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Design and *In silico* Screening of Combinatorial Library of New Herbicidal Analogs of Cycloalka[d]quinazoline-2,4dione-Benzoxazinones Inhibiting Protoporphyrinogen IX Oxidase

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Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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ABSTRACT

We virtually design here an important new weedkiller Cycloalka[d]quinazoline-2,4dione-Benzoxazinones (CQB), inhibitors of Protoporphyrinogen IX Oxidase (PPO). Based on computer-assisted combinatorial chemistry techniques, docking, 3D-QSAR and pharmacophore models we first enumerate, focus and *in silico* screen a virtual library of CQB analogs substituted at

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positions R₁, R₂ and R₃. By docking inhibitors into the target active site from the crystal structure (PDB ID: 1SEZ) of PPO in complex with a CQB ligand, 3D models of 29 PPO:CQBx complexes with known observed activity (K_i^{exp}) were prepared to establish a quantitative structure–activity (QSAR) model and linear correlation between relative Gibbs free energy (GFE) of receptor-ligand complex formation ($\Delta\Delta G_{com}$) and K_i^{exp}: pK_i^{exp} = -0.1664 $\star\Delta\Delta G_{com}$ + 8.306 (1); R² = 0.94. A 3D QSAR pharmacophore model (PH4) derived from the QSAR directed our effort to design novel CQB analogs. During the design, an initial virtual library of 118 CQB was focused down and PH4 screened to identify 28 promising novel analogs. Their K_i (K_i^{pre}) values were predicted by means of equation (1). The most active analog namely CQB22 display K_i^{pre} 22 times superior to that of the reported most active training set ligand 17i. Our survey proposes this compounds to the synthesis and to the assessment on herbicidal.

Keywords: CQB; PPO; docking; 3D-QSAR models; pharmacophore; in silico screening.

1. INTRODUCTION

Due to their detrimental impact on green plants and cultures survival bad herbs have drawn attention of research community in the design of herbicides addressing the United Nations sustainable development goal N°2 "Zero Hunger" [1]. "Photosynthesis is the process by which green plants and some other organisms use sunlight to synthesize nutrients from carbon dioxide and water. Photosynthesis in plants generally involves the green pigment chlorophyll and generates oxygen as a product. In the plant's life cycle, protoporphyrin IX is an important substrate involved in the biosynthesis of chlorophyll. It results from the oxidation of protoporphyrinogen IX" [2,3] and its inhibition is lethal for the plants making it one of the most important herbicidal targets.

Recently Da Wei and al. [4] suggest 36 compounds belonging to the class of the Cycloalka[d]quinazoline-2,4dione-Benzoxazino-

nes (CQB) with the best active namely 17i (Fig. 1) displaying a one digit nanomolar concentration range ($K_i = 6.7$ nM). The interactions between PPO and 17i displayed in the 2D diagram (Fig. 2) has served as starting structural information for analogs docking, QSAR modeling. Others weedkiller [5-7], inhibitors of Protoporphyrino-gen Oxidase (PPO), novel diphenyl ether IX derivatives have been proposed by Li-Xia Zhao et al. [8]. In this work we elaborate a 3D-QSAR pharmacophore model [9] derived from the QSAR model to serve as virtual library screening tool in the search of new herbicides. Despite the limitation that these new analogs did not undergo and biological evaluation. svnthesis this simulation-based design of novel analogs shortens the way in comparison with traditional intuition process. Agrochemical industry is the main target prospect of this computer-aided design of herbicidal compounds keeping in mind that the capacity of an economy to design molecule to satisfy its own need is more and more admitted as emergence indicator.



Fig. 1. Chemical structure of herbicidal agents: Cycloalka[d]quinazoline-2,4dione-Benzoxazinones core (17i)

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Fig. 2. PPO-17i – ligand interactions at active site depicted in 2D for the most active CQB

2. MATERIALS AND METHODS

2.1 Training and Validation Sets

Chemical structures and biological activities (K_i^{exp}) of training and validation sets of CQB derivatives ligands of PPO used in this study were taken from literature [3]. The potencies of these compounds cover a sufficiently broad range of half-maximal effective concentrations $(6.7 \le K_i^{exp} \le 2500 \text{ nM})$ to allow construction of a QSAR model. The training set (TS) containing 29 CQBs ligands and the validation set (VS) including 7 CQBs were taken from the ref [3].

2.2 Model Building

The structure of NtPPO (PDB ID: 1SEZ) [3] was downloaded (www.pdb.org) and prepared with Discovery Studio. Two monomers are present in the protein (A and B), we have used only A. No crystallographic water molecules were included in the model. The 3D structures of the cycloalka[d]quinazolinedione-benzoxazinones were constructed by Discovery Studio [10], on the basis of the crystal structure of 17a [3], and subsequently optimized with methods of algorithm *Smart Minimizer* before docking.

2.3 Molecular Mechanics

Modeling of ligands CQB and P-L complexes was carried out by molecular mechanics using

CHARMm force field [11] as described earlier [12].

2.4 QSAR Model

Training set of 29 CQBs derivatives compounds with known inhibitory potencies towards the PPO [3], were docked to the binding site model of the PPO receptor model using the CDOCKER docking procedure [9]. Complexation QSAR models were elaborated for training set and a linear correlation was established between the computed Gibbs free energies of binding (GFE: $\Delta\Delta G_{com}$) and observed enzyme inhibition constants (Kiexp) for each training set Were computed an QSAR models, which relate the K_i^{exp} to the computed scores, were prepared by regression analysis equation linear (1) $pK_i = f(Gibbs free energies)$ and GFE = bindingEnergy of Complex($\Delta G_{binding}$) - Entropic Term $(T\Delta S)$ + solvatation Energy (ΔG_{sol}) of Receptor, which correlates the computed. The predictive power of Eq. (1), which was then used as the target-specific scoring function for the in silico screening of the designed virtual library CQB analogs, was verified by applying it to a validation set of 7 similar CQB inhibitors with known Ki^{exp} values, which were not included into the training set. The ratio of predicted activities K_i^{pre} obtained from the regression equation (1) and observed Kiexp was used to evaluate the performance of the QSAR model.

Table 1. Training and validation set of PPO inhibitors for QSAR model



Training set	n	R ¹	R ²	R ³	Ki(µM)
8a	1	Н	Н	$CH_2C_6H_4(4-OCH_3)$	210
8b	1	Н	Н	Н	340
8c	1	Н	Н	CH ₃	270
8d	1	Н	Н	CH ₂ CH ₃	98
8e	1	Н	Н	CH ₂ CH ₂ CH ₃	69
8g	1	Н	Н	$CH_2CH(CH_3)_2$	170
8h	1	Н	Н	$CH_2CH=CH_2$	20
8j	1	Н	Н	CH ₂ C≡CSi(CH ₃) ₃	2500
8k	1	Н	Н	$CH_2CO_2CH_2CH_3$	48
9a	2	Н	Н	$CH_2C_6H_4(4-OCH_3)$	200
9c	2	Н	Н	CH ₃	160
9e	2	Н	Н	$CH_2CH_2CH_3$	44
9f	2	Н	Н	CH ₂ CH ₂ CH ₂ CH ₃	46
9g	2	Н	Н	$CH_2CH(CH_3)_2$	100
9h	2	Н	Н	$CH_2CH=CH_2$	11
9j	2	Н	Н	CH₂C≡CSi(CH ₃) ₃	260
9k	2	Н	Н	$CH_2CO_2CH_2CH_3$	370
91	2	Н	Н	CH₂CN	110
17a	2	F	F	$CH_2C_6H_4(4-OCH_3)$	200
17b	2	F	F	Н	480
17c	2	F	F	CH₃	140
17d	2	F	F	CH ₂ CH ₃	110
17e	2	F	F	CH ₂ CH ₂ CH ₃	58
17f	2	F	F	CH ₂ CH ₂ CH ₂ CH ₃	21
17g	2	F	F	$CH_2CH(CH_3)_2$	84
17h	2	F	F	$CH_2CH=CH_2$	9.8
17i	2	F	F	CH₂C≡CH	6.7
17j	2	F	F	CH₂C≡CSi(CH ₃) ₃	170
17m	2	F	F	CH ₂ CH ₂ OCH ₃	120
19	3	F	F	CH₂C≡CH	72
Validation set	n	R^1	R^2	R ³	Ki(µM)
9i	2	Н	Н	CH₂C≡CH	7.8
8i	1	Н	Н	CH₂C≡CH	14
17k	2	F	F	$CH_2CO_2CH_2CH_3$	14
9d	2	Н	Н	CH ₂ CH ₃	98
8f	1	Н	Н	CH ₂ CH ₂ CH ₂ CH ₃	120
171	2	F	F	CH ₂ CN	200
9b	2	Н	Н	Н	210

2.5 Pharmacophore Generation

"Bound conformations of inhibitors taken from the models of E-I complexes were used for constructing of 3D-QSAR pharmacophore (PH4) by using Catalyst HypoGen algorithm implemented in Discovery Studio" [9] as described earlier [13].

2.6 Virtual Library Generation

The virtual library generation was performed as described earlier [13].

2.7 Pharmacophore-Based Library Searching

"The pharmacophore model (PH4) described in Section 2.5 and derived from the bound conformations of CQBs at the active site of PPO served as a library searching tool" as described earlier [13].

3. RESULTS

3.1 Calculation of GFE and QSAR Model

The binding energy, Entropic Term and solvation Energy of the complex formation (E:I), equation (1) [14], was computed for the 29 complexes from docking and the ratio of predicted and observed inhibition constants (pKi^{pre}/pKi^{exp}) for the validation set of 7 CQBs (not included into the training set) are listed, show in Table 2. The QSAR model explained variation in the (pK^{exp}=experimental CQBs potencies $log_{10}(K_i^{exp}))$ [13] by correlating it with computed GFE $\Delta\Delta G_{com}$ through linear regression (Equation (1) [12] Table 2. In addition, significant correlation obtained in this QSAR relationship permitted to identify the CQBs active bound conformation at the PPO binding site and enabled definition of the PH4 pharmacophore. This correlation explained about 94% of the pKi exp data variation and underlined the role of the enthalpic contribution to the binding affinity of the ligand. Relatively high values of the regression coefficient R^2 , the leave-one-out cross-validated regression coef-ficient R²_{xv} and Fischer F-test of the correlation suggest strong relationship between the 3D model of ligand binding and the observed activation potencies of the CQBs.

The statistical data of the regression are presented in Table 3.

Therefore. structural information derived PPO:CQBx from the 3D models of complexes can be expected to lead to reliable PPO prediction of activation potencies for new CQBs analogs based on the QSAR model.

The statistical data confirmed validity of the correlation Equations (1) plotted on Fig. 3. The ratio pK^{pre}/pK^{exp} (the pK_i^{pre} values were estimated using correlation Equation (1), Table 3) calculated for the validation set documents the substantial predictive power of the complexation QSAR model from Table 2. Thus, the regression Equation (1) (Table 3) and computed $\Delta\Delta G_{com}$ GFEs can be used for K^{, pre} prediction of activator potencies against PPO for novel CQB analogs, provided they share the same binding mode as the training set.

3.2 QSAR Pharmacophore Model

"PPO activation 3D-QSAR pharmacophore was generated from the active conformation of 29 TS CQBx and evaluated by 7 VS CQB covering a large range of experimental activity (6.7-2500 nM) spanning more than two orders of magnitude. The generation process is divided into three main steps: (i) the constructive step, (ii) the subtractive and (iii) the optimization step step" [9].

During the constructive phase. 17i alone was retained as the lead (since only the activity of 17i fulfilled the threshold criterion. $K_i^{exp} \le 1.2 \times 8$ and used to generate the starting nM) features. In the subtractive phase, PH4 compounds for which $K_i^{exp} > 8 \times 103.5$ nM = 25.298 nM were considered inactive. Accordingly none of the training set CQBx was inactive and no starting PH4 features were removed. Finally, during the optimization phase, the score pharmacophoric hypotheses of the was improved. Hypotheses were scored according to errors in activity estimates from regression and complexity via a simulated annealing approach. At the end of the optimization, the top scoring 10 unique pharmacophore hypotheses were kept, all displaying five-point features. The cost values, correlation coefficients, root-mean square deviation (RMSD) values, the pharmacophore features, and the max-fit value of the top 10 ranked hypotheses (Hypo1- Hypo10) are listed in Table 4.

Training set	Ki (µM)	ΔG _{binding} (kcal/mol)	ΔG_{sol} (kcal/mol)	T∆S (kcal/mol)	ΔG_{compl} (kcal/mol)	ΔΔG _{compl} (kcal)	рКі
17i	0.0067	-58.79	-383.92	20.19	-462.90	0.00	8.17
17h	0.0098	-56.99	-383.42	20.20	-460.62	2.28	8.01
9h	0.011	-57.56	-382.94	19.98	-460.48	2.42	7.96
8h	0.02	-55.37	-383.61	20.36	-459.33	3.56	7.7
17f	0.021	-55.27	-382.60	20.36	-458.22	4.67	7.68
9k	0.037	-55.26	-382.90	20.24	-458.40	4.50	7.43
9e	0.044	-54.29	-382.90	20.02	-457.21	5.69	7.36
9f	0.046	-54.90	-382.56	20.15	-457.61	5.29	7.34
8k	0.048	-54.36	-382.26	20.14	-456.77	6.13	7.32
17e	0.058	-53.09	-382.99	20.22	-456.30	6.60	7.24
8e	0.069	-52.32	-383.16	19.91	-455.39	7.51	7.16
19	0.072	-53.94	-382.85	20.36	-457.15	5.75	7.14
17g	0.084	-52.66	-382.61	20.30	-455.57	7.33	7.08
8d	0.098	-51.19	-382.61	19.78	-453.57	9.33	7.01
9g	0.1	-51.88	-382.83	20.10	-454.81	8.09	7
91	0.11	-51.06	-383.04	19.99	-454.09	8.81	6.96
17d	0.11	-51.17	-382.89	20.12	-454.18	8.72	6.96
17m	0.12	-51.00	-383.88	20.36	-455.24	7.65	6.92
17c	0.14	-51.45	-383.55	20.04	-455.03	7.86	6.85
9c	0.16	-49.96	-383.33	19.80	-453.10	9.80	6.8
17j	0.17	-49.69	-382.60	20.01	-452.30	10.60	6.77
8g	0.17	-51.97	-381.34	20.01	-453.32	9.57	6.77
9a	0.2	-51.07	-382.35	20.51	-453.93	8.97	6.7
17a	0.2	-50.55	-382.12	20.69	-453.35	9.55	6.7
8a	0.21	-51.18	-382.11	20.41	-453.69	9.21	6.68
9j	0.26	-49.73	-381.67	20.50	-451.90	11.00	6.59
8c	0.27	-49.35	-383.32	19.68	-452.36	10.54	6.57
8b	0.34	-49.04	-383.76	19.57	-452.37	10.52	6.47
17b	0.48	-47.26	-384.00	19.94	-451.20	11.70	6.32
8j	2.5	-45.55	-384.04	19.10	-448.69	14.21	5.60

Table 2. Gibbs free energy (binding affinity) and its components for the training set of PPO ligands CQBs and validation set ligands

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Validation set	Ki	∆Gbinding (kcal/mol)	∆Gsol (kcal/mol)	T∆S (kcal/mol)	∆Gcompl (kcal/mol)	pKi_théo	pki_théo/pki_exp
9i	0.0078	-56.88	-383.52	19.97	-460.38	7.91	0.974
8i	0.014	-57.57	-383.00	19.86	-460.44	7.92	1.009
17k	0.014	-57.87	-382.75	20.54	-461.16	8.05	1.025
9d	0.098	-52.90	-382.98	19.90	-455.78	7.13	1.017
8f	0.12	-52.27	-382.93	20.01	-455.21	7.03	1.016
171	0.2	-50.02	-383.01	20.22	-453.26	6.70	1.000
9b	0.21	-49.94	-383.79	19.71	-453.44	6.73	1.008

Table 3. Analysis of computed binding affinities ΔΔGcom and experimental activity effective concentration of CQBs towards PPO [13]

Statistical Data of Linear Regression	
$pK_{i}^{exp} = -0.1664 \times \Delta\Delta G_{com} + 8.306$	(1)
Number of compound n	29
Squared correlation coefficient of regression R ²	0.94
LOO cross-validated squared correlation coef R^2_{xv}	0.94
Standard error of regression σ	0.132
Statistical significance of regression, Fisher F-test	449.36
Level of statistical significance α	>95%
Range of activities K _i ^{exp} [nM]	6.7-2500



Fig. 3. Plot for relative complexation Gibbs free energies of the PPO-CQBx complex formation $\Delta\Delta G_{com}$ [kcal.mol⁻¹] of the training set [15]. The validation set data points are shown in red color

Table 4. Parameters of 10 generated PH4 pharmacophoric hypotheses for PPO ligand after
Cat-Scramble validation procedure (49 scrambled runs for each hypothesis at the selected
level of confidence of 98%)

Hypothesis ^a	RMSD⁵	R²	Total costs ^c	Costs Difference ^d
Нуро 1	5.35	0.91	475.78	1994.95
Нуро 2	5.52	0.90	503.08	1967.65
Нуро З	5.68	0.89	529.64	1941.09
Hypo 4	5.76	0.89	544.18	1926.55
Hypo 5	5.79	0.89	549.03	1921.70
Нуро 6	5.79	0.89	549.23	1921.50
Нуро 7	5.81	0.89	551.80	1918.92
Нуро 8	5.86	0.88	559.82	1910.91
Нуро 9	5.87	0.88	562.11	1908.62
Hypo 10	6.08	0.87	597.58	1873.15

^aroot mean square deviation; ^b squared correlation coefficient; ^c overall cost parameter of the PH4 pharmacophore; ^d cost difference between Null cost and hypothesis total cost; ^e lowest cost from 49 scrambled runs at a selected level of confidence of 98%. The Fixed Cost = 43.48 with RMSD = 0, the Null Cost = 2470.73 with RMSD = 12.76 and the Configuration cost = 14.78

"The generated pharmacophore models were then assessed for their reliability based on the calculated cost parameters ranging from 475.78 (Hypo1) to 597.58 (Hypo10). The relatively small gap between the highest and lowest cost parameter corresponds well with the homogeneity of the generated hypotheses and consistency of the TS of CQBx. For this PH4 model, the fixed cost (43.48) is lower than the null cost (2470.73) by a difference $\Delta = 2427.25$. This difference is a major quality indicator of the PH4 predictability ($\Delta > 70$ corresponds to an excellent chance or a probability higher than 90% that the model represents a true correlation" [9]. "To be statistically significant, a hypothesis has to be as close as possible to the fixed cost and as far as possible from the null cost. For the set of 10 hypotheses, the difference $\Delta \ge 597.58$ which attests to the high quality of the pharmacophore model. The standard indicators such as the RMSD between the hypotheses ranged from 5.35 to 6.08, and the squared correlation coefficient (R^2) falls to an interval from 0.91 to 0.87" [13]. The first PH4 hypothesis with the total costs (475.78) and best RMSD and R^2 was retained for further analysis. The statistical data for the set of hypotheses (costs, RMSD, R^2) are listed in Table 4. The configuration cost (14.78 for all hypotheses) below 17 confirms this pharmacophore as a reasonable one. The evaluation of Hypo 1 is the mapping of the best active training set 17i (Fig. 4 (D)) displaying the geometry of the Hypo1 pharmacophore of PPO activation. The regression equation for pK_i^{exp} vs.

pK_i^{pre} estimated from Hypo1: pK_i^{exp} = 0.947x pK_i^{pre} + 0.41 (2) (n = 29. R² = 0.91. R²_{xv} = 0.91. F-test = 292.32. σ = 0.16, α > 95 %) is also plotted on Fig. 4 (E). The ratio pK_i^{pre}/pK_i^{exp} (the pK_i^{pre} values were estimated using correlation Equation ((2), Table 5) calculated for the training set and validation set is near of the 1. Therefore the PH4 is good potentially to choice the new CQB analogs.

We can carry out computational design and selection of new CQB analogs with elevated activation potencies against PPO.

3.3 Virtual Screening

In silico screening of a virtual (combinatorial) library can lead to hit identification as it was shown in our previous works on inhibitors design [14,16-20]. An initial virtual library (VL) was generated by substitutions at positions for R₁, R₂ and R₃ (Table 5) on the CQB scaffold. During the virtual library enumeration the R-groups listed in Table 5 were attached to in positions R₁.R₂ and R₃ of the CQB scaffold to form a combinatorial library of the size:

 $R_1 \times R_2 \times R_3 = 1 \times 1 \times 118 = 118$ analogs.

Table 5. The ratio pKi ^{pre} /pKi ^{ex}	$^{\circ}$ calculated for the training se	t and validation set
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Training set	рК _i ^{pre}	рК ^{ехр}	pKi ^{pre} /pK ^{exp}
17i	8.02	8.17	0.981
17h	7.84	8.01	0.979
9h	7.77	7.96	0.976
17f	7.76	7.68	1.011
8h	7.74	7.7	1.006
9f	7.74	7.34	1.055
9e	7.35	7.36	0.999
17e	7.33	7.24	1.013
19	7.27	7.14	1.019
9k	7.27	7.43	0.978
8k	7.19	7.32	0.982
8e	7.18	7.16	1.003
9g	7.05	7	1.006
17g	7.02	7.08	0.992
17d	6.97	6.96	1.001
8d	6.84	7.01	0.975
8g	6.75	6.77	0.997
91	6.71	6.96	0.965
17a	6.70	6.7	1.001
9a	6.70	6.7	1.000
9j	6.69	6.59	1.015
17c	6.68	6.85	0.975
17j	6.68	6.77	0.986
17b	6.67	6.32	1.056
17m	6.65	6.92	0.961
9c	6.55	6.8	0.964
8a	6.51	6.68	0.974
8b	6.40	6.47	0.990
8c	6.35	6.57	0.966
8j	5.60	5.6	1.001
Validation set	pKi_theo	pKi_exp	pKi_theo/pki_exp
9i	8.10	8.12	0.998
8i	7.85	7.85	1.000
17k	7.85	7.85	1.000
9d	7.05	7.01	1.006
8f	6.97	6.92	1.007
171	6.76	6.7	1.009
9b	6.74	6.68	1.009

The focused library of 118 analogs was further screened for molecular structures matching the 3D-QSAR PH4 pharmacophore model Hypo1 of PPO activation, 28 best fitting analogs (PH4 hits) then underwent complexation QSAR model screening. The computed GFE of PPO-CQBx complex formation, their components and predicted activity Ki^{pre} calculated from the correlation Equation (1) (Table 3) are listed in Table 6.

3.4 Novel CQB Analogs

The design of virtual library of novel analogs was guided by structural information retrieved from the CQBx active conformation and the pharmacophore model, were used for the selection of appropriate substituents. The hydrophobic feature of PH4 at the position R_3 show clearly the type of group.

4. DISCUSSION

4.1 Binding Mode Cycloalka [d]quinazoline-2,4dione-Benzoxazinones

In more the good quality of the QSAR model, pharmacophore and the focusing catalytic pocket of PPO interactions between the CQB and active site residues such as Leu334, Phe392, Leu372, and Leu356 are revealing. The key interactions responsible for the CQB affinity to PPO, such as hydrogen bonds, van der Waals interactions, and hydrophobic contacts, etc. As displayed in the 2D of Fig. 2, the binding of 17i most active ligand in the TS to the active site of PPO is supported by this interactions. This analysis indicates that compounds with hydrophobic groups may be advantageous to PPO-inhibiting activity.



(A)

(B)



(C)

(D)

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(E)

Fig. 4. (A) Distances between centers, (B) angles between centers of pharmacophoric features
(C) features, (D) mapping of pharmacophore of ligand with the most partial agonist 17i. Feature legend: HYDA = Hydrophobic Aliphatic (blue), HBA = Hydrogen bond Acceptor (green). (E) Correlation plot of experimental vs. predicted activation activity







(B)

Fig. 5. (A) 2D schematic interaction diagram of the most potent inhibitor CQB22 at the active site of PPO and (B) 3D schematic interaction













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Table 7. CQB analogues inhibitors of PPO

Analogues	ΔGbinding	∆Gsol (kcal)	T∆S (kcal)	∆Gcompl (kcal)	ΔΔGcompl (kcal)	Ki_theo(nM)
17i	-58.79	-383.92	20.19	-462.90	0.00	6.7
CQB1	-61.43	-385.56	20.81	-467.80	-4.90	0.76
CQB2	-56.15	-383.95	20.58	-460.67	2.22	11.6

Analogues	ΔGbinding	∆Gsol	TΔS	∆Gcompl	∆∆Gcompl	Ki_theo(nM)
_	_	(kcal)	(kcal)	(kcal)	(kcal)	
CQB3	-51.25	-383.64	20.52	-455.41	7.49	87.1
CQB4	-51.25	-384.16	20.29	-455.70	7.20	78.4
CQB5	-54.24	-384.04	20.41	-458.69	4.21	24.8
CQB6	-56.74	-384.02	20.41	-461.17	1.73	9.6
CQB7	-53.13	-384.35	20.33	-457.81	5.09	34.7
CQB8	-51.26	-383.74	20.57	-455.58	7.32	81.7
CQB9	-51.35	-383.70	20.61	-455.66	7.24	79.1
CQB10	-49.48	-384.05	20.51	-454.04	8.86	147.4
CQB11	-63,28	-383.44	21.24	-467.96	-5.06	0.7
CQB12	-62.47	-382.09	20.76	-465.32	-2.43	1.9
CQB13	-62.64	-380.04	20.80	-463.48	-0.59	3.9
CQB14	-57.81	-385.56	20.81	-464.18	-1.28	3.03
CQB15	-60.46	-383.37	20.68	-464.52	-1.62	2.7
CQB16	-61.44	-380.12	20.65	-462.21	0.68	6.4
CQB17	-63.13	-378.77	21.24	-463.14	-0.25	4.5
CQB18	-60.61	-380.06	21.27	-461.94	0,95	7.1
CQB19	-60.06	-383.50	20.98	-464.55	-1.65	2.6
CQB20	-55.99	-383.80	20.81	-460.60	2.30	11.9
CQB21	-53.72	-384.39	20.82	-458.92	3.97	22.7
CQB22	-69.39	-380.01	20.86	-470.26	-7.36	0.29
CQB23	-54.31	-379.73	20.39	-454.42	8.48	127.2
CQB24	-58.82	-379.58	20.71	-459.10	3.79	21.1
CQB25	-61.84	-379.71	20.65	-462.20	0.70	6.5
CQB26	-61.78	-379.78	20.81	-462.37	0.52	6.04
CQB27	-58.60	-380.12	20.67	-459.39	3.51	18.9
CQB28	-50.35	-380.47	20.26	-451.08	11.82	458.5

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Table 8. CQB analogues inhibitors of PPO, depicted in 2D,





CQB25 (6.45nM)

CQB28 (458.5nM)

CQB26 (6.04nM)

CQB27 (18.9nM)

4.2 Analysis of New Ligands from *In* silico Screening

An analysis of structural requirement for PPO activation at the level of hydrophobic contacts with the active site revealed that the substituent at R_3 Table 6. In this job, the use of the pharmacophore model new method to select the best analogs. The theoretical activities calculated by equation (1) are show in Table 7. Top scoring virtual hits analogs of CQB are: CQB1 (0.76nM), CQB11 (0.7nM) and CQB22 (0.29nM). The predicted activity of the best designed CQB analog CQB22 (Fig. 5) reached approximately 23 times lower than that of the most active ligand of the training set 17i with $K_i^{exp} = 6.7$ nM (Table 8).

5. CONCLUSION

Structural investigation of the SAR of Cycloalka [d]quinazoline-2, 4dione- Benzoxazinones as partial PPO agonists from the crystal structure of PPO: CQB complex guided us during preparation of a reliable QSAR model of activation of PPO which correlated computed Gibbs free energies upon complex formation with observed PPO activation potencies. In addition we have derived a 3D-QSAR PH4 pharmacophore model for CQB activation using a training set of 29 and validation set of 7 CQBs with known activation activities [7]. Careful analysis of interactions between the PPO's active site residues and CQBs directed us in the design of an initial diversity virtual combinatorial library of new CQB analogs with multiple substitutions hydrophobic group in R₃. A library screened by matching of the analogs to the PH4 pharmacophore permitted selection of a library subset of CQBs. This subset of 28 best virtual hits was submitted to computation of predicted activation potencies bv the complexation QSAR model. The hit analogs reached predicted activities in the nanomolar concentration range. The hit designed CQB analogs CQB1 (0.76 nM), CQB11 (0.7 nM) and CQB22 (0.29 nM) are recommended for synthesis and subsequent activity evaluation in PPO activation assays and may lead to a discovery of novel weedkiller potent partial PPO agonists. Virtual design method is an excellent trail already used in medicinal chemistry for discovery a good diseases targets therefore the researchers must apply it more in agropastoral.

CONSENT AND ETHICAL APPROVAL

It is not applicable.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES

- 1. Available:https://www.un.org/sustainablede velopment/sustainable-development-goals/
- Wang B, Wen X, Xi Z. Molecular simulations bring new insights into protoporphyrinogenIX oxidase / protoporphyrinogen IX interaction modes. Mol Inf. 2016;35(10):476–482.
- Hao G F, Tan Y, Xu W F, Cao R J, Xi Z, Yang G F. Understanding resistance mechanism of protoporphyrinogen oxidase inhibiting herbicides: Insights from computational mutation scanning and sitedirected mutagenesis. J Agric Food Chem. 2014;62(29):7209–7215.
- Wang D W, Rui-Bo,Yu S Y,Liang L, Ismail I, Li Y H et al. Design, herbicidal Activity, and QSAR Analysis of cycloalka[d]quinazoline-2,4-dione-BenzoxazinoneS as Protoporphyrinogen IX oxidase. J Agric. Food Chem. 2019; 67(33):9254-9264.
- 5. Zuo Y, Wu Q, Su SW, Niu CW, Xi Z, Yang GF. Synthesis, herbicidal activity, and QSAR of novel N-benzothiazolyl pyrimidine-2,4-diones as protoporphyrinogen oxidase inhibitors. J Agric Food Chem. 2016;64(3):552–562.
- Wang DW, Li Q, Wen K, Ismail I, Liu DD, Niu CW, et al. Synthesis and herbicidal activity of pyrido[2,3-d]pyrimidine-2,4dione-benzoxazinone hybrids as proto porphyrinogen oxidase inhibitors. J. Agric. Food Chem. 2017;65(24):5278–5286.
- Jiang LL, Tan Y, Zhu XL, Wang ZF, Zuo Y, Chen Q, et al. Design, synthesis, and 3D-QSAR analysis of novel 1,3,4-oxadiazol-2(3H)-ones as protoporphyrinogen oxidase inhibitors. J Agric Food Chem. 2010;58(5): 2643–2651.
- Zhao LX, Jiang MJ, Hu JJ, Zou YL, Yuan C, Tao Ren et al. Design, synthesis, and herbicidal activity of novel diphenyl ether derivatives containing fast degrading tetrahydrophthalimide. J Agric Food Chem. 2020;68(12):3729–3741.
- 9. Li H, Sutter J, Hoffmann R. Pharmacophore perception, development and use in drug design. Güner O F ed, International University Line: La Jolla, CA, USA; 2000.

- 10. Discovery Studio Molecular Modeling and Simulation Program, version 2.5; Accelrys, Inc.: San Diego, CA, USA; 2009.
- Maple J R, Hwang M-J, Stockfisch T P et al. Derivation of class II force fields. I. Methodology and quantum force field for the alkyl functional group and alkane molecules. J Comput Chem. 1994; 15(2):162-182.
- 12. Kouassi A, Kone M, Keita M, et al. Computer-aided design of orally bioavailable pyrrolidine carboxamide inhibitors of enoyl-acyl carrier protein reductase of mycobacterium tuberculosis with favorable pharmacokinetic profiles. Int J Mol. Sci. 2015;16(12):29744-29771.
- Okra G M, N'Guessan R K, Dali B, N'Guessan H, Kouassi A. Conformational analysis and molecular design of anthranilic acid derivatives as partial agonist of the Farnesoid X Receptor (FXR) with favorable predicted pharmacokinetic profiles. J Comput Chem Mol Model. 2021; 5(2):585-605.
- Kouman KC, Keita M, Kre N'Guessan R et al. Structure-based design and *in Silico* screening of virtual combinatorial library of benzamides inhibiting 2- trans enoyl-acyl carrier protein reductase of mycobacterium tuberculosis with favorable predicted pharmacokinetic profiles. Int J Mol Sci. 2019;20(19):4730.

PMid:31554227 View Article PubMed/ NCBI

 Merk D, Lamers C, Ahmad K et al. Extending the structure-activity relationship of anthranilic acid derivatives as farnesoid x receptor modulators: development of a highly potent partial farnesoid x receptor agonist. J Med Chem. 2014;57(19):8035-8055.
PMid:25255039 View Article PubMed/ NCBI

- N'Guessan H, Megnassan E. Design of phosphonic arginine and hydroxamic acid inhibitors of *Plasmodium falciparum* M17 leucyl aminopeptidase with favorable pharmacokinetic profile. J Drug Des Med Chem. 2017;3(6):98.
- Allangba KNPG, Keita M, N'Guessan K R, Megnassan E, Frecer V, Miertus S. Virtual design of novel *Plasmodium falciparum* cysteine protease falcipain-2 hybrid lactone-chalcone and isatin-chalcone inhibitors probing the S2 active site pocket. J Enzyme Inhib Med Chem. 2019;34(1): 547-561.

PMid:30696325

View Article PubMed/NCBI

- Esmel A, Keita M, Megnassan E, Beguemsi T, Frecer V, Miertus S. Insight into the binding mode of nitrile inhibitors of *Plasmodium falciparum* falcipain-3, QSAR and pharmacophore models, virtual design of new analogs with favorable pharmacokinetic profiles. J Comput Chem Mol Model. 2018;2(1):103-124.
- Kouakou K J L, Keita M, Esmel A E, Dali 19. B, N'Guessan A, Kouassi A F. In silico Desian of Novel Nhydrosulfonylbenzamides inhibitors of dengue RNA-dependent RNA polymerase predicted showing favorable profiles. pharmacokinetic Journal of Computational Chemistry & Molecular Modeling. 2021;5(2):561-584.
- 20. Yéo Y, Keita M, Bibila M, Esmel A E, Dali N'Guessan Molecular Β, Η. and thermodynamic the modeling of protein-ligand interaction. Application to design computer-assisted of anticompetitive inhibitors of human histone deacetylase 2 (HDAC2). SDRP J. Comput. Chem. Mol. Model. 2021;5(2): 585-609.

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