

Supplementary material to:

REDOX STATUS, DNA AND HSA BINDING STUDY OF NATURALLY OCCURRING NAPHTHOQUINONE DERIVATIVES

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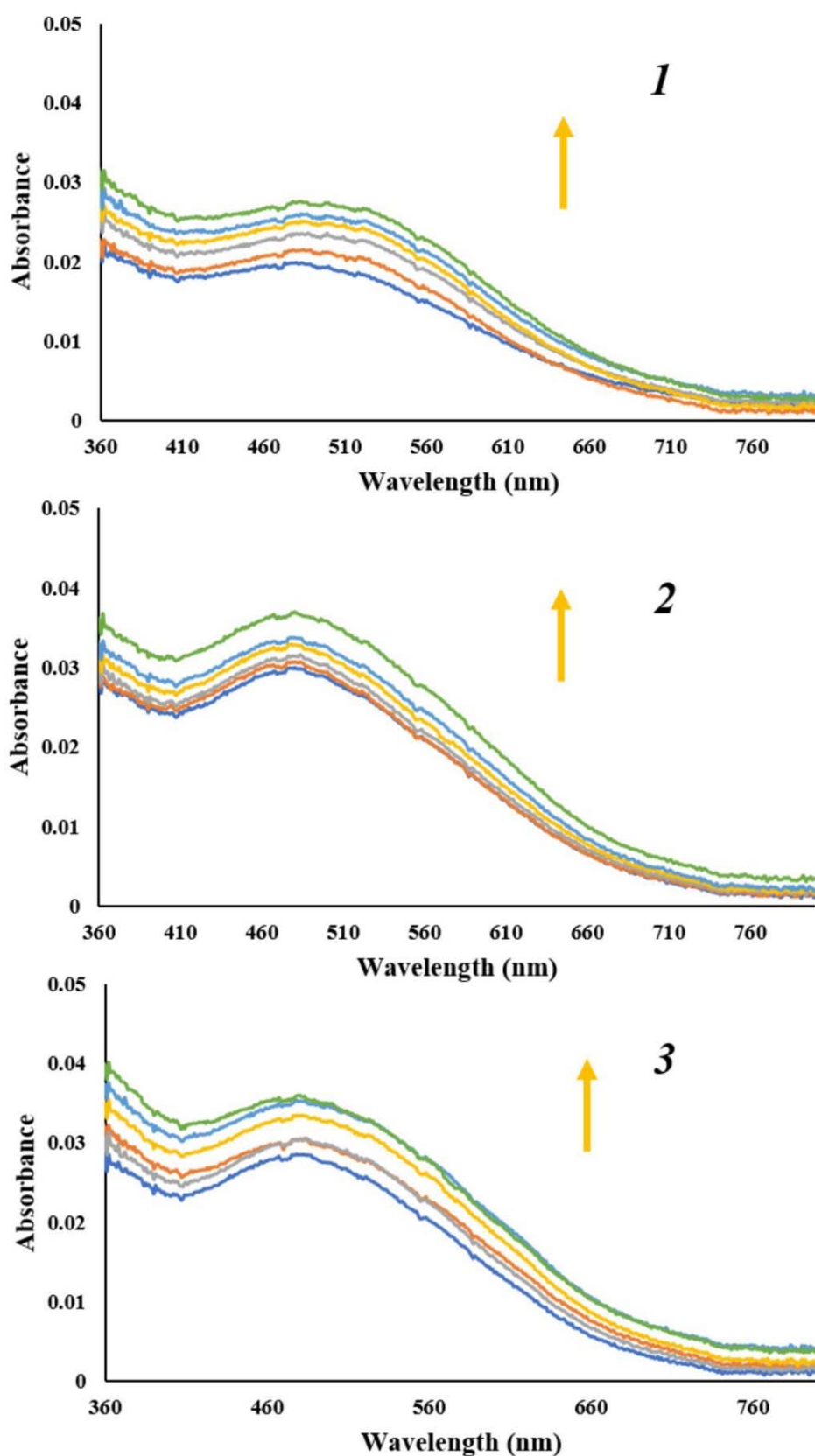
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Supplementary Figure 1: Absorption spectra of α -methylbutyrylshikonin (**1**), acetylshikonin (**2**) and β -hydroxyisovalerylshikonin (**3**) (concentration of each was 8.00×10^{-5} M) before (blue line) and after addition of CT-DNA (0.00 - 1.73×10^{-4} M). Arrow shows the absorbance changes upon increasing concentration of CT-DNA.

Supplementary Table 1: Results of docking analysis of the α -methylbutyrylshikonin (**1**), acetylshikonin (**2**) and β -hydroxyisovalerylshikonin (**3**) with DNA: corresponding purine bases with its position in the structure of DNA, type of interaction during bond formation, distance in Å between respective active sites of naphthoquinones and purine bases, pairwise interaction energy (E_i) in kJ mol^{-1} , (ΔG_{bind}) in kJ mol^{-1} , binding energy, estimated inhibition constant (K_i) in μM .

Complexes	Interaction	Type of interaction	Distance (Å)	E_i (kJ mol^{-1})	ΔG_{bind} (kJ mol^{-1})	K_i (μM)
1-DNA	DA5:H-LIG:O	Conventional Hydrogen Bond	2.08	-0.60	-22.99	93.14
	DA6:H-LIG:O	Conventional Hydrogen Bond	1.87	-0.60		
	DA17:H-LIG:O	Conventional Hydrogen Bond	3.03	-0.05		
	DA16:C -LIG:O	Carbon Hydrogen Bond	3.39	-0.03		
	DA17:C-LIG:O	Carbon Hydrogen Bond	2.04	-0.03		
	DA18:H-LIG	Pi-Donor Hydrogen Bond	2.05	-0.09		
	DA18 -LIG	Pi-Pi T-shaped	5.93	-0.01		
	DA18 -LIG	Pi-Pi T-shaped	5.35	-0.01		
	DT19 -LIG	Pi-Pi T-shaped	5.54	-0.01		
	DA5-LIG	Pi-Alkyl	4.13	-0.02		
	DA6-LIG	Pi-Alkyl	4.84	-0.02		
	DA16-LIG	Pi-Alkyl	4.54	-0.02		
2-DNA	DA5:H-LIG:O	Conventional Hydrogen Bond	2.56	-0.20	-20.98	209.04
	DA5:H-LIG:O	Conventional Hydrogen Bond	2.15	-0.60		
	DA18:H-LIG:O	Conventional Hydrogen Bond	2.65	-0.20		
	DA18:H-LIG:O	Conventional Hydrogen Bond	1.84	-0.60		
	DT19 - LIG	Pi-Pi T-shaped	5.73	-0.01		
	DT19 - LIG	Pi-Pi T-shaped	5.90	-0.01		
	DA16 - LIG	Pi-Alkyl	4.02	-0.02		
	DA17 - LIG	Pi-Alkyl	4.52	-0.02		
3-DNA	DA5:H-LIG:O	Conventional Hydrogen Bond	1.82	-0.60	-20.31	274.69
	DA6:H-LIG:O	Conventional Hydrogen Bond	2.19	-0.60		
	DA6:H-LIG:O	Conventional Hydrogen Bond	2.25	-0.60		
	DA17:H-LIG:O	Conventional Hydrogen Bond	2.93	-0.07		
	DA17:H-LIG:O	Conventional Hydrogen Bond	2.03	-0.60		
	DA18:H- LIG:O	Conventional Hydrogen Bond	1.92	-0.60		
	LIG:H - DT7:O4	Conventional Hydrogen Bond	2.20	-0.60		
	DT19 - LIG	Pi-Pi T-shaped	5.55	-0.01		
	DA16 - LIG	Pi-Alkyl	4.17	-0.02		

Supplementary Table 2: Results of docking analysis of the α -methylbutyrylshikonin (**1**), acetylshikonin (**2**) and β -hydroxyisovalerylshikonin (**3**) with HSA: corresponding amino acid with its position in the structure of protein, type of interaction during bond formation, distance in Å between respective active sites of naphthoquinones and amino acids, pairwise interaction energy (E_i) in kJ mol^{-1} , (ΔG_{bind}) in kJ mol^{-1} binding energy, estimated inhibition constant (K_i) in μM .

Complexes	Interaction	Type of interaction	Distance (Å)	E_i (kJ mol^{-1})	ΔG_{bind} (kJ mol^{-1})	K_i (μM)
1-HSA	LYS199:H - LIG:O	Conventional Hydrogen Bond	2.35	-2.00	-13.17	4.91
	ARG222:H - LIG:O	Conventional Hydrogen Bond	3.01	-0.21		
	ARG257:H - LIG:O	Conventional Hydrogen Bond	2.61	-0.08		
	LIG:H - GLU153:O	Conventional Hydrogen Bond	2.05	-2.51		
	LYS195:N - LIG	Pi-Cation	4.22	-0.08		
	LYS195:N - LIG	Pi-Cation	4.11	-0.08		
	ALA291:C - LIG	Pi-Sigma	3.33	-0.17		
	ALA291 - LIG:C	Alkyl	3.74	-0.08		
	LIG:C - ARG257	Alkyl	3.85	-0.08		
	LIG:C - LEU260	Alkyl	4.34	-0.08		
	LIG:C - ARG218	Alkyl	3.84	-0.08		
	LIG:C - ARG218	Alkyl	4.67	-0.08		
	TRP214 - LIG1:C	Pi-Alkyl	4.47	-0.08		
	LIG - LYS195	Pi-Alkyl	4.96	-0.08		
LIG - ALA291	Pi-Alkyl	4.43	-0.08			
2-HSA	ARG222:H - LIG:O	Conventional Hydrogen Bond	2.24	-2.14	-29.08	8.08
	LIG:H - LYS195:O	Conventional Hydrogen Bond	2.57	-1.13		
	TRP214 - LIG	Pi-Pi Stacked	4.32	-0.08		
	TRP214 - LIG	Pi-Pi Stacked	5.24	-0.04		
	TRP214 - LIG	Pi-Pi Stacked	4.03	-0.08		
	TRP214 - LIG	Pi-Pi Stacked	3.90	-0.08		
	LIG:C - ARG218	Alkyl	3.97	-0.08		
	LIG:C - LEU238	Alkyl	4.38	-0.08		
	LIG:C - LEU219	Alkyl	4.21	-0.08		
	LIG - LYS199	Pi-Alkyl	3.74	-0.08		
	LIG - LYS199	Pi-Alkyl	4.44	-0.08		
3-HSA	LIG:H - GLU292:O	Conventional Hydrogen Bond	1.93	-2.51	-7.74	44.21
	LIG:H - ASP451:O	Conventional Hydrogen Bond	1.97	-2.51		
	ALA191:C - LIG	Pi-Sigma	3.71	-0.08		
	ALA191:C - LIG	Pi-Sigma	3.28	-0.08		
	TYR452 - LIG	Pi-Pi T-shaped	4.40	-0.08		
	ALA191 - LIG:C	Alkyl	3.47	-0.08		