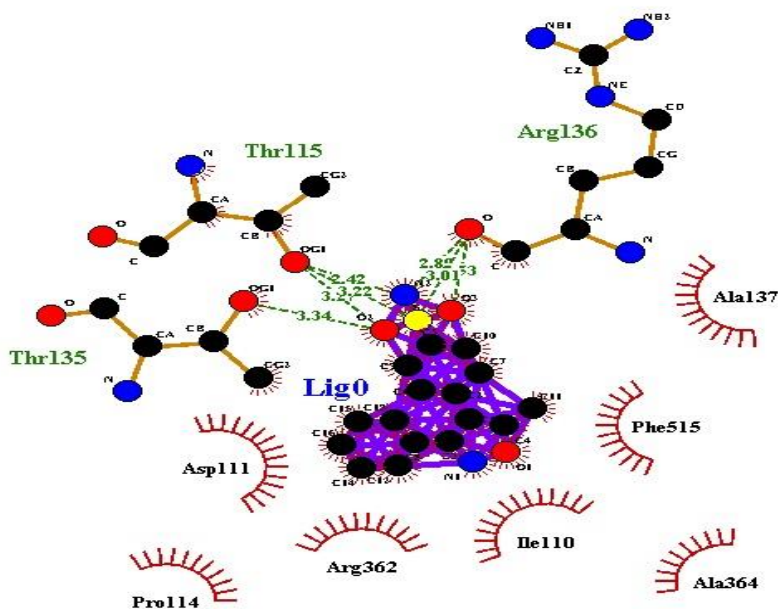
Ligplot



Hydrogen Bond Between 1HT8 and Tenoxicam



Hydrogen Between 3MQE and Tenoxicam





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