Quantum Chemical Studies and Pharmacophore Modeling for Designing Novel Keap1 Antagonists that Enhance Nrf2 Mediated Neuroprotection

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1 Abstract

In recent years, the significance of oxidative stress in the pathophysiology of Neurodegenerative/developmental diseases like Attention Deficit Hyperactivity Disorder, Parkinson's and Alzheimer's is being studied at an accelerating pace. Nrf2 activation via Keap1 inhibition is an established strategy for enhancing the activity of the cellular antioxidant mechanism. In this study pharmacophore modeling was employed to design efficient Keap1 inhibitors from well-known polypharmacological phytochemicals after extensive structural modifications to improve their pharmacodynamic, pharmacokinetic and drug-likeness qualities. Quantum chemical calculations at the B3LYP/6-31G (d, p) level of theory were performed for geometry optimization of the novel ligands and for computing their electronic properties. Molecular docking studies and comparative interaction analysis ranked the Resveratrol-4 derivative as the best multi-domain antagonist of the Keap1 protein. The following study presents the application of novel, modified, phytochemical derivatives, as efficient antagonists of the Keap1 protein for enhancing neuroprotection from redox insults.

Keywords: Oxidative stress; Attention Deficit Hyperactivity Disorder; Structural Modification; Quantum chemical studies; Nrf2-Keap1.

2 Introduction

Attention Deficit Hyperactivity Disorder or as described by Dr. Russell Barkley as the "Diabetes of the human brain" is a seriously underrated neurological disorder, if untreated, has lifelong socio-economic repercussions like poorer self-regulation for deferred gratification, ergo higher selfishness and lower stability in relationships, increased impulsivity, poor executive and cognitive function, poor decision-making ability, lower levels of general happiness and quality of living (Barkley 2014). Parental virility is a major contributing factor to the phycological health of children who have Attention Deficit Hyperactivity Disorder. Parental age, lack of virility, socio-economic instability and marital discord are all triggers that induce unhappiness-based negative reinforcement leading to the early onset of Attention Deficit Hyperactivity Disorder in genetically predisposed children, due to the consistent snowballing effects of catecholaminergic imbalance and neuronal oxidative stress. Etiological studies suggest that Attention Deficit Hyperactivity Disorder is physiologically aggravated due to stress (oxidative and otherwise caused due to lack of physical/mental wellbeing and maintenance) among several other factors.

Catecholaminergic imbalance and oxidative stress are some of the key interdependent factors that cause neurodegeneration. The combinatorial effects of excess / unmitigated nitrosative and oxidative stress caused by the dysregulation of the neuronal antioxidative mechanism can cause a variety of biochemical aberrations that eventually lead to neuronal cytotoxicity and death. Under excessive oxidative stress from ROS and RNS, the catecholaminergic catabolism proceeds in undesirable secondary oxidative pathways that culminate in the deposition of neurotoxic byproducts, by-products which accumulate in the neuroplasm and causes its death and eventual necrosis. Some of the neurotoxins produced from such aberrant pathways are, dopamine-oquinone, dopaminochrome and 6-hydroxydopamine from dopamine and norepinephrine-oquinone, norepinephrine semiquinone and 6-nitronorepinephrine from norepinephrine. Further, The neurotoxic by-products from the secondary oxidative pathways with epinephrine as the substrate resemble the by-products of norepinephrine (Napolitano et al. 1999a, 1999b, 2011).

Additionally, other substrates for neuronal redox imbalance and oxidative stress-induced catabolism are polyunsaturated fatty acids (PUFA), purines among several others. Unmitigated redox imbalance results in the excessive catabolism of cerebral docosahexaenoic acid (DHA is a very important fatty acid for normal brain development and function (Lauritzen et al. 2016; Hashimoto et al. 2017)) which terminates in the production of copious amounts of the nontoxic expiratory Attention Deficit Hyperactivity Disorder biomarker, ethane (Ross et al. 2003; Yakubenko and Byzova 2017). Likewise, nucleic acid bases (Adenine and Guanine) under similar conditions produce 8-oxo-7, 8-dihydroguanine (8-oxoG) or 8-oxodeoxyguanosine (8-oxodG) leading to neuronal DNA damage (which could be the etiological basis for the genetic component of Attention Deficit Hyperactivity Disorder) (Chovanová et al. 2006). The accumulation of these toxins lead to neuronal cell death and thereby contribute to the pathophysiology and progression of Attention Deficit Hyperactivity Disorder (ND's).

The Nuclear factor erythroid 2-related factor 2 (Nrf2) pathway is the parent cascade that regulates the proteins and enzymes which control the mechanism of neuronal antioxidation. The plethora of products of the Nrf2 associated genes that aid in the neuronal redox homeostasis, makes

the Nrf2 gene cascade the master regulator of neuronal redox equilibrium. Under redox balanced, housekeeping states, the highly potent Nrf2 proteins, are found to be in low cytosolic numbers by the action of the Kelch-like ECH associated protein 1 (Keap1) proteins. The Keap1 protein ubiquitinates Nrf2 and targets it for eventual proteolytic degradation by the 26S proteasome. But, under oxidative duress, due to the neuroplasmic accumulation of electrophiles (toxins, ROS and RNS), the keap1 activity is diminished (sensory inhibition) and the Nrf2 proteins, through a network of biomolecular interactions, activate the transcription and translation of the genes and RNA, whose products (Superoxide dismutase, catalase, GSH peroxidases) ultimately return the system to redox homeostasis (Johnson et al.; Kaspar et al.; Ma 2013). A recent study on Protandim mediated Nrf2 activation in erythrocytes reports a 34 % increase in the amount of superoxide dismutase produced (Nelson et al. 2006). Therefore, Keap1 inhibition for the activation of Nrf2 cascade for enhanced neuronal antioxidant activity could be a lucrative option for ameliorating the damage caused by oxidative stress in neurons that are inherently flawed, as in the case of genetically acquired Attention Deficit Hyperactivity Disorder.

Structural modification for drug design has been carried out extensively by repurposing existing drugs (Uzzaman and Mahmud 2020), phytochemicals (Gordaliza 2007), peptides (Di 2015), siRNA (Shukla et al. 2010) among other biomolecules as therapeutics. The novel coronavirus disease 2019 pandemic is showing the world the dire need for developing efficient drugs in a shorter time frame, all the while utilizing fewer resources. With that thought, Pharmacophore modeling of pre-existent bio-chemicals has significant advantages over their traditional ab initio counterparts, as it requires fewer iterations of computation and also due to the sheer abundance of information that is already available regarding the basal molecule, like its potency, drug-likeness properties, toxicity, pharmacodynamics and mode of action. Further, quantum chemical studies based on density functional theory are an excellent computational chemistry technique for studying the environment (solvent, cytoplasm, gas etc.) specific structure and electrochemical properties of novel drugs. DFT studies provide ample insight regarding the charge localization, molecular electrostatic potential, reactivity, ionization potential and electron affinity of the ligands among others. These quantum chemical descriptors are highly beneficial for understanding the redox chemistry of the ligands during biomolecular interactions (ligandreceptor) and is also beneficial in comprehending the behavior of the ligand in solvated micro/macro environments.

Herein, structurally modified, novel, antioxidant, phytochemical derivatives that induce Nrf2 activation by inhibiting keap1 activity, have been designed and presented after a comprehensive investigation by quantum chemical techniques and molecular modeling studies that are augmented by drug-likeness screening and pharmacokinetic analysis.

3 Materials and Methods

3.1 Library Generation and Structural Modification

A ligand library, comprising of four parent phytochemicals known for their therapeutic and most importantly, antioxidant properties, namely Decussatin (Karsten et al. 2012), Loliolide (Yang et al. 2011), Resveratrol (stilbene derived polyphenolic phytoalexin found in grapes and raisins) (Gülçin 2010), Scopoletin (Shaw et al. 2003), were selected for further investigation. The structures, physical and chemical parameters of the phytochemicals were obtained from the PubChem database (Kim et al. 2019). Further, ADMET-SAR (Cheng et al. 2012) software tool and the CB-ligand platform (Liu et al. 2014) (https://www.cbligand.org/BBB/) were employed to predict the blood-brain barrier permeability index of these four phytochemicals. The blood-brain barrier permeability index of the phytochemicals was predicted to be strongly positive by both of these tools. Therefore, these were finalized as the parent phytochemicals for further ligand library generation via structural modification. Structural modifications were performed primarily using ChemDraw Ultra 12.0 (2D) (Cousins 2011). To increase efficacy, antioxidant activity, and binding efficiency to the active sites of Keap1, the hydroxyl functional groups present in the parent molecules were targeted for modification via replacement as they can contribute to post catabolic oxidative stress further. The hydroxyl groups were serially replaced with trifluoromethyl (-CF₃) groups (commercially approved selective serotonin reuptake inhibitors (antidepressants), Fluvoxamine (Brand name: Luvox) (Wilde et al. 1993) and Fluoxetine (Brand name: Prozac) (Altamura et al. 1994) contain one trifluoromethyl moieties each) (Uzzaman and Mahmud 2020) and amino (-NH₂) groups. It is widely known that; the hydroxyl groups contribute to excess redox imbalance and oxidative damage as compared to nitrogen groups (Hybertson et al. 2011). The modified 2D structures were converted into a 3-dimensional format using Chem3D Pro 12.0, and the final structures were saved in .sdf and .pdb file formats. Modifications to the Decussatin molecule was not performed as none of its derivatives passed the blood-brain barrier threshold index as set by the two previously mentioned tools. A total of eleven derivatives were designed with a total of 15 ligands in the final ligand library (Figure 1). These ligands were subjected to further studies.

Figure 1: 2D structures of the ligand library: (A) Decussatin, (B-G): Loliolide (L, L1-6), (H-L): Resveratrol (R, R1-4), (M-O): Scopoletin (S, S1-2).

The ADMET (Absorption, Distribution, Metabolism, Excretion, and Toxicity) assessment of the prepared library was conducted using AdmetSAR (Cheng et al. 2012) tool. A preliminary screening was conducted based on Lipinski's rule of five (Lipinski 2000). Further, the blood-brain barrier permeability parameter was prioritized as the most stringent rule (BBB > 0.80). Additional parameters like Human intestinal absorption, Caco-2 permeability, P-glycoprotein substrate and inhibitor, Subcellular localization index, Renal organic cation transporter, HERG inhibition, AMES toxicity, and carcinogenicity were computed for a pervasive assessment of pharmacokinetic desirability. Ligands with severe outliers were simply discarded from the library, and the ligand derivatives with outliers that could merit from minor structural modifications were subjected to the same.

3.3 Quantum Chemical Calculations

Quantum chemical calculations of the drug-likeness screened library were performed using the General Atomic and Molecular Electronic Structure System (GAMESS-US) software (Gordon and Schmidt 2005). The Avogadro molecule editor and visualizer software (Hanwell et al. 2012) was used to generate input simulation files for the GAMESS program. MacMolPlt software (Bode and Gordon 1998) was utilized for visualizing the output (.log) files. Initial ligand optimization was performed using the Avogadro software. The pH of the system was adjusted to that of cerebrospinal fluid, i.e., 7.2 (Albrecht et al. 2020). The Merck molecular force field 94s (MMFF94s) was set up for primary optimization of the ligand geometry with water as the solvent system. Secondary optimization using GAMESS was performed using the unrestricted density functional theory with Becke's (B) three-parameter hybrid model (Becke 1988) along with the Lee, Yang and Parr's (LYP) non-local correlation functional (B3LYP) (Lee et al. 1988) along with Pople 6-31G (d, p) as the basis set for the geometry optimization and electronic property calculation of all ligands. The energies of the frontier molecular orbitals, highest occupied molecular orbital (HOMO), lowest unoccupied molecular orbital (LUMO), and bandgap were also calculated at the B3LYP/6-31G (d, p) level of theory. Further, electronic properties like electronic energy, free energy, dipole moment, molecular electrostatic potential along with bond angles and distances, were calculated. The pictorial illustrations of the electronic spin density distributions of the HOMO and LUMO represented ligand structures were generated and color-coded to differentiate the positive and negative electrostatic potentials by using the MacMolPlt visualizer. The Parr and Pearson's interpretation (Parr and Pearson 1983; Lee et al. 1988; Parr and Yang 1995) along with the Koopman's theorem (Koopmans 1934) were utilized for calculating the reactivity parameters viz electron affinity (EA), ionization energy (IA), chemical potential (µ), hardness (η), softness (σ), electronegativity (χ), electrophilicity (ω), electron-accepting power (ω^+) and electron-donor power (ω) of the ligands.

3.4 Preparation of Ligands

Ligand preparation was performed using the UCSF Chimera 1.13.1 suite (Pettersen EF, Goddard TD, Huang CC, Couch GS, Greenblatt DM, Meng EC 2004). The ligands were minimized via the structural editor tool with 100 steepest descent steps and ten conjugate gradient steps with a step size of 0.02 Å. Missing hydrogens and Gasteiger partial charges were added. AMBER force field 14SB (AMBERFF14SB) was set up during the whole procedure. The structures were saved in pdbqt format for further studies.

3.5 Receptor Selection and Analysis of Active Site

The Broad complex, Tramtrack, and Bric à brac (BTB) (50-184 amino acid residues) domain of the Keap1 protein is crucial for its homo-dimerization, activity, and structural integrity. Further, the Kelch domain (327-611 amino acid residues) in Keap1 (one domain per monomer) is essential for the capture and binding of Nrf2 proteins. The DLG and ETGE domains (23-82 amino acid residues) in the Neh2 region of the Nrf2 protein is fundamental for it to bind simultaneously to two of the Kelch domains of the Keap1 homodimer (Jnoff et al. 2014; Canning et al. 2015). Molecular docking analysis was performed with these two domains (BTB and Kelch) as the active sites to investigate the prepared library for dual-site binding efficiency. The structure of the Broad complex, Tramtrack, and Bric à brac (BTB) domain (PDB Code: 5DAD) (Resolution: 2.61 Å) and the Kelch domain (PDB Code: 4L7B) (Resolution: 2.41 Å) of the Keap1 protein were obtained from the Protein Data Bank (RCBS) (Berman 2000). Ramachandran Plot analysis was performed to assess the stereochemical suitability and torsional stability of the amino acids of the selected protein receptors via the MolProbity server (Davis et al. 2007) and the PROCHCECK tool (Laskowski et al. 1996).

3.6 Preparation of Receptor

'Dock Prep' of the UCSF Chimera 1.13.1 suite was employed to prepare the selected receptors for molecular docking simulation. Initially, the co-crystalized, (1S,2R)-2-[(1S)-1-[(1,3-dioxo-2,3-dihydro-1H-isoindol-2-yl)methyl]1,2,3,4tetrahydroisoquinolincarbonyl]cyclohexane-1-carboxylic acid co-crystallized ligand from the kelch domain of the receptor (4L7B) and the (6aS,7S,10aS)-8-hydroxy-4-methoxy-2,7,10a-trimethyl-15,6,6a,7,10,10a-hexahydrobenzo[h] quinazoline-9-carbonitrile (TX6201) ligand from the BTB domain of the keap1 receptor (5DAD) were removed. Secondly, the Dunbrack rotamer library was used to add the incomplete side chains of the receptors. Next, missing hydrogens were added for the completed sidechains. Further, Gasteiger partial charges were added, and the AMBER force field 14SB (AMBERFF14SB) was set up for the preparation process. The prepared receptors were saved in pdbqt format and were subject to further analysis.

Molecular docking simulation of the prepared ligands into the prepared receptor was carried out using the widely accepted AutoDock Vina (Trott and Olson 2009) software tool using the PyRx version 0.8 software (Dallakyan and Olson 2015). Rigid docking was performed to investigate the ligands-receptors binding efficiency at the lowest degree of freedom for higher reliability. Water molecules in the system were ignored and were set to not interfere with the simulation in any way. The grid box of the BTB domain (PDB Code: 5DAD) was fixed at X: Y: Z :: -25.4541 : -2.91812: 10.0365 and likewise the grid box for the Kelch domain (PDB Code: 4L7B) was fixed at X: Y: Z:: -1.50946: 3.16751: -27.1727 coordinates. The size of the grid box was heuristically adjusted during each simulation to completely incorporate the ligand and the domain that is under purview. Exhaustiveness of the search was set to 10 and the simulation of all ligands including the co-crystalized ligands were conducted. The binding energy (kcal/mol) of the docked conformation with the highest binding energy coupled with the lowest RMSD (upper and lower bound) was identified, and the same conformation was exported to Accelrys discovery studio (Accelrys Software, Inc., San Diego, CA, USA) for interaction analysis. Further, the ligandreceptor interaction analysis was performed, and docking attributes like type of bonds, interacting atoms-residues, the bond length was recorded, and pictorial illustrations were obtained. Additionally, LigPlot (Laskowski and Swindells 2011) program was also used to assess the ligandreceptor interactions for scaffolding the interaction analysis results obtained from the discovery studio by acting as an auxiliary validator.

4 Results and Discussion

4.1 Druglikeness Screening and Pharmacokinetic Desirability Assessment

The complete ligand library qualified the Lipinski's rule of five, which was the primary parameter considered for drug-likeness screening. Further, the blood-brain barrier permeability was the second parameter that was prioritized for drug-likeness screening. The top five ligands with the highest BBB permeability were, R4 > L5 > L1 > L > L2. Likewise, the top five ligands with the highest human intestinal absorption were R4 > L5 > L > L2 > L4. Further, Caco2 cell line permeability prediction resulted in the following descending order of penetration in ligands, R4 > L5 > S1 > R1 > S2. It is noteworthy to observe that R4, L5, and S1 are the top three different ligands (with three different parent molecules) with the highest pharmacokinetic desirability. The additional pharmacokinetic analysis resulted in desirable scores for these three ligands concerning toxicity, metabolism, and distribution. From the results obtained from the conducted drug-likeness screening and pharmacokinetic analysis, it can be envisaged that the derivatives, R4, S1, L5 would successfully permeate through to the systemic circulatory system and also penetrate past the blood-brain barrier in potent quantities. Upon analysis of the bioavailability radar representations of the ligands, it was found that R4 had minor outliers pertaining to its lipophilicity (XLOGP3; -0.7 < XLOGP3 < +5), unsaturation (Fraction Csp3; 0.25 < (Fraction Csp3) < 1) and insolubility (LogS (ESOL); 0 < LogS (ESOL) < 1) values. Likewise, S1 had one outlier in its level of unsaturation. Further, the values of L5 was well within the desirable (pink)

region of the bioavailability radar. The bioavailability radar illustrations of these ligands, as obtained from the SwissADME software (Daina et al. 2017), are represented in (Figure 2). The results for the complete pharmacokinetic (absorption, distribution, metabolism, elimination) analysis for the entire ligand library is tabulated and presented (Table 1).

Figure 2: Bioavailability radar illustrations: (A) Loliolide (L5); (B) Resveratrol derivative-4 (R4); and (C) Scopoletin (S1).

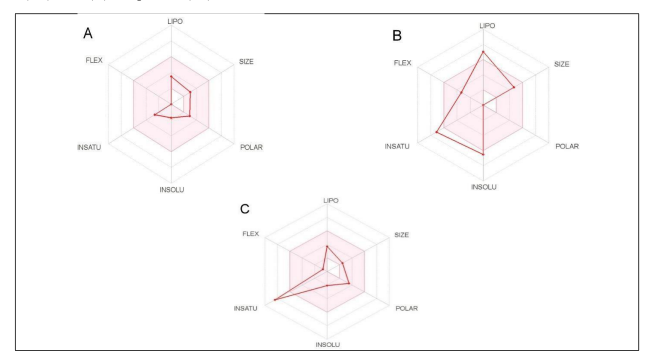


Table 1: Pharmacokinetic analysis of the designed ligand library.

Sl. No	Ligands	Blood- Brain Barrier	Human Intestinal Absorption	Caco-2 Permeability	HERG- Gene Inhibition	AMES Toxicity	Carcinogenicity (Three-class)
1	Decussatin	0.5499	0.852	0.8865	0.9463	0.8251	0.5420
2	Loliolide	0.9384	1	0.5978	0.9723	0.9498	0.5326
3	L1	0.9408	0.9937	0.5309	0.9863	0.9331	0.5731
4	L2	0.9264	1	0.5237	0.9877	0.9118	0.591
5	L3	0.8345	1	0.5223	0.9553	0.9357	0.6692
6	L4	0.8692	1	0.5065	0.9511	0.9439	0.6637
7	L5	0.9668	1	0.602	0.9591	0.924	0.5729
8	Resveratrol	0.5900	0.9952	0.8915	0.8933	0.9462	0.6825
9	R1	0.5352	0.9831	0.5853	0.8707	0.8629	0.8109
10	R2	0.694	0.9651	0.5549	0.9423	0.8781	0.7352
11	R3	0.7351	0.9426	0.5599	0.9031	0.8500	0.8179
12	R4	0.9917	1	0.8014	0.9633	0.8399	0.4747
13	Scopoletin	0.7975	0.9399	0.8944	0.9672	0.9431	0.8059
14	S 1	0.7751	0.9892	0.6018	0.9927	0.6303	0.5913
15	S 2	0.5968	0.9885	0.5737	0.9232	0.8653	0.7800

4.2 Quantum Chemical Calculations: DFT Studies

Although the Aufbau principle is used to estimate the allotment of electrons into the orbitals of an atom, DFT studies help understand the charge localization and charge migration in a multiatomic molecular system for varied ionic states (-2, -1, 0, +1, +2 and so on). Quantifiable electrochemical properties of the molecule at different ionic states are crucial for understanding its reactivity, behavior, binding affinity (if in a complex), and most certainly, its redox chemistry and antioxidant potential. Geometry optimization of the ligands using DFT studies alleviated the steric hindrance in the molecules, and the most stable coordinates of the ligands were obtained. DFT studies conducted on the constructed ligand library revealed the highest electron affinity for Loliolide-5 at 1.986 eV, followed by Resveratrol-4 at 1.8775 eV. Scopoletin-5 was placed at ninth place among all ligands. Likewise, the electron-donating power (ω -) of Loliolide and its derivatives were the highest among all the other ligands (L5 > L > L2 > L4 > L1 > L3). This was followed by Resveratrol-4 with the second-best electron-donating power (ω-) at 108.74 eV (R4 > R > R1 > R2 > R3). Scopoletin and its derivatives displayed the lowest transition energy (ΔE = LUMO - HOMO), indicating the highest bioactivity among the rest (S2 > S > S1). Meanwhile, resveratrol-4 was ranked at 11^{th} place with $\Delta E = 4.24497$ eV. Murugavel et al., after experimental and computational studies, have reported a ΔE value of 4.6431 eV along with an electron affinity of 1.4438 eV as an excellent energy parameter for their novel human topoisomerase IIα inhibitor (Murugavel et al. 2019). Hence, it can be speculated that the Scopoletin derivatives (S, S1-2) and R4 have the highest bioactivity and the potential for being an efficient inhibitor since lower transition energies and higher electron affinities (Top five ligands (including L5, R4, S1) with

electron affinities > 1.68710 eV and top eleven ligands with $\Delta E < 4.24497$ eV) was observed for the ligands designed in this study. The LUMO was found to be evenly distributed on both the benzene rings of the R4 molecular structure. In contrast, the HOMO orbitals were found to be majorly distributed in and around the central C3=C4 doubly unsaturated linker bond (Figure 3(A)). Likewise, the HOMO orbitals of the S1 derivative were found to be majorly distributed on the heterocyclic (-O) pyran (C10-C12-C11) ring. Whereas, the LUMO orbitals were found to accumulate onto the homo (C7-C6-C14) hexacyclic ring which is in between the pyran-2-one ring and the adjunct Nitrite (-NO2) and methoxy (-OCH3) functional moieties and to a medium extent over the adjacent heterocyclic pyran ring (Figure 3(B)). The color-coded contour maps of the Molecular electrostatic potentials that represent the electronegative and electropositive centers of these ligands were generated and analyzed (Figure 4 (A, B)). The contour maps of the remaining ligands are represented in (Figure (S2)). Finally, the R4 and S1 derivatives of Resveratrol and Scopoletin were found to be the best candidates with the highest potential for both antioxidant and inhibitory binding functions. Hence, these molecules were given more focus for further studies. The quantum reactivity parameters of the ligands are tabulated in (Table (2)). The electronic properties and energy profile of the ligands as computed by GAMESS are tabulated in (Table (3)).

Figure 3: The color-coded frontier molecular orbitals depicting the Highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO): (A) Resveratrol derivative-R4; (B) Scopoletin derivative-S1.

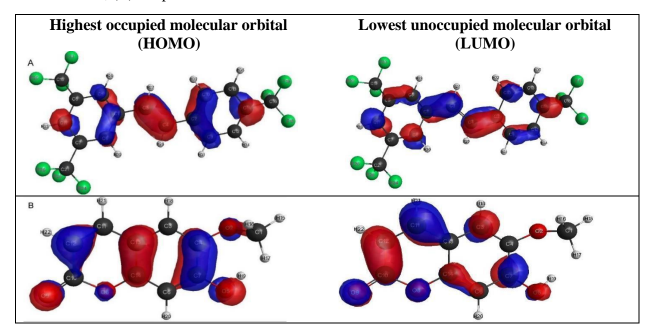


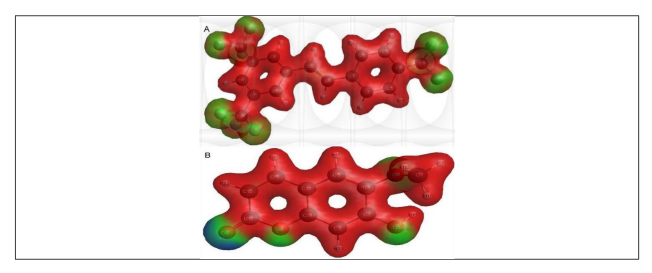
Table 2: The quantum reactivity parameters (eV) computed for the ligand library based on Koopman's theorem and Parr and Pearson's interpretation. (IE: Ionization energy; EA: Electron Affinity; EN: Electronegativity; CP: Chemical Potential; EAP: Electron Accepting Power; EDP: Electron Donating Power).

Sl. No	Ligand	EHOMO (eV)	ELUMO (eV)	ΔE (eV)	IE (eV)	E A (eV)	Ε N (χ) (eV)	C P (μ) (eV)	Hardness (η) (eV)	Softness (σ) (eV)	Electro philicity (ω) (eV)	E A P (ω+) (eV)	E D P (ω-) (eV)
1	Decussatin	-5.88	-1.82	4.05	5.88	1.82	3.85	-3.85	2.03	0.49	15.03	32.63	95.92
2	Loliolide (L)	-7.13	-1.03	6.10	7.13	1.03	4.08	-4.08	3.05	0.33	25.39	39.88	191.53
3	L1	-6.48	-0.71	5.77	6.48	0.71	3.59	-3.59	2.88	0.35	18.61	26.66	146.19
4	L2	-6.61	-0.63	5.99	6.61	0.63	3.62	-3.62	2.99	0.33	19.60	26.97	156.67
5	L3	-6.34	-0.98	5.36	6.34	0.98	3.66	-3.66	2.68	0.37	17.95	28.85	134.02
6	L4	-6.61	-0.63	5.99	6.61	0.63	3.62	-3.62	2.99	0.33	19.60	26.97	156.67
7	L5	-7.81	-1.99	5.82	7.81	1.99	4.90	-4.90	2.91	0.34	34.93	69.00	235.09
8	Resveratrol (R)	-5.33	-1.20	4.14	5.33	1.20	3.27	-3.27	2.07	0.48	11.03	20.59	76.46
9	R1	-5.20	-1.14	4.05	5.20	1.14	3.17	-3.17	2.03	0.49	10.19	18.86	70.97
10	R2	-5.14	-1.06	4.08	5.14	1.06	3.10	-3.10	2.04	0.49	9.82	17.69	69.37
11	R3	-4.98	-0.98	4.00	4.98	0.98	2.98	-2.98	2.00	0.50	8.88	15.68	63.35
12	R4	-6.12	-1.88	4.24	6.12	1.88	4.00	-4.00	2.12	0.47	16.98	36.66	108.74
13	Scopoletin (S)	-6.01	-1.69	4.33	6.01	1.69	3.85	-3.85	2.16	0.46	16.04	33.17	105.25
14	S1	-6.01	-1.69	4.33	6.01	1.69	3.85	-3.85	2.16	0.46	16.04	33.17	105.25
15	S2	-5.47	-1.50	3.97	5.47	1.50	3.48	-3.48	1.99	0.50	12.05	24.63	79.60

Table 3: Molecular formula, Molecular weight (g/mol), Total Energy, Electronic Energy in Hartree, and dipole moment (Debye) of the studied polypharmacological phytochemicals and its modified derivatives.

Sl. No	Name	Molecular formula	Molecular weight (g/mol)	Total Energy (Hartree)	Electronic energy (Hartree)	Dipole Moment (Debye)
1	Decussatin	$C_{16}H_{16}O_{6}$	304.29	-1068.875956	-2858.402387	3.094964
2	Loliolide	$C_{11}H_{16}O_3$	196.24	-654.1009001	-1641.739902	8.796275
3	L1	$C_{11}H_{17}NO_2$	195.26	-634.2080522	-1620.996895	8.114084
4	L2	$C_{11}H_{17}NO_2$	195.26	-634.2319977	-1617.550874	7.418502
5	L3	$C_{11}H_{17}NO_2$	195.26	-634.233399	-1619.974298	8.868181
6	L4	$C_{11}H_{15}$ NO_2	193.24	-633.0297468	-1581.563645	4.599633
7	L5	$C_9H_8F_4O_3$	240.15	-633.0335648	-1602.554482	4.75625
8	Resveratrol	$C_{14}H_{12}O_3$	228.24	-765.9331476	-1801.148688	3.285841
9	R1	$C_{14}H_{13}NO_2$	227.26	-746.0713	-1780.358745	3.200884
10	R2	$C_{14}H_{14}N_2O$	226.27	-726.2084	-1759.946242	2.088747
11	R3	$C_{14}H_{14}N_2O$	226.27	-726.2079	-1762.419272	3.360219
12	R4	$C_{17}H_9F_9$	384.24	-1551.106	-3770.281371	1.187759
13	Scopoletin	$C_{10}H_8O_4$	192.17	-686.2291	-1540.026918	7.633748
14	S 1	$C_{10}H_9NO_3$	191.18	-686.35691	-1540.10211	7.633748
15	S2	$C_{10}H_9NO_3$	191.18	-666.112597	-1516.636141	9.267832

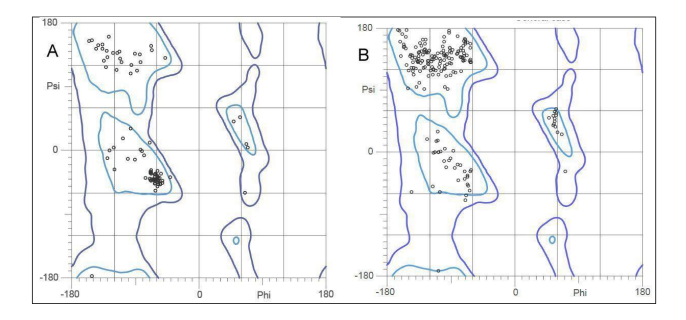
Figure 4: The color-coded contour maps illustrating the molecular electrostatic potential, the red color depicts electropositive centers, the blue regions represent the neutral potentials, and the green areas represent the electronegative centers: (A) Resveratrol derivative-R4; (B) Scopoletin derivative-S1.



4.3 Protein Receptor Analysis by Ramachandran Plot

Ramachandran Plot analysis was performed to assess the structural suitability and torsional stability of the selected receptor proteins (PDB: 4L7B / 5DAD). The allowed values of ψ (Psi) against ϕ (Phi) angles for a particular amino acid in a protein was assessed. There are numerous structures of the Keap1 protein deposited into the Protein Data Bank. For the suitability assessment of proteins for further in-silico studies, Ramachandran plot plays a critical role (Hooft et al. 1997). For the Kelch inhabiting Keap1 protein structure (PDB: 4L7B), 100% (290/290) of the amino acid residues were reported to be in the allowed region. Further, 96.2% (279/290) amino acids were reported to be in the favored area. Likewise, for the BTB domain harboring Keap1 protein structure (PDB: 5DAD), 100% (119/119) of the amino acid residues were reported to be in the allowed region. Further, 95.8% (114/119) of the amino acids were in the favored area. Hence, these two crystal structures were considered as the most suitable, and further studies were conducted on them. The general case representations of the Ramachandran Plots of the individual proteins are depicted in (Figure 5 (A, B)).

Figure 5: General case representations of the Ramachandran plot analysis representations of the selected protein receptor structures depicting the quadrant allocations of the constituent amino acid residues, as obtained from the MolProbity server: (A) BTB domain (PDB: 5DAD); (B) Kelch Domain (PDB: 4L7B).



4.4 Molecular Docking Simulation

The entire ligand library was subjected to molecular docking and was studied for their binding efficiency with both the BTB domain and Kelch domain of the Keap1 protein. An efficient lead should have excellent binding affinities with both the proteins. Such a ligand would prevent the homo-dimerization of the keap1 protein and also would inhibit the binding and eventual ubiquitination and proteolytic degradation of the Nrf2 protein. Upon analysis, it was found that the Resveratrol derivative R4 ranked in the first place (lowest root mean square deviation: upper and lower bound) for binding affinities with both the BTB (-7.3 kcal/mol) and Kelch (-8 kcal/mol) domains. The top five ligands concerning their binding relationships with the BTB domain were R4 > R > R1 > R2 > R3. Unfortunately, the Scopoletin derivatives were ranked at the 7th place in the order S > S2 > S1, respectively. Likewise, the top five ligands concerning their binding affinities with the Kelch domain were R4 > D > L5 > R > R3 > S1. The therapeutic outcome by inhibitory prevention of the Keap1-Nrf2 complex formation is relatively more significant for enhancing the neuronal antioxidant mechanism as compared to the prevention of the Keap1 homodimerization. Additionally, Pharmacokinetic analysis and quantum chemical studies based on density functional theory have regarded the R4, S1 and L5 derivatives as the best ligands with more promise for exerting a desirable level of therapeutic outcome in the paradigm of this study. Hence, the binding/docking of R4, L5, and S1with the BTB and Kelch domains were studied in depth. The interaction analysis obtained from LigPlot and Accelrys discovery studio, for some

ligands, yielded deviant results from one another; this may be because the algorithms employed by these software for predicting/identifying interactions are different from one another. However, an extensive literature review promotes Accelrys discovery studio to be much more efficient compared to LigPlot. Therefore, the interaction analysis obtained discovery studio was given more prominence. Nevertheless, the pictorial representations obtained from LigPlot for the interaction analysis of the ligands that are not under the spotlight of this study with the selected Keap1 and BTB receptors are presented in (Figure S3 (A-O) and Figure S4 (A-O)).

4.5 Interaction Analysis: BTB Domain

The Bis(trifluoromethyl)benzene moiety of the R4 derivative was found to be sequestered into the BTB active site, which apparently is sandwiched between the alpha helices (H131-A143 and K150-Y162) (Figure 6 (A, B)). Also, The R4 Bis(trifluoromethyl)benzene moiety displayed a Pi donor- hydrogen bond and a hydrophobic bond with the Cysteine 151 residue of the BTB active site. In-depth studies, conducted on the significance of the cysteine residues in the Keap1 protein by Saito et al., report that the Cysteine 151 residue is indispensable for the keap1 inhibition mediated 'turn-on' mechanism of the cellular antioxidant system as it is one of three significant cellular oxidative stress sensing cysteine residues (Saito et al. 2016). However, Huerta et al., claim that the cysteine 151 binding mediated inhibition itself is not that significant as compared to the remaining residues that line the BTB binding pocket for keap1 inhibition mediated Nrf2 accumulation (Huerta et al. 2016). Holding that thought, R4 was found to interact with Valine 155, Histidine 154 and 129, Arginine 135, Lysine 131, Tyrosine 85 and Glutamine 86 (Good agreement with LigPlot results (Figure 6 (C))). Further, the fluoride substituted heterocyclic ring adjoining the furan-2-one ring of the L5 derivative was also found to interact with Cysteine 151 residue with a hydrogen bond. L5 also interacted with Lysine 131 and Histidine (154, 129) residues of the BTB domain (Figure 7 (A, B)) (Good agreement with LigPlot results (Figure 7 (C))). Finally, S1 was found to interact with Arginine 135, Lysine 131, Histidine 129, Histidine 154, and most importantly Cysteine 151 with various types of bonds (Figure 8 (A, B)) (Good agreement with LigPlot results (Figure 8 (C))). The complete interaction profile is tabulated in (Table (4)). The remaining pictorial illustrations from PyMol and Discovery studio for the BTB domain is represented in (Figure S5).

Figure 6: The pictorial illustrations for the Resveratrol-4 derivative in the BTB domain (PDB: 5DAD), including the surface view of the top-ranked docked pose (A), interaction analysis from Accelrys discovery studio (B) and the same from LigPlot (C).

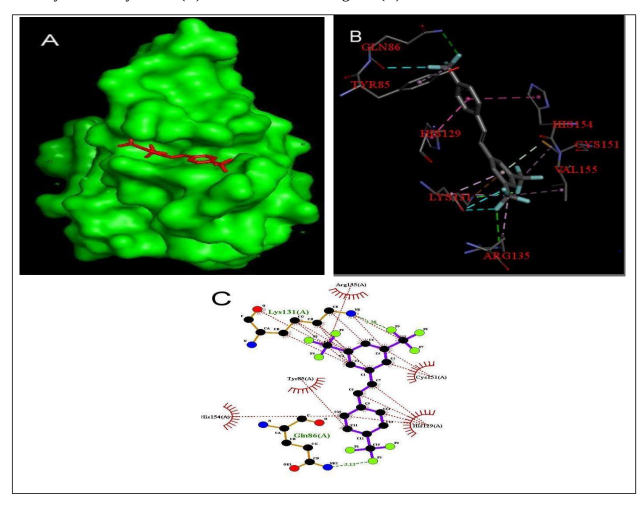


Figure 7: The pictorial illustrations for the Loliolide-5 derivative in the BTB domain (PDB: 5DAD), including the surface view of the top-ranked docked pose (A), interaction analysis from Accelrys discovery studio (B) and the same from LigPlot (C).

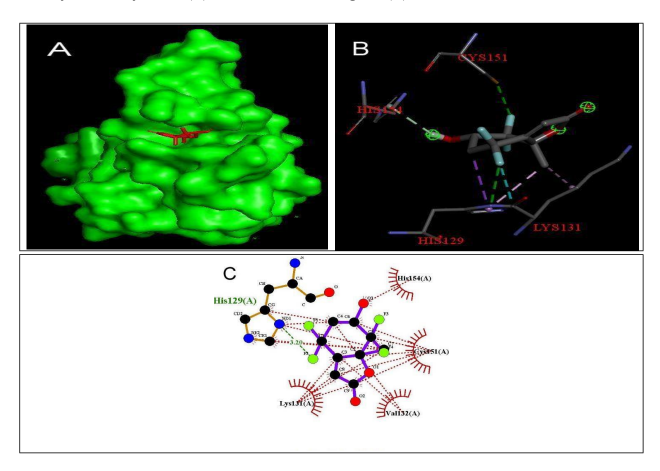
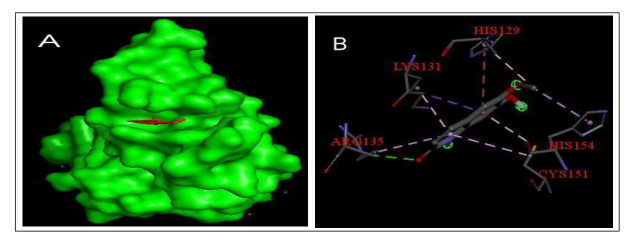


Figure 8: The pictorial illustrations for the Scopoletin-1 derivative in the BTB domain (PDB: 5DAD), including the surface view of the top-ranked docked pose (A), interaction analysis from Accelrys discovery studio (B) and the same from LigPlot (C).



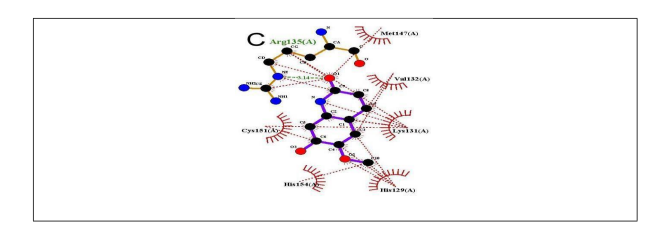


Table 4: Post dock Interaction profile obtained from Accelrys Discovery Studio for the Ligands-BTB domain interactions.

Name	Binding Affinity (Kcal/Mol)	Distance	Bond Category	Interaction Type	Interacting Residues
Decussatin	-6.1	3.53981	Hydrogen Bond	СНВ	Lys131
		3.67302	Hydrogen Bond	CHB	His154
		3.9301	Hydrogen Bond	PDHB	Cys151
		3.65578	Hydrogen Bond	PDHB	Cys151
		4.145	Hydrophobic	PPS	His129
		4.31251	Hydrophobic	A	Cys151
		4.62053	Hydrophobic	A	Lys131
		4.63892	Hydrophobic	A	Arg135
		5.25922	Hydrophobic	PA	Tyr85
		4.54163	Hydrophobic	PA	His154
		5.37109	Hydrophobic	PA	Lys131
		4.3827	Hydrophobic	PA	Lys131
		3.01588	Hydrogen Bond	HHB	Lys131
		3.53981	Hydrogen Bond	CHB	Lys131
		3.67302	Hydrogen Bond	CHB	His154
		3.9301	Hydrogen Bond	PDHB	Cys151
		3.65578	Hydrogen Bond;Other	P-HB; P- SB	Cys151
Loliolide (L)	-4.6	3.82849	Hydrophobic	A	Lys131
		4.41392	Hydrophobic	A	Ays151
		3.98677	Hydrophobic	A	Lys131
		4.62529	Hydrophobic	A	Val132
		4.72689	Hydrophobic	A	Val132

		3.44747	Hydrophobic	A	Ays151
		4.77965	Hydrophobic	A	Val155
		5.43301	Hydrophobic	Α	Met147
		3.3297	Hydrophobic	Α	Ays151
		4.09246	Hydrophobic	PA	His129
Loliolide (L1)	-4.6	3.82918	Hydrophobic	A	Lys131
,		3.45466	Hydrophobic	A	Cys151
		4.77775	Hydrophobic	A	Val155
		5.43854	Hydrophobic	A	Met147
		3.3275	Hydrophobic	A	Cys151
Loliolide (L2)	-4.2	3.89011	Hydrophobic	A	Lys131
		4.40328	Hydrophobic	A	Cys151
		5.25261	Hydrophobic	A	Met147
		3.30096	Hydrophobic	A	Cys151
		4.69165	Hydrophobic	A	Val132
		3.48989	Hydrophobic	A	Cys151
		4.82821	Hydrophobic	A	Val155
		3.82816	Hydrophobic	A	Lys131
		4.77718	Hydrophobic	A	Val132
		4.08718	Hydrophobic	PA	His129
Loliolide (L3)	-4.3	3.55522	Electrostatic	P-C	His129
		4.7201	Electrostatic	P-C	His154
		4.89091	Hydrophobic	A	Val132
		3.74529	Hydrophobic	A	Ays151
		5.06961	Hydrophobic	A	Val155
		3.60548	Hydrophobic	A	Lys131
		4.38336	Hydrophobic	A	Val132
		4.13823	Hydrophobic	A	Lys131
		3.90624	Hydrophobic	PA	His129
		4.07472	Hydrophobic	PA	His129
Loliolide (L4)	-3.8	4.81785	Electrostatic	P-C	His129
		2.68942	Hydrogen Bond; Electrostatic	P-C; PDHB	His154
		4.90609	Hydrophobic	A	Val132
		3.62228	Hydrophobic	A	Cys151
		5.07086	Hydrophobic	A	Val155
		3.41561	Hydrophobic	A	Lys131

		4.39384	Hydrophobic	A	Val132
		3.94794	Hydrophobic	A	Lys131
		4.00906	Hydrophobic	PA	His129
		4.18422	Hydrophobic	PA	His129
Resveratrol (R)	-6.7	4.46708	Electrostatic	P-C	Lys131z
		3.7492	Hydrophobic	PPS	His129
		3.87561	Hydrophobic	PA	Lys131
		5.41155	Hydrophobic	PA	Arg135
		4.57043	Hydrophobic	PA	Cys151
		4.46708	Electrostatic	P-C	Lys131z
		3.7492	Hydrophobic	PPS	His129
		3.87561	Hydrophobic	PA	Lys131
		5.41155	Hydrophobic	PA	Arg135
		4.57043	Hydrophobic	PA	Cys151
		3.18767	Hydrogen Bond	HHB	Arg135
		4.46708	Electrostatic	P-C	Lys131z
		3.7492	Hydrophobic	PPS	His129
		5.07597	Hydrophobic	PA	His129
		3.87561	Hydrophobic	PA	Lys131
		5.41155	Hydrophobic	PA	Arg135
		4.57043	Hydrophobic	PA	Cys151
		4.46708	Electrostatic	P-C	Lys131z
		3.7492	Hydrophobic	PPS	His129
		3.87561	Hydrophobic	PA	Lys131
		5.41155	Hydrophobic	PA	Arg135
		4.57043	Hydrophobic	PA	Cys151
Resveratrol (R1)	-6.7	2.80703	Hydrogen Bond	ННВ	Arg135
		4.43187	Electrostatic	P-C	Lys131
		3.7267	Hydrophobic	PPS	His129
		3.94196	Hydrophobic	PA	Lys131
		5.39084	Hydrophobic	PA	Arg135
		4.58869	Hydrophobic	PA	Cys151
Resveratrol (R2)	-6.7	2.47259	Hydrogen Bond	ННВ	Tyr85
		4.61343	Electrostatic	P-C	Lys131
		3.8035	Hydrophobic	PPS	Ais129
		4.74585	Hydrophobic	P-P-TS	Ais154
		3.86137	Hydrophobic	PA	Lys131
		5.31935	Hydrophobic	PA	Arg135
		4.59489	Hydrophobic	PA	Cys151

Resveratrol (R3)	-6.4	2.91884	Hydrogen Bond	ННВ	Tyr85
(113)		2.86593	Hydrogen Bond	ННВ	Ser146
		4.55058	Electrostatic	P-C	Lys131
		3.81372	Hydrophobic	PPS	Ais129
		3.7706	Hydrophobic	PA	Lys131
		5.49754	Hydrophobic	PA	Arg135
		4.5229	Hydrophobic	PA	Cys151
Scopoletin (S)	-5.8	3.12433	Hydrogen Bond	ННВ	Arg135
		3.88118	Hydrogen Bond	PDHB	Cys151
		3.74339	Hydrophobic	P-S	Lys131
		5.0608	Hydrophobic	PPS	His129
		3.95133	Hydrophobic	PA	His129
		4.07152	Hydrophobic	PA	His154
		3.77458	Hydrophobic	PA	Lys131
		5.11458	Hydrophobic	PA	Arg135
		4.77899	Hydrophobic	PA	Cys151
Scopoletin (S1)	-5.7	3.14201	Hydrogen Bond	ННВ	Arg135
		3.87395	Hydrogen Bond	PDHB	Cys151
		3.77804	Hydrophobic	P-S	Lys131
		5.14951	Hydrophobic	PA	Arg135
		4.74865	Hydrophobic	PA	Cys151
		4.02594	Hydrophobic	PA	His154
		3.77424	Hydrophobic	PA	Lys131
Scopoletin (S2)	-5.8	3.14201	Hydrogen Bond	ННВ	Arg135
		3.87395	Hydrogen Bond	PDHB	Ays151
		3.77804	Hydrophobic	P-S	Lys131
		5.0017	Hydrophobic	PPS	His129
		3.98552	Hydrophobic	PA	His129
		4.02594	Hydrophobic	PA	His154
		3.77424	Hydrophobic	PA	Lys131
		5.14951	Hydrophobic	PA	Arg135
		4.74865	Hydrophobic	PA	Ays151
Resveratrol (R4)	-7.3	3.12891	Hydrogen Bond	ННВ	Gln86
		3.25628	Hydrogen Bond	HHB	Lys131
		3.22821	Hydrogen Bond	ННВ	Arg135
		4.01622	Hydrogen Bond; Electrostatic	P-C; PDHB	Lys131

		3.85983	Hydrogen Bond	PDHB	Ays151
		4.19043	Hydrophobic	PPS	His129
		5.08778	Hydrophobic	P-P-TS	His154
		4.06628	Hydrophobic	A	Lys131
		4.10535	Hydrophobic	A	Arg135
		5.14967	Hydrophobic	A	Val155
		4.18138	Hydrophobic	A	Ays151
		5.3425	Hydrophobic	PA	Tyr85
		3.97664	Hydrophobic	PA	Lys131
		3.12891	Hydrogen Bond; Halogen	CHB; X(F)	Gln86
		3.25628	Hydrogen Bond; Halogen	CHB; X(F)	Lys131
		3.22821	Hydrogen Bond; Halogen	CHB; X(F)	Arg135
Loliolide (L5)	-4.4	3.19794	Hydrogen Bond	ННВ	His129
		3.1168	Hydrogen Bond	HHB	Cys151
		2.65803	Hydrogen Bond	PDHB	His154
		3.86608	Hydrophobic	P-S	His129
		3.86608 3.89742	Hydrophobic Hydrophobic	P-S A	His129 Lys131
			• •		
		3.89742	Hydrophobic	A	Lys131
		3.89742 3.72478	Hydrophobic Hydrophobic Hydrogen Bond;	A PA	Lys131 His129
		3.89742 3.72478 3.19794	Hydrophobic Hydrophobic Hydrogen Bond; Halogen Hydrogen Bond;	A PA CHB; X(F)	Lys131 His129 His129
		3.89742 3.72478 3.19794 3.1168	Hydrophobic Hydrophobic Hydrogen Bond; Halogen Hydrogen Bond; Halogen	A PA CHB; X(F) CHB; X(F)	Lys131 His129 His129 Cys151
		3.89742 3.72478 3.19794 3.1168 3.63923	Hydrophobic Hydrophobic Hydrogen Bond; Halogen Hydrogen Bond; Halogen Halogen	A PA CHB; X(F) CHB; X(F) X(F)	Lys131 His129 His129 Cys151 Lys131
		3.89742 3.72478 3.19794 3.1168 3.63923 2.65803	Hydrophobic Hydrophobic Hydrogen Bond; Halogen Hydrogen Bond; Halogen Halogen Hydrogen Bond	A PA CHB; X(F) CHB; X(F) X(F) PDHB	Lys131 His129 His129 Cys151 Lys131 His154

CHB: Carbon Hydrogen Bond; PDHB: Pi-Donor Hydrogen Bond; PPS: Pi-Pi Stacked; A: Alkyl; PA: Pi-Alkyl; HHB: Conventional Hydrogen Bond; P-HB;P-SB: Pi-Donor Hydrogen Bond; Pi-Sulphur; P-C: Pi-Cation; P-C;P-HB: Pi-Cation; Pi-Donor Hydrogen Bond; P-P-TS: Pi-Pi T-shaped; P-S: Pi-Sigma; CHB;X(F): Conventional Hydrogen Bond; Halogen (Fluorine); X(F): Halogen (Fluorine).

The R4 (-8 kcal/mol) Bis(trifluoromethyl)benzene moiety displayed numerous interactions with the active site residues of the Kelch domain. Noteworthy among them were the various types of bonds formed with Arginine 415 and Serine 508 residues deep in the kelch pore of the keap1. Jnoff et al., while depositing the protein structure for the kelch domain of the keap1 protein (PDB: 4L7B), have presented a comparative interaction analysis study between the docking results and their actual co-crystallized (Nrf2-Keap1 complex) inhibitor, in the kelch domain of the keap1 protein. In that study, the prominence of the binding induced inhibition facilitated by their novel inhibitor interaction with the Arginine 415 and Serine 508 residues in the kelch pore of the Keap1 protein is elucidated (Jnoff et al. 2014). With the strong interactions displayed by the R4 bis(trifluoromethyl)benzene moiety with the Arginine 415, Serine 508 and the surrounding residues that line the kelch pore, a scenario can be envisaged where the R4 bis(trifluoromethyl)benzene moiety occludes the kelch pore, thereby rendering the Keap1 protein incapable of complex formation with the Nrf2 protein. Meanwhile, the trifluoromethyl-benzene moiety past the double unsaturated hinge region was found to interact with Phenylalanine 577 and Tyrosine 572 residues of the kelch domain at the brim of the pore tunnel (Figure 9 (A, B, C)).

Further, the L5 derivative displayed interactions with kelch pore residues like Serine 602, Glycine 364, Asparagine 414, and 382 (Figure 10 (A, B, C)). The importance of these residues haven't been experimentally assessed for their binding mediated keap1 inhibition capability. Hence no definite conclusions were drawn with regard to their individual / synergistic importance for inhibiting Keap1 activity. Furthermore, S1 displayed interactions with Kelch residues like Asparagine 414, Arginine 415, Serine 363, Asparagine 382, Tyrosine 334, and Alanine 556. The hydrophobic interactions displayed by Asparagine 414 and Arginine 415 are noteworthy (Figure 11 (A, B, C)). The inter-residue interactions that did not warrant inclusion were excluded while obtaining pictorial illustrations of the interaction analysis. The complete interaction profile is tabulated in (Table (5)). The remaining pictorial illustrations from PyMol and Discovery studio for the Kelch domain is represented in (Figure S6).

Figure 9: The pictorial illustrations for the Resveratrol-4 derivative in the Kelch domain (PDB: 4L7B), including the surface view of the top-ranked docked pose (A), interaction analysis from Accelrys discovery studio (B) and the same from LigPlot (C).

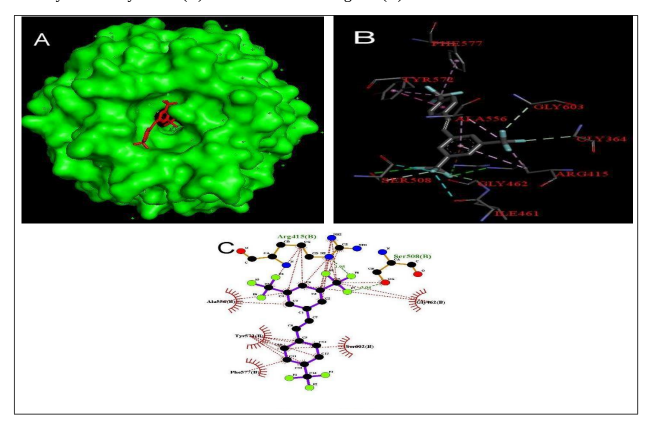
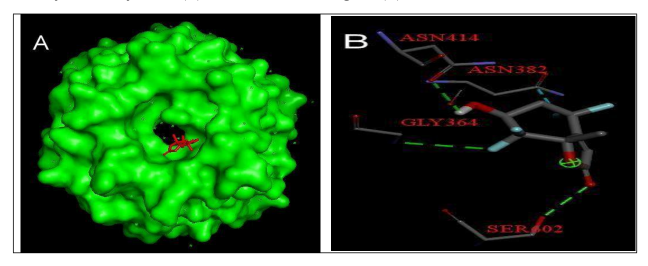


Figure 10: The pictorial illustrations for the Loliolide-5 derivative in the Kelch domain (PDB: 4L7B), including the surface view of the top-ranked docked pose (A), interaction analysis from Accelrys discovery studio (B) and the same from LigPlot (C).



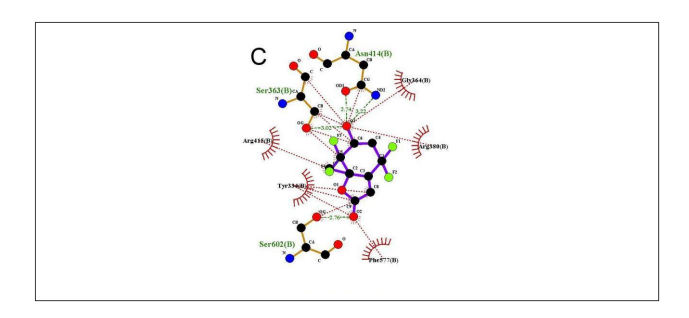
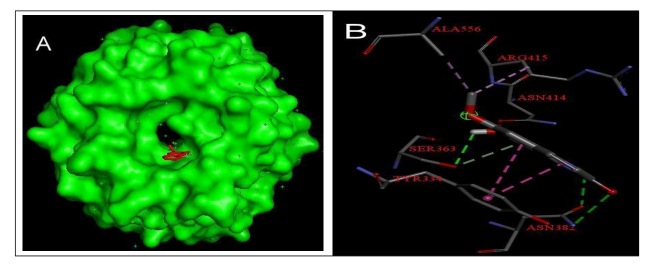


Figure 11: The pictorial illustrations for the Scopoletin-1 derivative in the Kelch domain (PDB: 4L7B), including the surface view of the top-ranked docked pose (A), interaction analysis from Accelrys discovery studio (B) and the same from LigPlot (C).



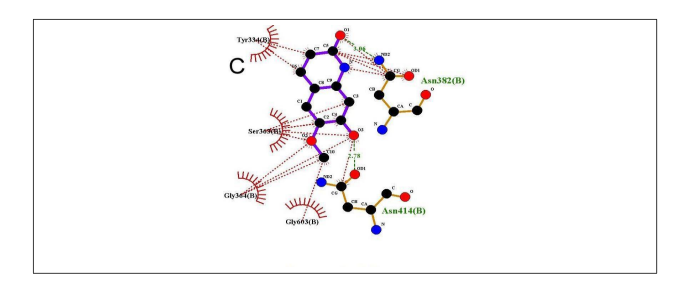


Table 5: Post dock Interaction profile obtained from Accelrys Discovery Studio for the Ligands-Kelch domain interactions.

Name	Binding Affinity (Kcal/mol)	Distance	Bond Category	Interaction type	Interacting Residues
Decussatin	-6.9	3.24218	Hydrogen Bond	ННВ	Arg415
		3.57364	Electrostatic	P-C	Arg415
		3.987	Hydrophobic	P-S	Tyr572
		3.95953	Hydrophobic	A	Arg415
		3.97514	Hydrophobic	A	Ala556
		5.01244	Hydrophobic	A	Ile461
		4.96178	Hydrophobic	PA	Ala556
		5.09825	Hydrophobic	PA	Ala556
		4.95345	Hydrophobic	PA	Phe577
Loliolide (L)	-6.2	3.0705	Hydrogen Bond	ННВ	Ser363g
		2.98659	Hydrogen Bond	ННВ	Ser363g
		5.1654	Hydrophobic	PA	Tyr334
		5.38432	Hydrophobic	PA	Tyr334
		4.56214	Hydrophobic	PA	Tyr572
		5.49526	Hydrophobic	PA	Phe577
Loliolide (L1)	-6.3	2.86297	Hydrogen Bond	ННВ	Ser363
, ,		2.75771	Hydrogen Bond	HHB	Ser602
		2.5552	Hydrogen Bond	HHB	Ser363
		2.08997	Hydrogen Bond	HHB	Asn414
		5.41112	Hydrophobic	A	Ala556
		3.51456	Hydrophobic	A	Ala556

		4.4082	Hydrophobic	A	Ala556
T -10-104-		4.58934	Hydrophobic	Α	Arg415
Loliolide (L2)	-6.1	3.14792	Hydrogen Bond	ННВ	Ser363
()		3.19834	Hydrogen Bond	ННВ	Arg415
		2.4382	Hydrogen Bond	HHB	Ser363
		4.98131	Hydrophobic	A	Ala556
		4.50989	Hydrophobic	PA	Tyr572
Loliolide (L3)	-6.1	2.13026	Hydrogen Bond	ННВ	Ser363
(- /		5.2945	Hydrophobic	PA	Tyr334
		4.0761	Hydrophobic	PA	Tyr334
Loliolide (L4)	-6.4	2.87467	Hydrogen Bond	ННВ	Ser363
,		3.25252	Hydrogen Bond	ННВ	Asn414
		3.03981	Hydrogen Bond	HHB	Ser602
		5.20126	Hydrophobic	A	Ala556
		3.89607	Hydrophobic	A	Ala556
		4.34819	Hydrophobic	A	Ala556
		4.26968	Hydrophobic	A	Arg415
		5.05972	Hydrophobic	A	Arg415
Loliolide (L5)	-6.8	3.31077	Hydrogen Bond	ННВ	Gly364
(- /		2.75642	Hydrogen Bond	ННВ	Ser602
		2.55491	Hydrogen Bond	HHB	Asn414
		3.31077	Hydrogen Bond	HHB	Gly364
		2.75642	Hydrogen Bond	HHB	Ser602
		2.55491	Hydrogen Bond	HHB	Asn414
		3.31077	Hydrogen Bond; Halogen	HHB; $X(F)$	Gly364
		2.75642	Hydrogen Bond	ННВ	Ser602
		2.55491	Hydrogen Bond	ННВ	Asn414
		3.26649	Halogen	X(F)	Asn382
Resveratrol (R)	-6.5	2.79219	Hydrogen Bond	ННВ	Ser363
()		1.98037	Hydrogen Bond	ННВ	Asn414
		4.54006	Electrostatic	P-C	Arg415
		4.21464	Electrostatic	P-C	Arg415
		3.8606	Hydrophobic	P-S	Ala556
		5.43894	Hydrophobic	PA	Arg380
Resveratrol (R1)	-6.4	2.84504	Hydrogen Bond	ННВ	Ser363
,		2.13222	Hydrogen Bond	ННВ	Ser508
		1.99622	Hydrogen Bond	HHB	Asn414
			Hydrogen Bond	CHB	

		4.66159 4.34841 3.72588	Electrostatic Electrostatic Hydrophobic	P-C P-C P-S	Arg415 Arg415 Ala556
Resveratrol		5.47801	Hydrophobic	PA	Arg380
(R2)	-6.4	2.15698	Hydrogen Bond	HHB	Ser363
, ,		2.06752	Hydrogen Bond	HHB	Ser555
		4.07268	Electrostatic	P-C	Arg415
		5.63874	Hydrophobic	P-P-TS	Tyr334
		4.69447	Hydrophobic	PA	Ala556
Resveratrol (R3)	-6.5	2.31515	Hydrogen Bond	ННВ	Ser363
· - /		2.37749	Hydrogen Bond	HHB	Ser555
		4.15495	Electrostatic	P-C	Arg415
		5.75362	Hydrophobic	P-P-TS	Tyr334
		4.6338	Hydrophobic	PA	Ala556
Resveratrol (R4)	-8	2.95147	Hydrogen Bond	ННВ	Arg415
,		3.15531	Hydrogen Bond	HHB	Arg415
		3.03519	Hydrogen Bond	HHB	Ser508
		2.94774	Hydrogen Bond	CHB	Gly364
		3.07229	Hydrogen Bond	CHB	Gly462
		3.12396	Hydrogen Bond	CHB	Ser508
		3.58099	Hydrogen Bond	CHB	Gly603
		3.94395	Electrostatic	P-C	Arg415
		3.69051	Hydrophobic	PPS	Tyr572
		3.92247	Hydrophobic	A	Ala556
		3.72052	Hydrophobic	A	Arg415
		4.61022	Hydrophobic	PA	Tyr572
		5.21239	Hydrophobic	PA	Phe577
		4.54073	Hydrophobic	PA	Arg415
		4.10525	Hydrophobic	PA	Ala556
Scopoletin (S)	-6.4	3.16699	Hydrogen Bond	ННВ	Asn382
` ,		2.62422	Hydrogen Bond	ННВ	Ser363
		3.55323	Hydrogen Bond	PDHB	Ser363
		5.12251	Hydrophobic	PPS	Tyr334
		4.00382	Hydrophobic	PPS	Tyr334
		4.03552	Hydrophobic	A	Ala556
		4.61713	Hydrophobic	A	Arg415
Scopoletin (S1)	-6.5	3.05648	Hydrogen Bond	ННВ	Asn382
• •		2.49493	Hydrogen Bond	HHB	Asn382
		2.7133	Hydrogen Bond	HHB	Ser363
		3.63098	Hydrogen Bond	PDHB	Ser363

		4.09363	Hydrophobic	PPS	Tyr334
		5.10585	Hydrophobic	PPS	Tyr334
		3.97733	Hydrophobic	A	Ala556
		4.85466	Hydrophobic	A	Arg415
Scopoletin (S2)	-6.2	3.58029	Hydrophobic	P-S	Tyr572
		3.8056	Hydrophobic	PPS	Tyr334
		4.47884	Hydrophobic	PPS	Tyr334
		4.94473	Hydrophobic	P-P-TS	Tyr572
		5.1015	Hydrophobic	A	Arg415

CHB: Carbon Hydrogen Bond; PDHB: Pi-Donor Hydrogen Bond; PPS: Pi-Pi Stacked; A: Alkyl; PA: Pi-Alkyl; HHB: Conventional Hydrogen Bond; P-HB;P-SB: Pi-Donor Hydrogen Bond; Pi-Sulphur; P-C: Pi-Cation; P-C;P-HB: Pi-Cation; Pi-Donor Hydrogen Bond; P-P-TS: Pi-Pi T-shaped; P-S: Pi-Sigma; CHB;X(F): Conventional Hydrogen Bond; Halogen (Fluorine); X(F): Halogen (Fluorine).

5 Conclusion

In this investigation, Resveratrol, Decussatin, Loliolide, and Scopoletin were subjected to in silico derivatization by structural modification via sequential replacement of the parent hydroxyl groups with trifluoromethyl and nitro groups resulting in a ligand library. The entire ligand library was subjected to pharmacokinetic analysis (ADME) and was further modified to obtain maximum drug-likeness qualities (Blood-brain-barrier permeability (BBB > 9.5), Human intestinal absorption among others (HIA > 0.7)). Quantum chemical calculations based on density functional theory were performed to optimize the geometries of the constructed library. DFT studies also revealed the electronic properties of the ligand (bond lengths, bond angles, HOMO, LUMO, bandgap, ionization potential, and electron affinity) in a water solvent system set to the pH (7.2) of the human cerebrospinal fluid. R4 (Resveratrol), L5 (Loliolide), and S1 (Scopoletin) derivatives were attributed as superior molecules concerning their potential as an antioxidant/free radical scavenger and also for binding mediated inhibition of the Keap1 protein. Further, molecular docking studies of the ligand library on the Kelch and BTB domains of the Keap1 protein was performed. The post dock comparative interaction analysis suggested remarkable active site residue-ligand bond formations based on which the R4 (Resveratrol) derivative was ranked in the first place (-8 Kcal/mol) as the best multi-domain inhibitor of the Keap1 protein. Overall, it can be concluded that the pharmacokinetic analysis, quantum chemical calculations, and molecular docking studies on the constructed ligand library have produced potent and efficient phytochemical inhibitors (R4, L5, and S1) that can successfully cross the blood-brain barrier and enhance neuronal cytosolic Nrf2 accumulation via Keap1 inhibition. However, the validity of these claims and findings is dependent on in vitro/in vivo experiments, along with the efficiency and reliability of the utilized software. Further, fluoride toxicity is a major issue posed by the novel derivatives presented here (although the rationale for including trifluoromethyl moieties are described previously in this work), but the potency of these candidates seems to be extremely high. Therefore, it can be assumed that small quantities of these novel candidates would possess high

therapeutic potential. Additionally, a hybrid prodrug formulation that would rectify the problem posed by fluoride toxicity can also be a future scope of work.

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7 References

- Albrecht B, Voronina E, Schipke C, Peters O, Parr MK, Díaz-Hernández MD, et al. Pursuing Experimental Reproducibility: An Efficient Protocol for the Preparation of Cerebrospinal Fluid Samples for NMR-Based Metabolomics and Analysis of Sample Degradation. Metabolites [Internet]. 2020 Jun 16 [cited 2020 Jun 28];10(6):251. Available from: https://www.mdpi.com/2218-1989/10/6/251
- Altamura AC, Moro AR, Percudani M. Clinical Pharmacokinetics of Fluoxetine [Internet]. Vol. 26, Clinical Pharmacokinetics. Springer; 1994 [cited 2020 Aug 23]. p. 201–14. Available from: https://link.springer.com/article/10.2165/00003088-199426030-00004
- Barkley RA. Attention-Deficit Hyperactivity Disorder: A HANDBOOK FOR DIAGNOSIS AND TREATMENT [Internet]. 3rd ed. New York: Guilford; 2014 [cited 2020 Jun 25]. Available from: www.guilford.com
- Becke AD. Density-functional exchange-energy approximation with correct asymptotic behavior. Phys Rev A [Internet]. 1988 Sep 1 [cited 2020 Jun 29];38(6):3098–100. Available from: https://journals.aps.org/pra/abstract/10.1103/PhysRevA.38.3098
- Berman HM. The Protein Data Bank / Biopython. Presentation [Internet]. 2000 [cited 2019 Aug 18];28(1):235–42. Available from: http://nar.oxfordjournals.org/content/28/1/235
- Bode BM, Gordon MS. MacMolPlt: A graphical user interface for GAMESS. J Mol Graph Model [Internet]. 1998 [cited 2020 Jun 28];16(3):133–8. Available from: https://pubmed.ncbi.nlm.nih.gov/10434252/
- Canning P, Sorrell FJ, Bullock AN. Structural basis of Keap1 interactions with Nrf2. 2015;
- Cheng F, Li W, Zhou Y, Shen J, Wu Z, Liu G, et al. AdmetSAR: A comprehensive source and free tool for assessment of chemical ADMET properties. J Chem Inf Model. 2012;52(11):3099–105.
- Chovanová Z, Muchová J, Sivoňová M, Dvořáková M, Žitňanová I, Waczulíková I, et al. Effect

- of polyphenolic extract, Pycnogenol®, on the level of 8-oxoguanine in children suffering from attention deficit/hyperactivity disorder. Free Radic Res. 2006;40(9):1003–10.
- Cousins KR. Computer review of ChemDraw ultra 12.0. Vol. 133, Journal of the American Chemical Society. 2011. p. 8388.
- Daina A, Michielin O, Zoete V. SwissADME: A free web tool to evaluate pharmacokinetics, druglikeness and medicinal chemistry friendliness of small molecules. Vol. 7, Scientific Reports. Nature Publishing Group; 2017.
- Dallakyan S, Olson AJ. Small-molecule library screening by docking with PyRx. Methods Mol Biol [Internet]. 2015 [cited 2020 Jun 29];1263:243–50. Available from: https://pubmed.ncbi.nlm.nih.gov/25618350/
- Davis IW, Leaver-Fay A, Chen VB, Block JN, Kapral GJ, Wang X, et al. MolProbity: All-atom contacts and structure validation for proteins and nucleic acids. Nucleic Acids Res [Internet]. 2007 [cited 2020 Jun 29];35(SUPPL.2). Available from: https://academic.oup.com/nar/article-abstract/35/suppl_2/W375/2920742
- Di L. Strategic Approaches to Optimizing Peptide ADME Properties. AAPS J. 2015;17(1):134–43.
- Gordaliza M. Natural products as leads to anticancer drugs. Clin Transl Oncol. 2007;9(12):767–76.
- Gordon MS, Schmidt MW. Advances in electronic structure theory: GAMESS a decade later. In: Theory and Applications of Computational Chemistry. Elsevier; 2005. p. 1167–89.
- Gülçin I. Antioxidant properties of resveratrol: A structure-activity insight. Innov Food Sci Emerg Technol. 2010 Jan 1;11(1):210–8.
- Hanwell MD, Curtis DE, Lonie DC, Vandermeerschd T, Zurek E, Hutchison GR. Avogadro: An advanced semantic chemical editor, visualization, and analysis platform. J Cheminform [Internet]. 2012 Aug 13 [cited 2020 Jun 28];4(8):17. Available from: https://jcheminf.biomedcentral.com/articles/10.1186/1758-2946-4-17
- Hashimoto M, Hossain S, Al Mamun A, Matsuzaki K, Arai H. Docosahexaenoic acid: one molecule diverse functions. Crit Rev Biotechnol. 2017;37(5):579–97.
- Hooft RWW, Sander C, Vriend G. Objectively judging the quality of a protein structure from a Ramachandran plot. Vol. 13, CABIOS. 1997.
- Huerta C, Jiang X, Trevino I, Bender CF, Ferguson DA, Probst B, et al. Characterization of novel small-molecule NRF2 activators: Structural and biochemical validation of stereospecific KEAP1 binding. Biochim Biophys Acta Gen Subj [Internet]. 2016;1860(11):2537–52. Available from: http://dx.doi.org/10.1016/j.bbagen.2016.07.026
- Hybertson BM, Gao B, Bose SK, McCord JM. Oxidative stress in health and disease: The therapeutic potential of Nrf2 activation. Mol Aspects Med [Internet]. 2011 Aug [cited 2020 May 23];32(4–6):234–46. Available from: https://linkinghub.elsevier.com/retrieve/pii/S0098299711000501
- Jnoff E, Albrecht C, Barker JJ, Barker O, Beaumont E, Bromidge S, et al. Binding mode and structure-activity relationships around direct inhibitors of the Nrf2-Keap1 complex. ChemMedChem. 2014;9(4):699–705.

- Johnson J, Johnson D, ... AK-A of the new, 2008 undefined. The Nrf2-ARE pathway: an indicator and modulator of oxidative stress in neurodegeneration. ncbi.nlm.nih.gov [Internet]. [cited 2020 Jun 23]; Available from: https://www.ncbi.nlm.nih.gov/pmc/articles/pmc2605641/
- Karsten H, Pratap Singh P, Singh A, Chauhan S, Murat Singh Chauhan S. Activity-guided isolation of antioxidant xanthones from Swertia chirayita (Roxb.) H. Karsten (Gentianaceae). Nat Prod Res [Internet]. 2012 [cited 2020 Jun 27];26(18):1682–6. Available from: http://www.tandfonline.com/action/journalInformation?journalCode=gnpl20
- Kaspar J, Niture S, Medicine AJ-FRB and, 2009 undefined. Nrf2: INrf2 (Keap1) signaling in oxidative stress. Elsevier [Internet]. [cited 2020 Jun 23]; Available from: https://www.sciencedirect.com/science/article/pii/S0891584909004699
- Kim S, Chen J, Cheng T, Gindulyte A, He J, He S, et al. PubChem 2019 update: Improved access to chemical data. Nucleic Acids Res. 2019 Jan 8;47(D1):D1102–9.
- Koopmans T. Über die Zuordnung von Wellenfunktionen und Eigenwerten zu den Einzelnen Elektronen Eines Atoms. Physica. 1934 Jan 1;1(1–6):104–13.
- Laskowski RA, Rullmann JAC, MacArthur MW, Kaptein R, Thornton JM. AQUA and PROCHECK-NMR: Programs for checking the quality of protein structures solved by NMR. J Biomol NMR [Internet]. 1996 [cited 2020 Jul 7];8(4):477–86. Available from: https://pubmed.ncbi.nlm.nih.gov/9008363/
- Laskowski RA, Swindells MB. LigPlot+: Multiple ligand-protein interaction diagrams for drug discovery. J Chem Inf Model [Internet]. 2011 Oct 24 [cited 2020 Jun 30];51(10):2778–86. Available from: https://pubmed.ncbi.nlm.nih.gov/21919503/
- Lauritzen L, Brambilla P, Mazzocchi A, Harsløf LBS, Ciappolino V, Agostoni C. DHA effects in brain development and function [Internet]. Vol. 8, Nutrients. MDPI AG; 2016 [cited 2020 Jun 30]. Available from: /pmc/articles/PMC4728620/?report=abstract
- Lee C, Yang W, Parr RG. Development of the Colle-Salvetti correlation-energy formula into a functional of the electron density. Phys Rev B [Internet]. 1988 Jan 15 [cited 2020 Jun 29];37(2):785–9. Available from: https://journals.aps.org/prb/abstract/10.1103/PhysRevB.37.785
- Lipinski CA. Drug-like properties and the causes of poor solubility and poor permeability. J Pharmacol Toxicol Methods [Internet]. 2000 Jul [cited 2020 Jun 27];44(1):235–49. Available from: https://linkinghub.elsevier.com/retrieve/pii/S1056871900001076
- Liu H, Wang L, Lv M, Pei R, Li P, Pei Z, et al. AlzPlatform: An Alzheimer's disease domain-specific chemogenomics knowledgebase for polypharmacology and target identification research. J Chem Inf Model [Internet]. 2014 [cited 2020 Jun 27];54(4):1050–60. Available from: https://pubs.acs.org/doi/full/10.1021/ci500004h
- Ma Q. Role of Nrf2 in Oxidative Stress and Toxicity. Annu Rev Pharmacol Toxicol. 2013 Jan 6;53(1):401–26.
- Murugavel S, Ravikumar C, Jaabil G, Alagusundaram P. Synthesis, crystal structure analysis, spectral investigations (NMR, FT-IR, UV), DFT calculations, ADMET studies, molecular docking and anticancer activity of 2-(1-benzyl-5-methyl-1H-1,2,3-triazol-4-yl)-4-(2-chlorophenyl)-6-methoxypyridine A novel poten. J Mol Struct [Internet]. 2019;1176:729–42. Available from: https://doi.org/10.1016/j.molstruc.2018.09.010

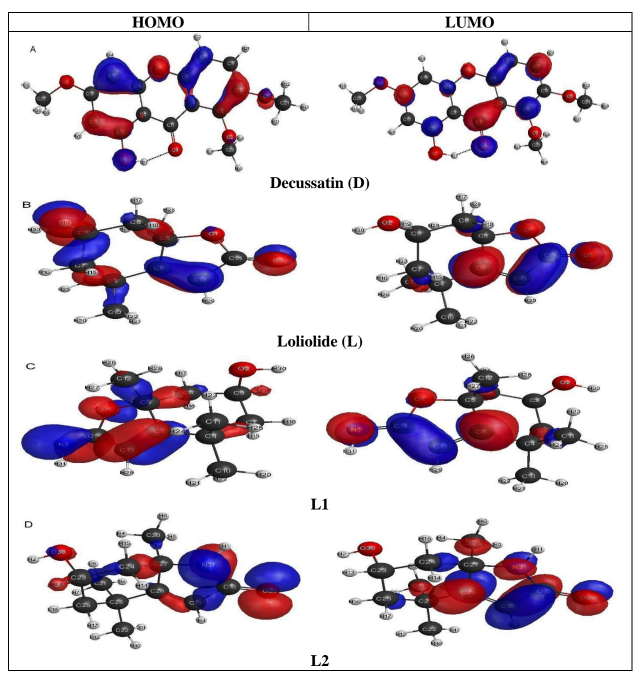
- Napolitano A, Manini P, D'Ischia M. Oxidation Chemistry of Catecholamines and Neuronal Degeneration: An Update. Curr Med Chem. 2011;18(12):1832–45.
- Napolitano A, Pezzella A, Prota G. 6,7-Dihydroxy-1,2,3,4-tetrahydroisoquinoline formation by iron mediated dopamine oxidation: A novel route to endogenous neurotoxins under oxidative stress conditions. Tetrahedron Lett. 1999a;40(14):2833–6.
- Napolitano A, Pezzella A, Prota G. New reaction pathways of dopamine under oxidative stress conditions: Nonenzymatic iron-assisted conversion to norepinephrine and the neurotoxins 6-hydroxydopamine and 6,7-dihydroxytetrahydroisoquinoline. Chem Res Toxicol. 1999b;12(11):1090–7.
- Nelson SK, Bose SK, Grunwald GK, Myhill P, McCord JM. The induction of human superoxide dismutase and catalase in vivo: A fundamentally new approach to antioxidant therapy. Free Radic Biol Med. 2006;40(2):341–7.
- Parr RG, Pearson RG. Absolute Hardness: Companion Parameter to Absolute Electronegativity. J Am Chem Soc [Internet]. 1983 [cited 2020 Jun 29];105(26):7512–6. Available from: https://pubs.acs.org/doi/abs/10.1021/ja00364a005
- Parr RG, Yang W. Density-Functional Theory of the Electronic Structure of Molecules. Annu Rev Phys Chem [Internet]. 1995 Oct [cited 2020 Jun 29];46(1):701–28. Available from: https://pubmed.ncbi.nlm.nih.gov/24341393/
- Pettersen EF, Goddard TD, Huang CC, Couch GS, Greenblatt DM, Meng EC F TE. UCSF Chimera--a visualization system for exploratory research and analysis. J Comput Chem. 2004.
- Ross BM, McKenzie I, Glen I, Bennett CPW. Increased levels of ethane, a non-invasive marker of n-3 fatty acid oxidation, in breath of children with attention deficit hyperactivity disorder. Nutr Neurosci. 2003;6(5):277–81.
- Saito R, Suzuki T, Hiramoto K, Asami S, Naganuma E, Suda H, et al. Characterizations of Three Major Cysteine Sensors of Keap1 in Stress Response. 2016 [cited 2020 Jul 5]; Available from: http://mcb.asm.org/
- Shaw C-Y, Chen C-H, Hsu C-C, Chen C-C, Tsai Y-C. Antioxidant properties of scopoletin isolated from Sinomonium acutum. Phyther Res [Internet]. 2003 Aug 1 [cited 2020 Jun 27];17(7):823–5. Available from: http://doi.wiley.com/10.1002/ptr.1170
- Shukla S, Sumaria CS, Pradeepkumar PI. Exploring chemical modifications for siRNA therapeutics: A structural and functional outlook. ChemMedChem. 2010;5(3):328–49.
- TrottO, Olson AJ. AutoDock Vina: Improving the speed and accuracy of docking with a new scoring function, efficient optimization, and multithreading. J Comput Chem [Internet]. 2009 Jan 30 [cited 2020 Jun 29];31(2):NA-NA. Available from: https://onlinelibrary.wiley.com/doi/full/10.1002/jcc.21334
- Uzzaman M, Mahmud T. Structural modification of aspirin to design a new potential cyclooxygenase (COX-2) inhibitors. Silico Pharmacol [Internet]. 2020;8(1). Available from: https://doi.org/10.1007/s40203-020-0053-0
- Wilde MI, Plosker GL, Benfield P. Fluvoxamine: An Updated Review of its Pharmacology, and Therapeutic Use in Depressive Illness. Drugs [Internet]. 1993 Oct 22 [cited 2020 Aug 23];46(5):895–924. Available from: https://link.springer.com/article/10.2165/00003495-

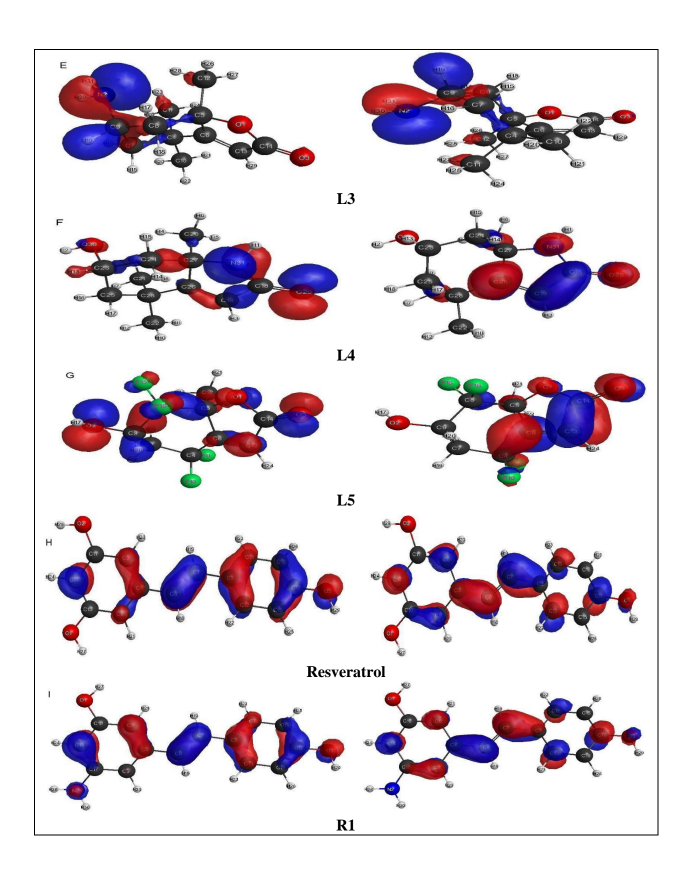
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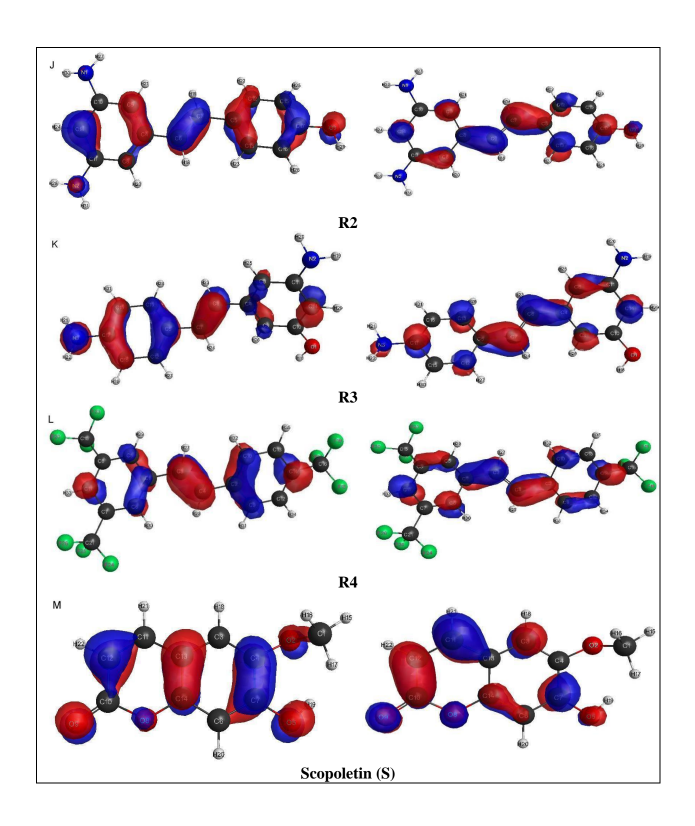
- Yakubenko VP, Byzova T V. Biological and pathophysiological roles of end-products of DHA oxidation. Biochim Biophys Acta Mol Cell Biol Lipids [Internet]. 2017;1862(4):407–15. Available from: http://dx.doi.org/10.1016/j.bbalip.2016.09.022
- Yang X, Kang M-C, Lee K-W, Kang S-M, Lee W-W, Jeon Y-J. Antioxidant activity and cell protective effect of loliolide isolated from Sargassum ringgoldianum subsp. coreanum. Algae [Internet]. 2011 [cited 2020 Jun 27];26(2):201–8. Available from: http://e-algae.kr

Supplementary information

Figure S 1: The color-coded contour maps illustrating the Highest occupied molecular orbitals(HOMO) and the lowest unoccupied molecular orbital (LUMO) of the entire ligand library.







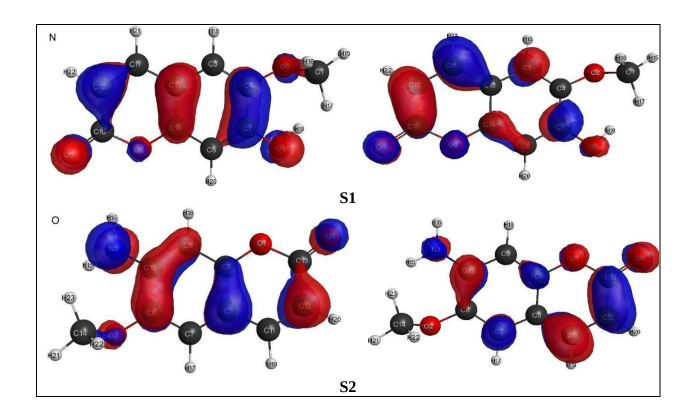
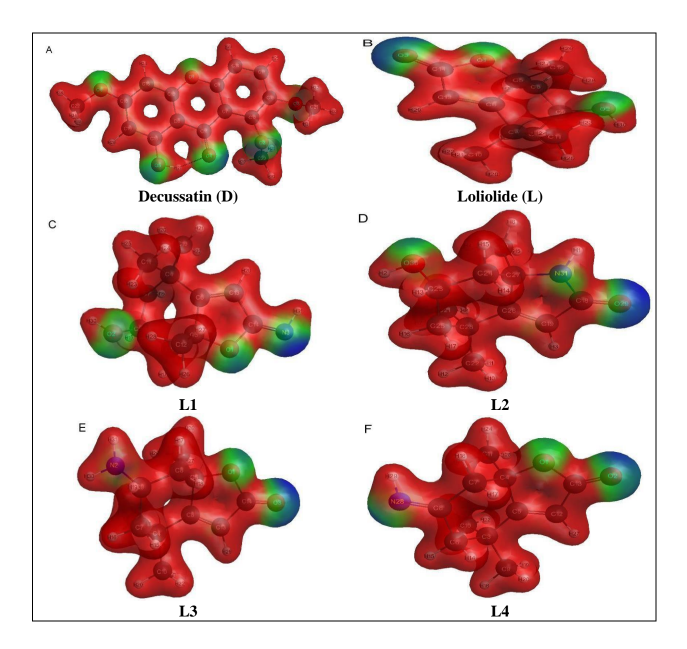
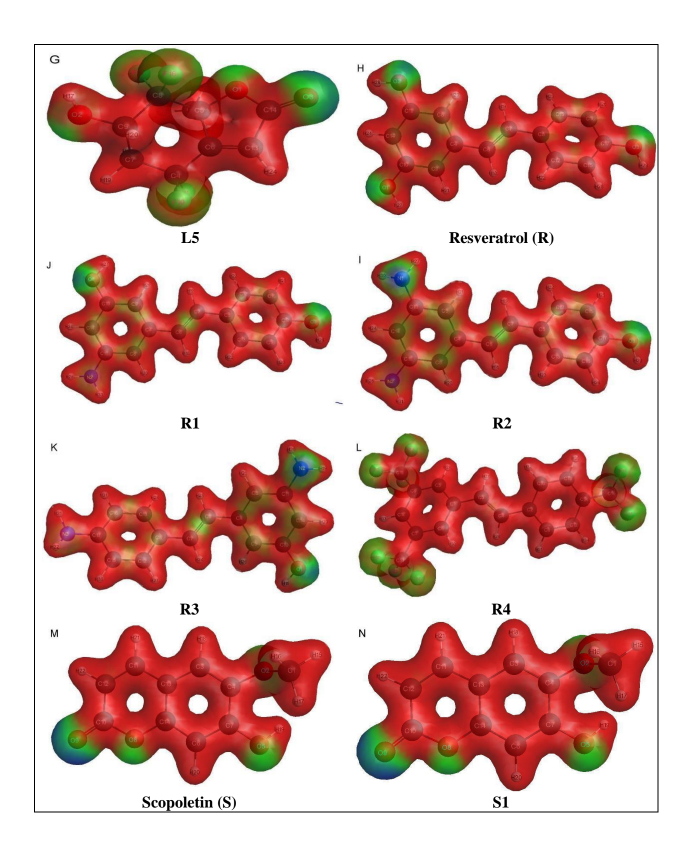


Figure S2: The color-coded contour maps illustrating the variation of molecular electrostatic potentials over the regions of the ligands in the constructed library.





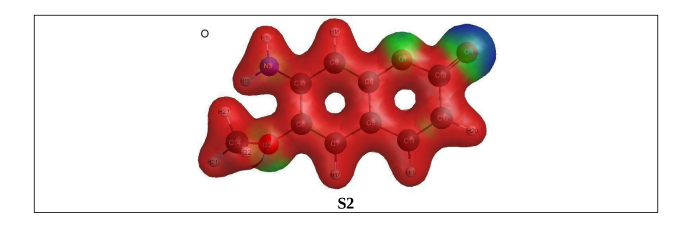
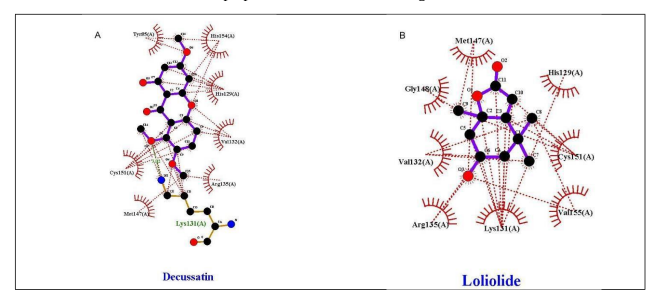
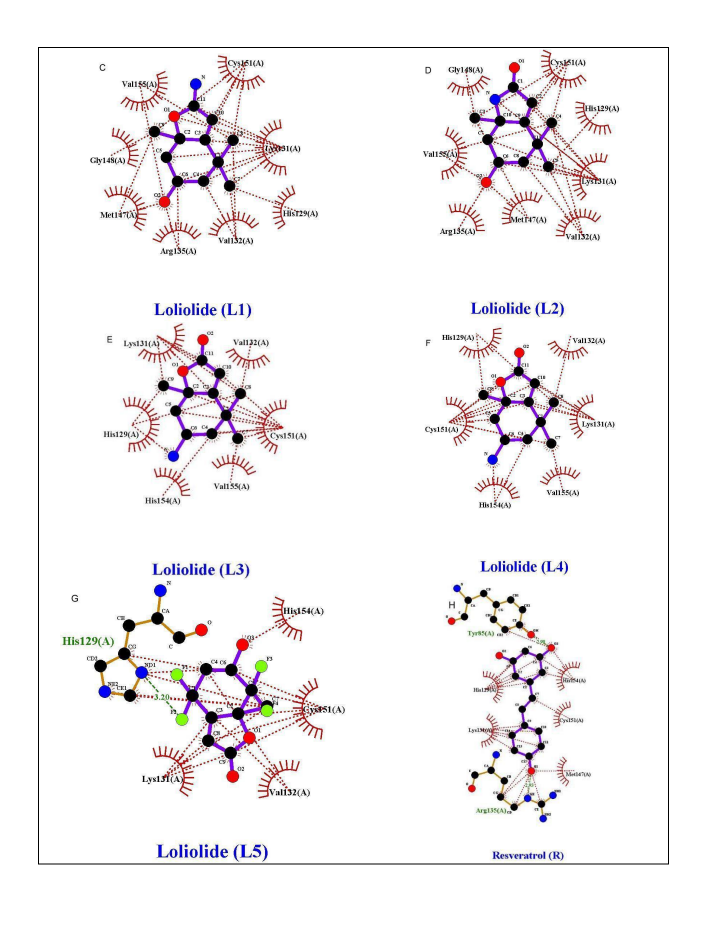
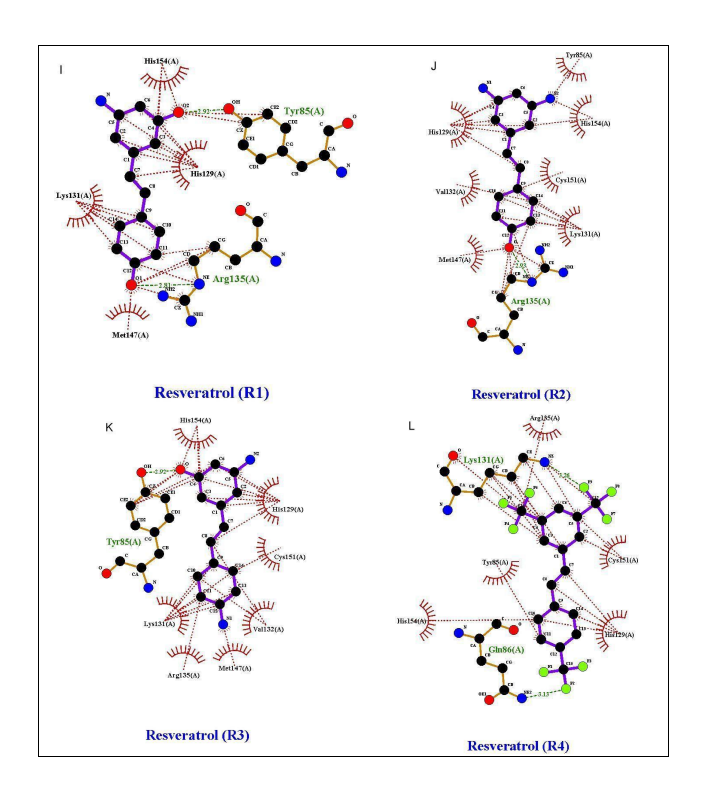


Figure S3: The two-dimensional illustrations of the interaction analysis of the individual ligands with the BTB domain of the Keap1 protein as obtained from LigPlot.







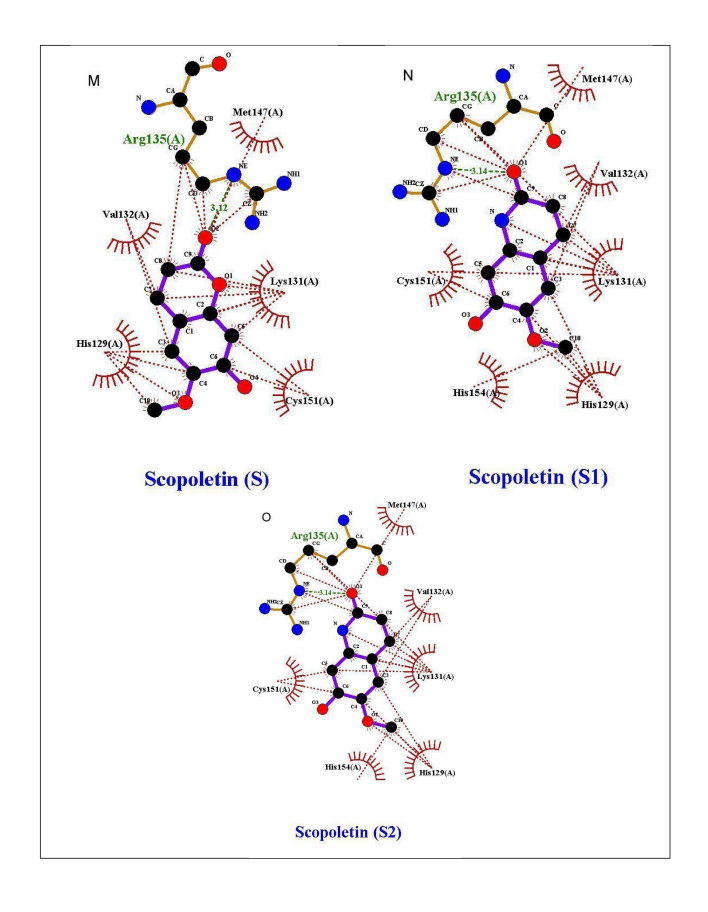
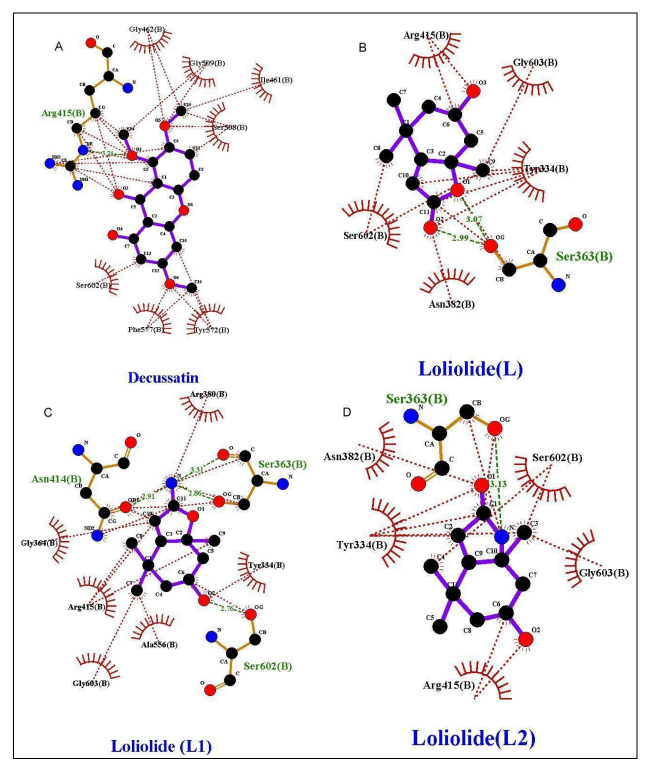
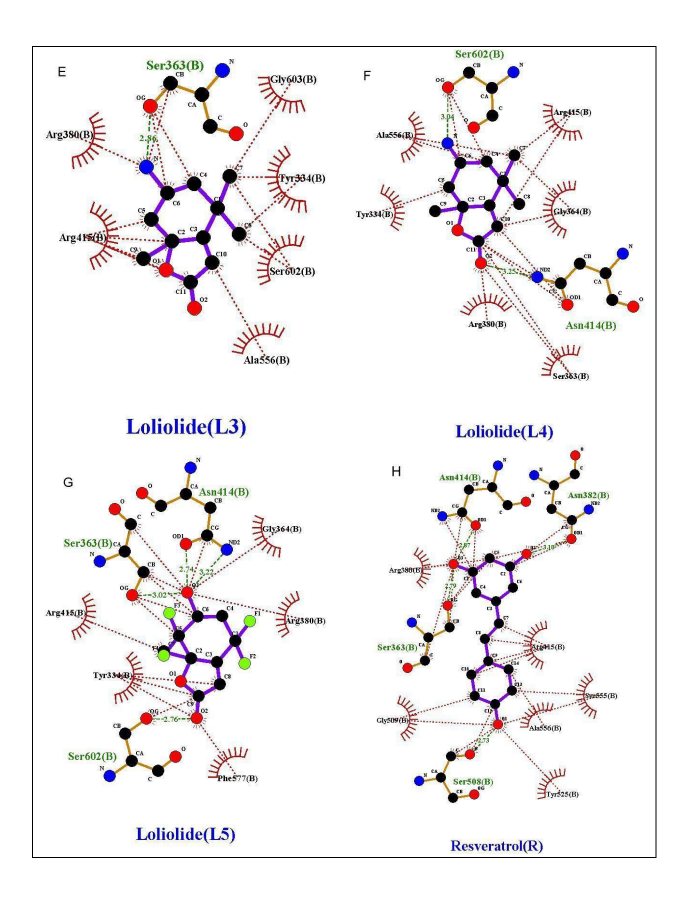
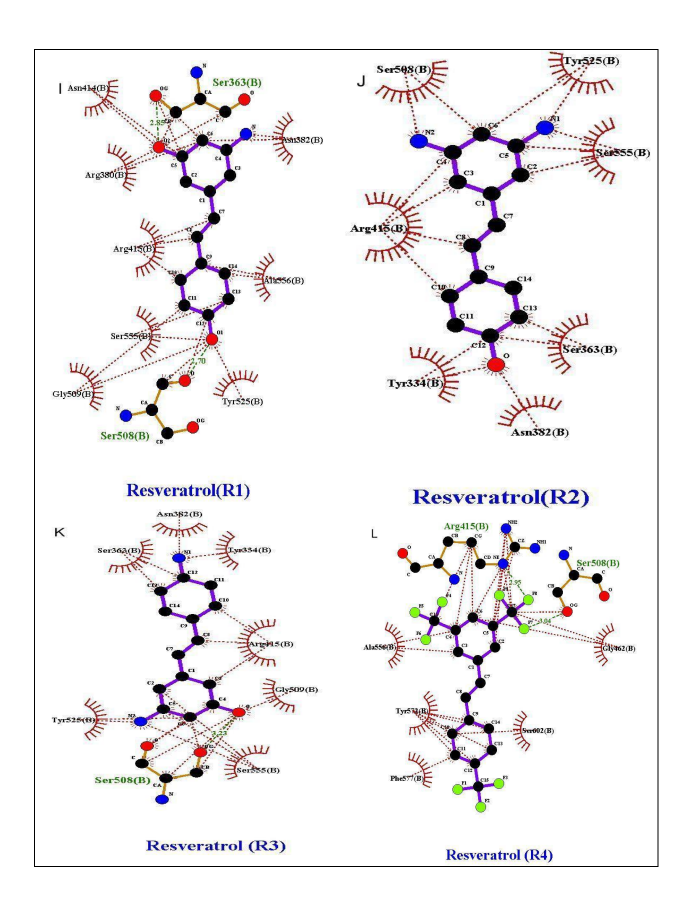


Figure S4: The two-dimensional illustrations of the interaction analysis of the individual ligands with the Kelch domain of the Keap1 protein as obtained from LigPlot.







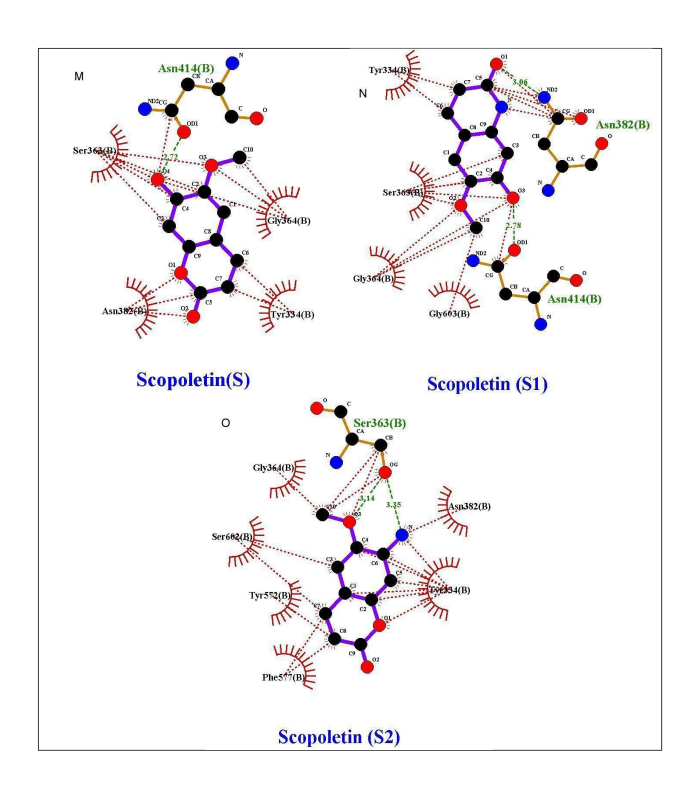
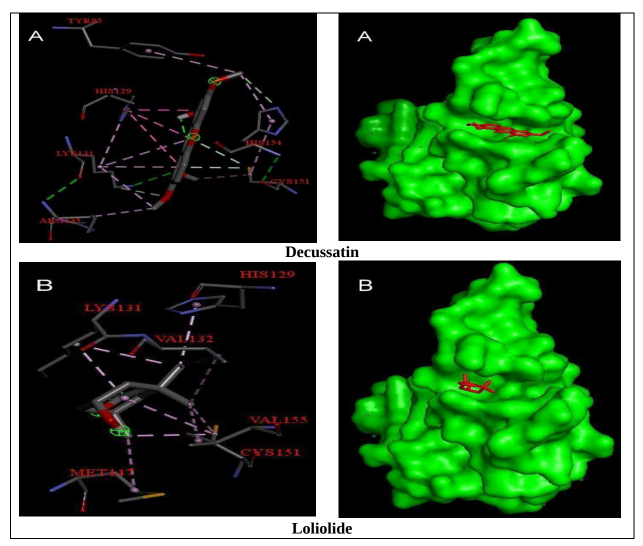
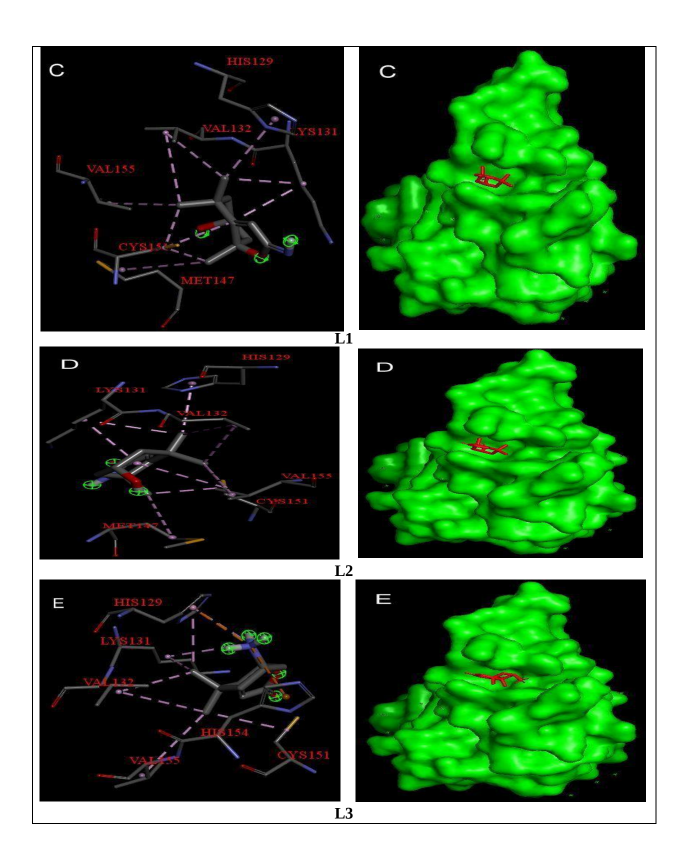
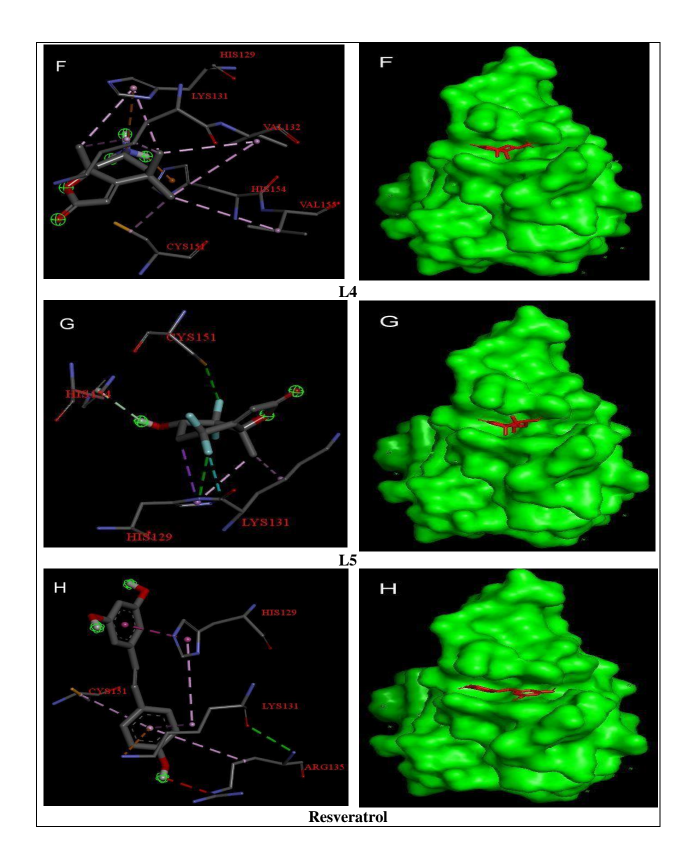
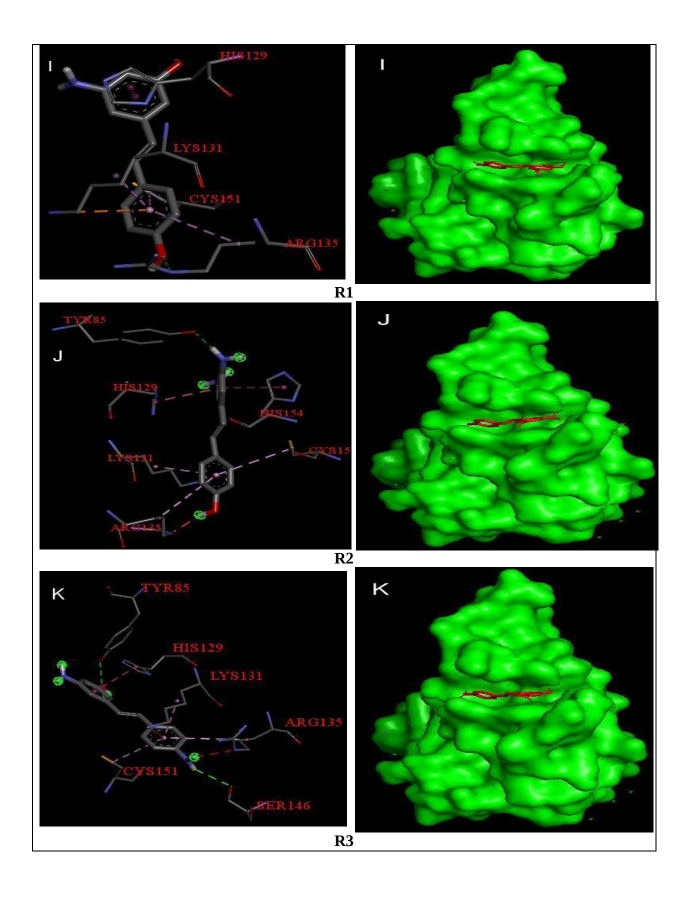


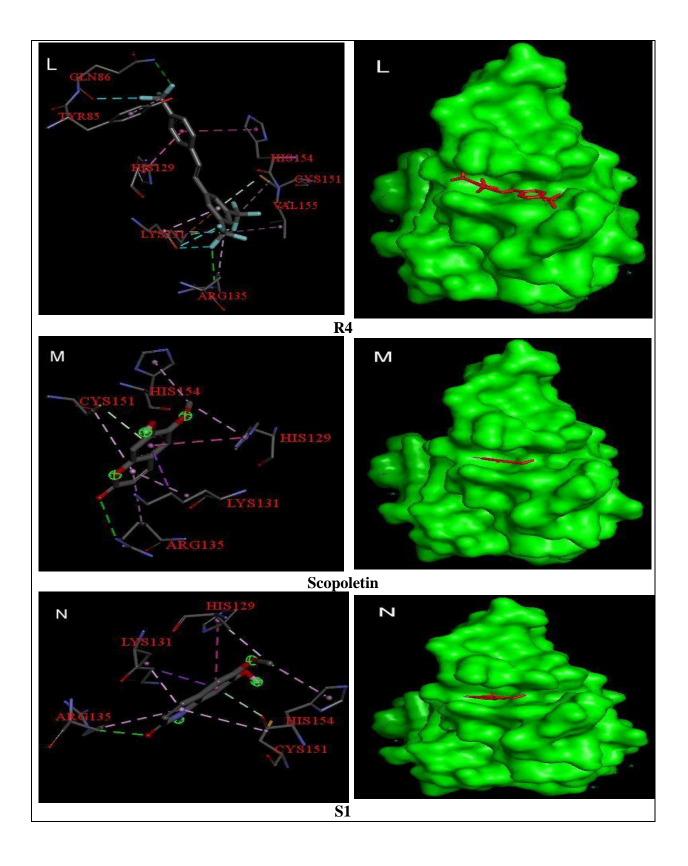
Figure S5: The three-dimensional illustrations of the interaction analysis of the individual ligands with the BTB domain of the Keap1 protein as obtained from the Discovery studio and the surface view of the docked ligand in the BTB active site as obtained from PyMol.











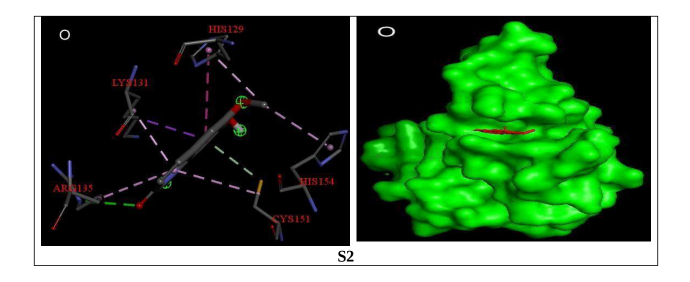


Figure S6: The three-dimensional illustrations of the interaction analysis of the individual ligands with the Kelch domain of the Keap1 protein as obtained from the Discovery studio and the surface view of the docked ligand in the BTB active site as obtained from PyMol.

