the world's leading publisher of Open Access books Built by scientists, for scientists

4,800

Open access books available

122,000

International authors and editors

135M

Downloads

154

TOP 1%

Our authors are among the

most cited scientists

12.2%

Contributors from top 500 universities



WEB OF SCIENCE™

Selection of our books indexed in the Book Citation Index in Web of Science™ Core Collection (BKCI)

Interested in publishing with us? Contact book.department@intechopen.com

Numbers displayed above are based on latest data collected.

For more information visit www.intechopen.com





Binding Mode Identification for 7-keto-8-Aminopelargonic Acid Synthase (AtKAPAS) Inhibitors

Nam Sook Kang, Jung-Sup Choi and In-Taek Hwang

Additional information is available at the end of the chapter

http://dx.doi.org/10.5772/60966

Abstract

In this study, we determined the 3D structure of Arabidopsis thaliana KAPAS by homology modeling. We then investigated the binding mode of compounds obtained from the in-house library using computational docking methods. From the flexible docking study, we achieved high dock scores for the active compounds denoted in this study as compound 3 and compound 4. Thus, we highlight the flexibility of specific residues, Lys 312 and Phe 172, when used in active sites.

Keywords: KAPAS, Herbicides, Homology modeling, Protein docking

1. Introduction

Agricultural research efforts for discovery of herbicides acting on new target sites are increasing due to demand from farmers and multinational companies in many countries. However, new modes of action have not been turned to commercial success for the past 10 years. We have recently reported 7-keto-8-aminopelargonic acid synthase (KAPAS, also known as 8-amino-7-oxononanoate synthase, ANONS) and have suggested the potential KAPAS inhibitor triphenyltin. Research on herbicides has advanced during the past 50 years to the point that herbicides can now protect crops and improve the quality and quantity of agricultural products. However, the successful development of herbicides has decreased recently owing to new environmental regulations and lack of discovery. To overcome this problem, there is an urgent need for new herbicidal targets and new techniques [1].



While the traditional approach to discover lead compounds heavily depends on serendipity given the poor understanding of biological modes of actions, the structure-based approach utilizes the structure of appropriate target proteins, which have well-known binding sites for possible rational designs. The introduction of new herbicides with either a new mode of action or of a novel chemical class has lingered. However, discovering a chemical structure that could enter the pest, be transported within it, inhibit a key target, get away from detoxification, and also be modified to allow it to fulfill increasing regulatory criteria with respect to environmental compatibility has been required. The structure-based approach uses only appropriate target proteins instead of the entire plant for *in vivo* testing [2]. In order to perform a structure-based assay, it is necessary to determine a potent target and to have a thorough understanding of the mechanism of action of the target.

Several enzymes in plants are known to be essential enzymes, meaning that they are crucial for the plant's survival. Disrupting a single essential enzyme leads to severe disorder of metabolic processes in the plant, ultimately causing a lethal outcome. The enzyme 7-keto-8-aminopelargonic acid synthase from *Arabidopsis thaliana* plants (*At*KAPAS), introduced in this research, and is a new potent herbicide target, which is involved in the early steps of the creation of the biotin biosynthesis pathway. *At*KAPAS as a pyridoxal 5'-phosphate-dependent enzyme catalyzes the decarboxylative condensation of L-alanine with pimeloyl-CoA in a stereospecific manner to form KAPA, coenzyme A, and carbon dioxide in the first committed step of biotin biosynthesis. Inhibiting *At*KAPAS leads to significant changes in the phenotype, such as growth inhibition, severe growth retardation, and the creation of lethal phenotypes [3].

Although the physiological systems of humans and plants are different in various ways, the misuse of agricultural chemicals can be extremely harmful to humans. Therefore, use of herbicides must follow strict toxicity regulations that are in place to prevent harm to humans and other life. As mentioned above, the novel herbicidal target 7-keto-8-aminopelargonic acid synthase functions in the initial steps of the biosynthetic pathways of biotin (vitamin H) in plants and microorganisms. Because biosynthetic steps of biotin exist only in plants, we expect that the inhibition of the potent target AtKAPAS will not affect the human metabolic system [1, 3]. A few publications have also reported beneficial effects of AtKAPAS as a potential herbicidal target. Hwang et al. described the possibility of AtKAPAS as a potential herbicide target enzyme and chemical validation of triphenyltin acetate as a lead compound for the AtKAPAS inhibition in vitro and in vivo [1]. They also suggested AtKAPAS can be a useful target for the rational design of inhibitors in the hope of developing new herbicides.

In this chapter, we aim to obtain potential AtKAPAS inhibitors using the knowledge-based computational informatics method in this research. We described the 3D-structure of AtKAPAS via theoretical method and the binding mode for AtKAPAS and its inhibitors obtained from $in\ vitro$ assay for in-house compounds.

2. Building a homology model of AtKAPAS

To apply the structure-based drug design (SBDD) method using current knowledge of protein and drug interactions, a three-dimensional protein structure is necessary [4]. Because the

known protein crystal structural information of 7-keto-8-aminopelargonic acid synthase from an experiment was absent, a homology model of *At*KAPAS was constructed from its amino acid sequence (Table 1).

60	50	40	30	20	10
evfdglcqwd	ksranggdgy	Msrqneeeiv	qilrslrpic	eeavnvlesr	madhswdktv
120	110	100	90	80	70
dylglsshpt	fkklllfsgn	dalaecrkgr	psngeeifsg	ptfqkwlhde	rtsvevsvsi
180	170	160	150	140	130
gfaanmaamv	kkedclvcpt	Llesslaqlk	licgyttyhr	eygmgpkgsa	isnaaanavk
240	230	220	210	200	190
yrhcdmyhln	erqgnvevfv	Asiidgvrla	vaifsdalnh	asgkplknek	aigsvaslla
300	290	280	270	260	250
fvcgengggv	llviddahgt	lsqlrkkygf	mdgdfapmee	kvvvtdslfs	sllsnckmkr
360	350	340	330	320	310
ipvpmaaaay	rgrsfifsta	skkwkqliqs	agchggfiac	dlcvgtlska	aeefnceadv
420	410	400	390	380	370
ryllksgfhv	gnqekalkas	Isspiislvv	kefkelsgvd	wrrkaiwerv	aavvvarkei
480	470	460	450	440	430
flfpkl	fdntathips	Litalsscld	aahttedvkk	nscrlrvtls	mairpptvpp

Table 1. Single letter amino acid sequence for *At*KAPAS.

To obtain the 3D-structure of AtKAPAS, a hierarchical protein structure modeling approach was adopted based on secondary-structure enhanced profile-profile threading alignment (PPA) and iterative implementation by the threading assembly refinement (TASSER) program [5]. We obtained five candidate models for the three-dimensional structure of AtKAPAS and then performed molecular dynamics simulations with the use of the CHARMM (Chemistry at HARvard Macromolecular Mechanics) force field (version 27.0) [6] with default parameters interfaced with Accelrys Discovery Studio 3.1. To identify binding sites, we collected the crystal structures of homologous proteins with AtKAPAS as templates from the RCSB Protein Data Bank (PDB) based on three different categories: structural similarity, binding site similarity and functional similarity. Ten different protein crystal structures originating from different species but structurally similar to AtKAPAS were selected. The binding site sequence and conformation of the target protein are particularly important here compared to other sites. Therefore, we found 10 different PDB hits which are similar in terms of their binding site to AtKAPAS. A comparison between each PDB hit and the AtKAPAS biding site sequence was done. The results are denoted using root mean square deviation (RMSD) values ranging from 1.83 to 3.51, as shown in Table 2.

PDB code	Sequence	RMSD
1FC4	36.3	1.83
1DJE	39.5	1.66
2BWO	39.6	1.59
2WKA	32.2	2.42
3KKI	30.3	2.64
3DXV	16.6	3.26
3DXW	16.8	3.25
2OAT	15.7	3.38
1GBN	15.6	3.43
1MLY	15.5	3.51

Table 2. Sequence similarity and RMSD between 10 reference PDB proteins.

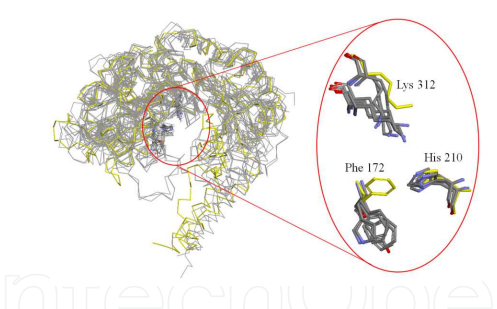


Figure 1. The superimposed 10 known enzymes (PDB code: 1FC4, 1DJE, 2WKA, 2BWO, 3KKI, 3DXV, 3DXW, 2OAT, 1GBN and 1MLY). The yellow-colored structure represents our homology model, and the overlapping predominantly important residues in active site are elaborated on the right side of the entire protein alignment.

To identify the binding mode of AtKAPAS, 10 different enzymes (PDB code; 1FC4 [7], 1DJE [8], 2WKA [9], 2BWO [10], 3KKI [11], 3DXV [12], 3DXW [12], 2OAT [13], 1GBN [14] and 1MLY [15]) with different functions originating from different species were superimposed (Fig. 1). Consequently, we found that several residues were crucial for protein-ligand interactions. His 210 mainly forms a π - π interaction or a π -cation interaction with the ligands, and all of the reference proteins contain histidine residue at a position homologous to the AtKAPAS. Adjacent to His 210, Phe 172 forms a π – π interaction with compounds that have an aromatic ring moiety as well. 50% of proteins have phenyl residue at a similar position; however, its

conformation was considerably different. As shown in Fig. 1, conformation of Phe 172 residue of our homology model, shown in yellow, is uniquely folded toward the active site cavity as opposed to the phenyl residues of the reference PDB. To confirm the effect of Phe 172, we undertook a flexible docking evaluation. Lys 312 also plays an important role as a hydrogen bond donor in the active site. All reference enzymes showed the lysine residue at an analogous location which led to π –cation interaction and hydrogen bonding interaction with its ligand. In addition, due to its comparatively free long aliphatic chain, lysine residue was very flexible. Lysine residues superimposed onto the analogous site verified its flexibility. By superimposing the binding sites of other similar proteins, we reached the conclusion that the residues His 210, Phe 172 and Lys 312 play major roles at the binding site. This research will therefore highlight the ligand binding site residues and its flexibility to search for potent hits.

3. Rigid docking

As a structure-based drug design, we used an automated docking method [16]. To dock the compounds as shown in Table 3 into the protein active site, we used the rigid docking method implemented in Discovery Studio 3.1 (Accelrys, Inc.), which adopts a Monte-Carlo algorithm to generate ligand conformations and docks the generated ligands into the active site using a shape-based filtering method. The rigid docking process consists of two main steps: defining a binding cavity and docking ligands onto the defined cavity [17]. The *At*KAPAS protein as obtained from the homology model needs to be prepared before the docking process is initiated. To prepare the protein, the CHARMM force field was assigned and the docking cavity was defined using the advanced define and edit binding site tool module. Ligands also should be prepared with low-energy conformations so as to be docked onto a 'clean' protein. The protocol 'Generate Conformations' in Discovery Studio 3.1 was used to obtain three-dimensional conformations of each ligand. In-house 17 ligands were generated by the protocol 'Generate Conformation' with the conformation method 'BEST', and the CHARMM force field was applied. For an interaction filter, Lys 312 was selected.

4. Flexible docking

Various docking methods are utilized by researchers. Each approach was developed by focusing on different aspects of docking. One of the factors determining the accuracy of docking is protein flexibility. Much emphasis has been placed on the conformational changes of protein binding sites, where different ligands form interactions. The Flexible Docking protocol of Discovery Studio 3.1 allows receptor flexibility during the docking of ligands [18-20]. To confirm the flexibility of the selected residues in the *At*KAPAS active site, three different sets of residues were defined as follows: flexible docking 1 (Lys 312), flexible docking 2 (Phe 172), and flexible docking 3 (Lys 312 and Phe 172). For the flexible docking 1 set, only the residue Lys 312 was assigned to move when flexible docking was underway. 17 preprocessed ligands were docked into the prepared *At*KAPAS homology model. The 'BEST'

conformation method was selected to generate three-dimensional ligands with stable energy levels, and other parameters were left with the default values. The residue Phe 172 was selected for the second flexible docking trial with the same parameters used with the flexible docking 1 set. To validate the effect of both residues, Lys 312 and Phe 172 were set as the flexible docking 3 group, and these residues were moved while flexible docking was underway.

Compounds	Structure	pIC ₅₀	Compounds	Structure	pIC ₅₀
1		5.48	10	CI	4.95
2		5.36	11		4.38
3		6.23	12		5.68
4		6.20	13		5.02
5	N—s C1 C1	4.25	14		5.36
6	N—S CI	4.27	15		5.97
7		4.39	16		5.90
8		4.33	17		5.36
9		4.95			

Table 3. The structures of the 17 compounds used and their biological activities.

5. *In Vitro* assay

Pimeloyl CoA was synthesized according to a previously described method [21]. AtKAPAS activity was determined according to the method described previously [22] using a linked assay by monitoring the increase in absorption of NADH at 340 nm using a microplate spectrophotometer (Benchmark Plus, Bio-rad, USA), thermostatically controlled at 30°C. A typical assay contained 20 mM potassium phosphate (pH 7.5), 1mM α -ketoglutarate, 0.25 mM thiamine pyrophosphate, 1 mM NAD⁺, 3 mM MgCl₂, 0.1 unit of α-ketoglutarate dehydrogenase, and 2 to 10 µg of AtKAPAS in a total volume of 200µL. L-Alanine and pimeloyl-Co A were added to give the desirable final concentrations. Prior to analysis, enzyme samples were dialyzed for 2 hours at 4°C against 20 mM potassium phosphate (pH 7.5) containing 100 µM PLP. The KAPAS concentration in all analyses was 10 µM in 20 mM potassium phosphate (pH 7.5) and the concentrations of each compound were 0.1 to 250 µM. Reference cuvettes contained all other compounds except inhibitor.

6. Results and discussion

Interesting results were obtained from the docking processes undertaken in this study. The rigid docking output scores of the in-house compounds are shown in Table 4. The most active compounds, in this case compound 3 and compound 4, obtained high dock scores of 104.21 and 105.47, respectively. Moreover, active ligands which have IC₅₀ (μ M) values 1.07 and 1.26 (compound 15 and compound 16, respectively) formed a stable docking pose (Fig. 2) with high dock scores 106.03 and 72.5, respectively. However, other active compounds, specifically compound 1, compound 9, and compound 10, showed rather low dock scores, as shown in Table 3.

Compound	Dock score	pIC ₅₀ a	Compound	Dock score	pIC ₅₀ a
1	40.40	5.48	10	45.79	4.95
2	67.64	5.36	11	49.91	4.38
3	104.21	6.23	12	63.13	5.68
4	105.47	6.20	13	52.84	5.02
5	55.73	4.25	14	52.99	5.36
6	42.30	4.27	15	106.03	5.97
7	50.67	4.39	16	72.51	5.90
8	51.67	4.33	17	52.31	5.36
9	58.72	4.95			

a: $pIC50 = -logIC_{50}$

Table 4. The result of rigid docking.

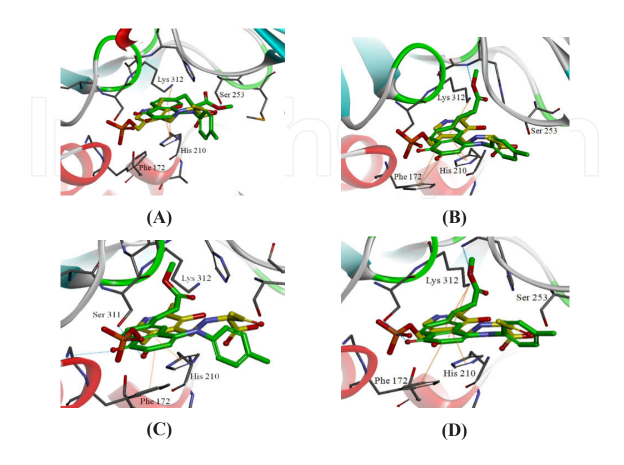


Figure 2. The docking pose from the process of rigid docking and flexible docking (A~D). The blue dashed line represents hydrogen bonding interaction and the orange dashed line represents the π -cation and π - π interaction. The docked ligand, compound 3, is colored with green and the crystal ligand of 1FC4 colored with yellow is overlapped as a reference compound. The docking pose from the process of rigid docking is shown in (A). From the process of flexible dock set 1, the docking pose of compound 3 was obtained as shown in (B). The docking pose shown in (C) achieved from the process of flexible docking set 2. In addition, the docking pose of compound 3 shown in (D) obtained from the process of flexible docking set 3.

To obtain a better docking result, we used a flexible docking strategy, as mentioned in the experimental section. In the result for flexible docking 1, which stipulated that the Lys 312 residue of the *At*KAPAS model is set to move, most of the compounds formed a stable conformation and formed interactions with the *At*KAPAS homology model (Table 5).

According to the docking result, the diverse conformation of Lys 312 directly affects the pose of the ligands and the related activity. The Lys 312 residue forms a hydrogen bond or undergoes the π -cation interaction mostly with the oxygen moiety of the ligand by flexibly moving through the protein cavity (Fig. 2). Interestingly, we obtained relatively high dock scores for compound 1, compound 9, and compound 10. This was unobtainable with the rigid docking process. The correlation coefficient between pIC50 and the dock score for the flexible docking 1 set was 0.72, as shown in Fig. 3.

Compound	Dock score	pIC ₅₀
1	74.00	5.48
2	83.14	5.36
3	134.03	6.23
4	125.27	6.20
5	53.40	4.25
6-1-	50.83	4.27
7	59.46	4.39
8	54.40	4.33
9	105.54	4.95
10	76.74	4.95
11	74.36	4.38
12	83.56	5.68
13	76.92	5.02
14	80.04	5.36
15	131.67	5.97
16	96.88	5.90
17	75.41	5.36

Table 5. The result of flexible docking set 1.

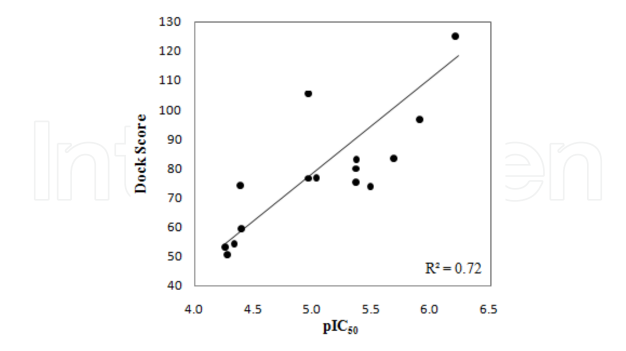


Figure 3. Correlation of \mbox{IC}_{50} and dock score for the flexible docking set 1.

By flexibly moving the residue Phe 172 while the flexible docking 2 process was underway, a better result than the flexible docking 1 process was obtained (Table 6). The flexibility of the Phe 172 residue has a significant effect on the ligand binding at the AtKAPAS active site. The phenyl ring is particularly important to form the π - π interaction with compounds. According to the docking result, high dock scores of 119.91 and 113.06 were the result with the most active compounds: compound 3 and compound 4. In addition, active compounds with IC $_{50}$ values lower than 1.3 μ M received dock scores higher than 90 with a stable conformation. Conversely, the inactive compounds of compound 5 through compound 8 as well as compound 11 obtained relatively low dock scores (Table 6).

Compound	Dock score	pIC ₅₀
1	75.60	5.48
2	85.84	5.36
3	119.05	6.23
4	113.06	6.20
5	48.11	4.25
6	44.80	4.27
7	53.49	4.39
8	49.60	4.33
9	72.97	4.95
10	74.43	4.95
11	57.66	4.38
12	85.24	5.68
13	79.99	5.02
14	67.73	5.36
15	113.29	5.97
16	90.84	5.90
17	65.49	5.36

Table 6. The result of flexible docking set 2.

The correlation coefficient between pIC50 and the dock score for the flexible docking 2 set was 0.85, as shown in Fig. 4.

The two specific *At*KAPAS residues, Lys 132 and Phe 172, can be moved mutually for the flexible docking 3 process. The result shows the clear discrepancy between an active compound and an inactive compound (Table 7). The flexible docking 3 results showed a higher correlation between pIC50 and the dock score compared to flexible docking 2, whereas it had a slightly lower R² value than the flexible docking 2 result (Fig. 5). Because the two residues were set to move at the same time, more diverse results could be obtained. As a result of several flexible docking processes, this study emphasizes the flexibility of several residues. Noticeably higher

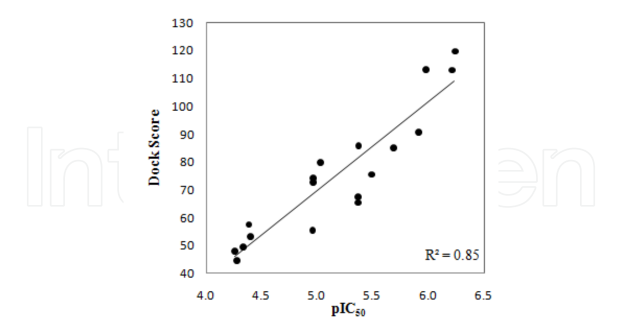


Figure 4. Correlation of IC_{50} and dock score for the flexible docking set 2.

dock scores were obtained from flexible docking as compared to rigid docking. The rigid Lys 312 residue mostly tends to form π –cation interaction with the aromatic moiety of compounds, allowing a certain amount of space for compounds during the rigid docking process. However, the flexible Lys 312 forms either the π –cation interaction or the hydrogen bonding interaction with hydrogen bond acceptors. The flexibility of the residue, including Lys 312 and Phe 172, allows more space for compounds, thus offering a better docking pose and dock scores.

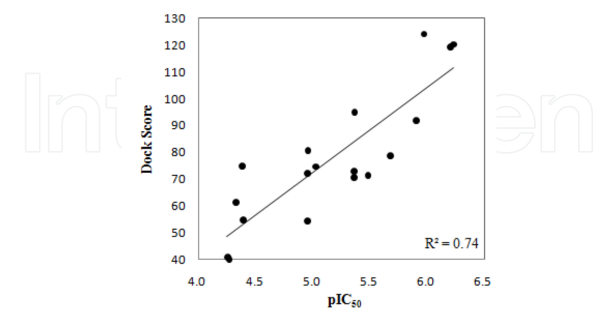


Figure 5. Correlation of IC₅₀ and dock Score for the flexible docking set 3.

Compound	Dock score	pIC ₅₀
1	71.47	5.48
2	95.12	5.36
3	120.48	6.23
4	119.46	6.20
5 5 7	40.92	4.25
6	40.03	4.27
7	54.90	4.39
8	61.54	4.33
9	80.87	4.95
10	72.13	4.95
11	75.00	4.38
12	78.83	5.68
13	74.82	5.02
14	73.04	5.36
15	124.28	5.97
16	91.92	5.90
17	70.70	5.36

Table 7. The result of flexible docking set 3.

7. Summary

In this study, we determined the 3D-structure of AtKAPAS by homology modeling. We then investigated the binding mode of our in-house library using computational docking methods. In the rigid docking of the in-house compounds as shown in Table 3, the most active compounds, in this case compound 3 and compound 4, obtained high dock scores of 104.21 and 105.47, respectively. However, some active compounds showed rather low dock scores. To obtain a better docking result, we used a flexible docking strategy. From the flexible docking study, we achieved high dock scores and stable binding conformations for the active compounds denoted in this study. Thus, we highlight the flexibility of specific residues, Lys 312 and Phe 172, when used in active sites. Furthermore, we are going to optimize compound 3 and compound 4 using this homology model for AtKAPAS. Following the initial identification of a lead chemical, intensive research and testing needs to be followed to optimize its structure to understand its action and provide data on its environmental compatibility.

Author details

Nam Sook Kang¹, Jung-Sup Choi² and In-Taek Hwang^{2,3*}

- *Address all correspondence to: ithwang@krict.re.kr
- 1 Graduate School of New Drug Discovery and Development, Chungnam National University, Deajeon, Republic of Korea
- 2 Korea Research Institute of Chemical Technology, Yusung, Daejon, Republic of Korea
- 3 Department of Green Chemistry and Environmental Biotechnology, University of Science & Technology, Yuseong-gu, Daejon, Republic of Korea

References

- [1] Hwang, I. T.; Choi, J. S.; Song, H. Y.; Cho, S. J.; Lim, H. K.; Park, N. J.; Lee, D. H. Validation of 7-keto-8-aminopelargonic acid synthase as a potential herbicide target with lead compound triphenyltin acetate, *Pestic. Biochem. Physiol.*, 97(2010), 24.
- [2] Hwang, I. T.; Lee, D. H.; Park, N. J. 7-Keto-8-aminopelagonic acid synthase as a potential herbicide target, *InTec.*, 471(2011), 22.
- [3] Choi, J. S.; Lim, H. K.; Seo, B. R.; Kim, J. S.; Choi, C. W.; Kim, Y. S.; Ryu, S. Y. Herbicidal activity and KAPAS inhibition of juglone with potential as natural herbicide, *Korean Soc. Weed Sci.*, 31(2011), 240.
- [4] Krieger, E.; Nabuurs, S. B.; Vriend, G. Homology modeling, *Struct. Bioinformatics*, 25(2003), 507.
- [5] Zhang, Y. Template-based modeling and free modeling by ITASSER in CASP7, *Proteins*, 69(2007), 108.
- [6] Brooks, B. R.; Bruccoleri, R. E.; Olafson, B. D.; States, D. J.; Swaminathan, S.; Karplus, M. CHARMM: a program for macro molecular energy, minimization, and dynamics calculations, J. Comp. Chem., 4(1983), 187.
- [7] Schmidt, A.; Sivaraman, J.; Li, Y.; Larocque, R.; Barbosa, J. A.; Smith, C.; Matte, A.; Schrag, J. D.; Cygler, M. Three-dimensional structure of 2-amino-3-ketobutyrate CoA ligase from *Escherichia coli* complexed with a PLP-substrate intermediate: inferred reaction mechanism, *Biochemistry*, 40(2001), 5151.
- [8] Webster, S. P.; Alexeev, D.; Campopiano, D. J.; Watt, R. M.; Alexeeva, M.; Sawyer, L.; Baxter, R. L. Mechanism of 8-amino-7-oxononanoate synthase: spectroscopic, kinetic, and crystallographic studies, *Biochemistry*, 39(2000), 516.

- [9] Jahan, N.; Potter, J. A.; Sheikh, M. A.; Botting, C. H.; Shirran, S. L.; Westwood, N. J.; Taylor, G. L. Insights into the biosynthesis of the *Vibrio cholerae* major auto inducer CAI-1 from the crystal structure of the PLP-dependent enzyme CqsA, *J. Mol. Biol.*, 3(2009), 763.
- [10] Astner, I.; Schulze, J. O.; van den Heuvel, J.; Jahn, D.; Schubert, W. D; Heinz, D. W. Crystal structure of 5-aminolevulinate synthase, the first enzyme of heme biosynthesis, and its link to XLSA in humans, *EMBO J.*, 18(2005), 3166
- [11] Kelly, R. C.; Bolitho, M. E.; Higgins, D. A.; Lu, W.; Ng, W. L.; Jeffrey, P. D.; Rabinowitz JD, Semmelhack MF, Hughson FM, Bassler BL. The *Vibrio cholerae* quorum-sensing autoinducer CAI-1: analysis of the biosynthetic enzyme CqsA, *Nat. Chem. Biol.*, 12(2009), 891.
- [12] Okazaki, S.; Suzuki, A.; Mizushima, T.; Kawano, T.; Komeda, H.; Asano, Y.; Yamane, T. The novel structure of a pyridoxal 5'-phosphate-dependent fold-type I racemase, alpha-amino-epsilon-caprolactam racemase from Achromobacter obae, *Biochemistry*, 5(2009), 941.
- [13] Storici, P.; Capitani, G.; Müller, R.; Schirmer, T.; Jansonius, J. N. Crystal structure of human ornithine aminotransferase complexed with the highly specific and potent inhibitor 5-fluoromethylornithine, *J. Mol. Biol.*, 1(1999), 297.
- [14] Shah, S. A.; Shen, B. W.; Brünger, A. T. Human ornithine aminotransferase complexed with L-canaline and gabaculine: structural basis for substrate recognition, *Structure*, 8(1997), 1067.
- [15] Sandmark, J.; Mann, S.; Marquet, A.; Schneider, G. Structural basis for the inhibition of the biosynthesis of biotin by the antibiotic amiclenomycin, *J. Biol. Chem.*, 227(2002), 43352.
- [16] Taylor, R. D.; Jewsbury, P. J.; Essex, J. W. A review of protein-small molecule docking methods, *J. Comput.-Aided Mol. Des.*, 16(2002), 151.
- [17] Montes, M.; Miteva, M. A.; Villoutreix, B. O. Structure-based virtual ligand screening with LigandFit: pose prediction and enrichment of compound collections. *Protein*, 15(2007), 712.
- [18] Welch, W.; Ruppert, J.; Jain, A. N. Hammerhead: fast, fully automated docking of flexible ligands to protein binding sites. *Chem. Biol.*, 3(1996), 449.
- [19] Koska, J.; Spassov, V. Z.; Maynard, A. J.; Yan, L.; Austin, N.; Flook, P. K.; Venkata-chalam, C. M. Fully automated molecular mechanics based induced fit protein-ligand docking method, J. Chem. Inf. Model, 48(2008), 1965.
- [20] Venkatachalam, C. M.; Jiang, X.; Oldfield, T.; Waldman, M. LigandFit: a novel method for the shape-directed rapid docking of ligands to protein active sites, *J. Mol. Graph. Model*, 21(2003), 289.

- [21] Ploux, O.; Marquet, A. The 8-amino-7-oxopelargonate synthase from *Bacillus sphaericus*. Purification and preliminary characterization of the cloned enzyme overproduced in *Escherichia coli*, *Biochem. J.*, 283(1992), 327.
- [22] Webster, S. P.; Alexeev, D.; Campopiano, D. J.; Watt, R. M.; Alexeeva, M.; Sawyer, L.; Baxter, R. L. Mechanism of 8-amino-7-oxononanoate synthase: spectroscopic, kinetic, and crystallographic studies, Biochemistry, 39(2000), 516.



IntechOpen

IntechOpen