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Affinity of estrogens for human progesterone receptor A and B monomers and risk of breast cancer: a comparative molecular modeling study

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Correspondence: Tarique N Hasan Molecular Cancer Biology Research Laboratory, Department of Food Science and Nutrition, College of Food and Agricultural Sciences, King Saud University, Riyadh 11451, Saudi Arabia Tel +96 654 564 8874 Fax +96 61 467 7122 Email tariquenh@gmail.com **Background:** The human progesterone receptor (hPR) belongs to the steroid receptor family. It may be found as monomers (A and B) and or as a dimer (AB). hPR is regarded as the prognostic biomarker for breast cancer. In a cellular dimer system, AB is the dominant species in most cases. However, when a cell coexpresses all three isoforms of hPR, the complexity of the action of this receptor increases. For example, hPR A suppresses the activity of hPR B, and the ratio of hPR A to hPR B may determine the physiology of a breast tumor. Also, persistent exposure of hPRs to nonendogenous ligands is a common risk factor for breast cancer. Hence we aimed to study progesterone and some nonendogenous ligand interactions with hPRs and their molecular docking.

Methods and results: A pool of steroid derivatives, namely, progesterone, cholesterol, testosterone, testolectone, estradiol, estrone, norethindrone, exemestane, and norgestrel, was used for this in silico study. Dockings were performed on AutoDock 4.2. We found that estrogens, including estradiol and estrone, had a higher affinity for hPR A and B monomers in comparison with the dimer, hPR AB, and that of the endogenous progesterone ligand. hPR A had a higher affinity to all the docked ligands than hPR B.

Conclusion: This study suggests that the exposure of estrogens to hPR A as well as hPR B, and more particularly to hPR A alone, is a risk factor for breast cancer.

Keywords: human progesterone receptor, breast cancer, steroid derivatives, estrogens, molecular docking

Introduction

The human progesterone receptor (hPR), like other members of the steroid receptor family of proteins, is a ligand-induced transcription factor which mediates the effects of progesterone.^{1–3} Progesterone is well known for its critical role in regulation of the normal physiology of the ovary, uterus, mammary gland, as well as brain development during childhood. Progesterone also plays an important role in the maintenance of the cardiovascular, central nervous, and skeletal systems.^{4–7} The importance of the hPR can therefore be understood from its extensive involvement in human physiological processes.

hPR is comprised of specific functional domains, including the central DNA-binding domain and a carboxyl-terminal ligand-binding domain. The central DNA-binding and ligand-binding domains are the sites of hPR activity. In addition to these domains, hPR have many elements with activator and inhibitory functions enhancing and repressing the transcriptional activation of hPR by their interaction with different transcriptional coregulators.^{3,8–12} Cytoplasmic hPR have been found as multiprotein chaperone

complexes which facilitate their inactive conformation, whereby a ligand can bind with hPR.^{13,14}

Two distinct isoforms (hPR A and hPR B) of hPR have been reported previously, and differ by an additional amino acid at position 164 in hPR B. These differences arise as a result of either alternate initiation of translation by the same mRNA or by transcription from alternate promoters within the same gene.¹⁵

Our understanding of the mechanisms underlying the different activities of these two isoforms of human hPR is limited. However, structural and functional studies indicate that the hPR B isoform contains an additional domain, AF3, that accounts for the transcriptional activity of hPR B by suppressing the activity of an inhibitory domain contained within the sequences common to hPR A and hPR B.^{11,16,17} The evidence suggests that the two isoforms acquire different conformations within the cell, and hence interact with distinct coregulators, ie, the coregulators of hPR A are different from those of hPR B.^{18–20}

Much of our knowledge regarding the structure and function of hPR A and hPR B comes from models of single hPR isoform expression and models where the hPR homodimer is the dominant molecular species. However, cell types which coexpress the hPR isoforms may possibly have all three kinds of molecules, ie, hPR A, hPR B, and hPR AB, which increases the complexity of the action of hPR.

hPR is a ligand-activated receptor, and its activation/ deactivation is associated with the pathogenesis of many diseases, including breast cancer.^{21,22} Hence, the specificity of hPR with regard to interaction with its ligand (progesterone) is an important area of research. It has been reported that prolonged exposure to certain nonendogenous ligands, particularly estrogens (which are steroid derivatives) is the strongest risk factor for breast cancer.23,24 In the absence of progesterone, the hPR A:hPR B ratio influences the biology of estrogen receptor-positive tumors and their response to treatment, and hPR A isoforms are functionally dominant in progesterone-deficient states. This explains why PR A-rich tumors are particularly aggressive.²⁵ We hypothesized that hPR A, hPR B, and hPR AB have different affinities for different ligands, based on conditions such as the availability of progesterone and the concentration of hPR monomers and dimers, which may be a possible cause for the development of breast cancer. The aim of our study was to test this hypothesis and find the affinities of the different steroid derivatives for hPRs. Hence, other steroids that can trigger the development of breast cancer by forming a ligand receptor complex with hPR can be predicted. A battery of steroid derivatives,

namely, progesterone (DB00396), cholesterol (DB04540), testosterone (DB00624), testolectone (DB00894), estradiol (DB00783), estrone (DB00655), norethindrone (DB00717), exemestane (DB00990), and norgestrel (DB00506), was used for this in silico study (Figure 1).

Methods and materials Software, data sources, and experiments

All the software used in this study is freely available for academic use. Table 1 provides the source of the data, and Table 2 lists the software and online servers used. The Protein Data Bank (www.rcsb.org) is a World Wide Web repository where data are processed and distributed in the form of three-dimensional biological macromolecular structures.²⁶ The Protein Data Bank now has more than 20,000 structures determined by x-ray diffraction. Protein structures can be accessed and downloaded using keywords or the Protein Data Bank alphanumeric file name.

The Drug Bank (http://redpoll.pharmacy.ualberta.ca/ drugbank/) database is a unique bioinformatics and cheminformatics resource that combines detailed drug (ie, chemical, pharmacological, pharmaceutical) data with comprehensive drug target information (ie, sequence, structure, pathway).²⁷ The database contains nearly 4300 drug entries, including more than 1000 small molecule drugs approved by the US Food and Drug Administration (FDA), 113 FDA-approved biotechnologic (protein/peptide) drugs, 62 nutraceuticals, and more than 3000 experimental drugs. Additionally, more than 6000 protein (ie, drug target) sequences are linked to these drug entries. Each drug card entry contains more than 80 data fields, with half of the information concerning drug and chemical data and the other half concerning drug target or protein data.

PyMOL is a Python-based visualization program (version1.1r1; Delano Scientific 2006, www.pymol.org).²⁸ PyMOL was used to obtain graphical representations of proteins in the Protein Data Bank and the amino acid residues are in contact with the ligand. PyMOL tools can measure distances as well as identify different amino acid residues of receptors that interact with the ligands. Docking results were also analyzed by PyMOL.

AutoDock 4.2 docking software (Scripps Research Institute, www.scripps.edu) and AutoDockTools was used to perform the docking experiments.^{29–31} AutoDockTools is an accessory program that allows the user to interact with AutoDock from a graphic user interface. AutoDock is a suite of automated docking tools designed to predict how small molecules/ligands bind to a receptor/protein of known



Figure I Structure of ligands (as retrieved from Drug Bank) used for docking study.

three-dimensional structure. AutoDock consists of three separate programs, ie, AutoDock, which docks the ligand to a set of grids describing the target protein, AutoGrid, which precalculates these grids, and AutoTors, which sets up bonds which will be treated as rotatable within the ligand. We calculated the binding energy of the different ligands for hPR A, hPR B, and hPR AB, and the molecular docking process was performed according to the method described

Table I Sources of data used in this study	Table I	Sources	of data	used	in this	study
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Protein or drug data	Source		
Human progesterone receptor	PDB ID 1A28		
Progesterone	DB00396		
Cholesterol	DB04540		
Testosterone	DB00624		
Testolectone	DB00894		
Estradiol	DB00783		
Estrone	DB00655		
Norethindrone	DB00717		
Exemestane	DB00990		
Norgestrel	DB00506		

Abbreviations: PDB, Protein Data Bank; DB, Drug Bank.

by Mukharjee and Majumder.³² The details of the procedure are as follows.

Preparing hPRs and ligand files for AutoDock

The Protein Data Bank files downloaded from the World Wide Web repository are often not perfect for docking studies because of missing hydrogen atoms, presence of multiple molecules, added waters, and related problems. Using the AutoDockTools graphic user interface, the files were prepared as follows.

For the hPRs file, all heteroatoms in the Protein Data Bank files were removed. Three separate Protein Data Bank files were then prepared for hPR A, hPR B, and hPR AB, and saved with an extension of .pdb. All three Protein Data Bank files were first read in AutoDockTools, any added waters removed, and polar hydrogens added. AutoDockTools was then used to check if the molecule had charges and, if not, AutoDockTools determined whether the molecule was a peptide (ie, whether all of the names of its residues appeared

Protein Data Bank file and FASTA format of hPR	RCSB server online	http://www.pdb.org/pdb/explore/explore.do?structureId=1A28
PyMOL image	PyMOL I.IrI	http://download.cnet.com/PyMOL/3000-2054_4-10914845.html
AutoDock	AutoDock v 4.2	The Scripps Research Institute, www.scripps.edu

in the standard set of 20 commonly occurring amino acids). If the molecule was found to be a peptide, Kollman charges were added, otherwise Gasteiger charges were added. Finally, solvation parameters were added, and the files were saved with .pdbqs extension (where q and s represent charge and solvation, respectively).

Ligand files

Protein Data Bank files of ligands obtained from the Drug Bank, were read in AutoDockTools, modified as necessary, and saved using .pdbqs extension. AutoDockTools was then used for automatic calculation of the best root, ie, the fixed portion of the ligand from which rotatable branches sprout. We also determined the rotatable bonds in the ligand, making all amide bonds nonrotatable, and set the number of active torsions to the least number of atoms. The ligand files were then saved with ligand.out.pdbq extension (*q* representing charge).

Grid parameter files

For the calculation of interaction energy, parameters were set to create a grid (ie, a three-dimensional box) capable of enclosing hPRs molecules. The grid volume was large enough to allow the ligand to rotate freely, even with its most fully extended conformation. Grid parameters were stored in a grid parameter file using molecule.gpf extension.

AutoGrid4 program

The AutoGrid4 program was run, creating a map for every type of atom in the ligand. For example, a molecule having carbon, nitrogen, oxygen, and hydrogen maps was created as a molecule.C.map, molecule.N.map, molecule.O.map, and molecule.H.map. These grid maps were saved in ASCII format for readability by AutoDock. AutoGrid also generates a corresponding output of the macromolecular file with the extension molecule.glg.

Docking parameter file

The docking parameter file and other properties defined for the ligand was created. A docking parameter file instructs AutoDock on which ligand to move and the map files to use. AutoDock's search methods include the Monte Carlo simulated annealing method, the genetic algorithm, local search, and the hybrid genetic algorithm with local search (GALS). The GALS is also considered as the Lamarckian genetic algorithm because the offsprings are allowed to inherit the local search adaptations of their parents. GALS was the chosen algorithm for hPR and ligand interaction analysis.

AutoDock program

Finally, the AutoDock program was run from the AutoDock graphic user interface, and the docked ligand files (.dlg extension) were used for this study. The .dlg files were read in Autodock as well as in PyMOL to calculate the binding energies in the docked ligand-protein complexes.

Results

Rigorous docking experiments were done to assess the cross-reactivity of various steroid derivatives (ligands) with hPR monomers and dimers. The ligands, receptors, and their corresponding docking results are listed in Table 3, and receptor-ligand interactions demonstrated in Figures 2, 3, and 4. In these docking experiments, all selected ligands were able to bind with hPR A. Estradiol showed the highest binding affinity (ΔG kcal/mol) at -3.68e+06, followed by estrone, progesterone, and cholesterol (3.38e+06, -3.54e+06, and -2.19e+06 kcal/mol, respectively). Testosterone, testolactone, norethindrone, exemestane, and norgestrel exhibited binding affinities in the range of -8.73to -7.14 kcal/mol. Ligand interactions with hPR B showed lower binding affinity than with hPR A. A similar pattern of binding affinity was found for ligands with hPR B, except for progesterone and cholesterol. Estradiol had the highest binding affinity at -3.64e+6 kcal/mol followed by estrone, cholesterol, and progesterone (3.19e+6, -2.53e+6, and 2.05e+6 kcal/mol, respectively). Testosterone, testolactone, norethindrone, exemestane, and norgestrel exhibited binding affinities in the range of -5.86 to -4.75 kcal/mol. All the ligand interactions with hPR homodimer (AB) had similar binding affinity, except for cholesterol, which had a binding affinity of -1.86e+6 kcal/mol, which was the highest binding affinity among all the ligand interactions with hPR AB. Progesterone had a binding affinity of -10.82 kcal/mol, and the remainder of the ligand binding affinities were in the range of -9.82 to -8.50 kcal/mol.

Using this molecular modeling study, we found that the estrogens, estradiol and estrone, had higher affinity

Table 3 [Docking results o	f steroid derivative	(ligand) n	nolecules at	human progesterone	receptor	chains (A, E	3, and AB, blind d	locking)
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Steroid derivative	hPR A		hPR B		hPR AB		
(ligands)	∆G (kcal/mol)	Figure number	$\Delta \mathbf{G}$ (kcal/mol)	Figure number	$\Delta \mathbf{G}$ (kcal/mol)	Figure number	
Progesterone	- 3.54 e+06	2A	-2.05e+06	3A	-10.82	4A	
Cholesterol	-2.19e+06	2B	-2.53e+06	3B	-1.86e+06	4B	
Testosterone	-7.14	2C	-4.75	3C	-9.39	4C	
Testolactone	-7.99	2D	-5.40	3D	-9.33	4D	
Estradiol	-3.68e+06	2E	-3.64e+06	3E	-8.50	4E	
Estrone	-3.38e+06	2F	-3.19e+06	3F	-9.03	4F	
Norethindrone	-8.73	2G	-5.86	3G	-9.52	4G	
Exemestane	-8.03	2H	-5.41	3H	-9.82	4H	
Norgestrel	-8.30	21	-5.48	31	-9.55	41	

Abbreviation: hPR, human progesterone receptor.

for the hPR monomers A and B. Therefore, we undertook binding site analyses for hPR A, B, and AB, for estradiol and estrone. Table 4 summarizes the amino acid residues involved in the binding of estradiol and estrone with hPR A, hPR B, and AB. Only the HIS 93 residue of hPR A created a single hydrogen bond (H-bond) with estradiol and estrone (bond length 2.0 and 2.3 Å, respectively). In the case of hPR B, estradiol created two H-bonds, one with TRP 78 (2.2 Å) and the other with ARG 89 (2.0 Å), whereas estrone made three H-bonds, two with GLN 48 (2.2 and 2.2 Å) and one with ARG 89 (2.1 Å). Interestingly, hPR AB was not involved in the formation of any H-bond with estradiol (1.7 Å). The rest of the amino acid residues listed in Table 4 were involved in hydrophobic interactions through their side chains.

Discussion

Specificity in biomolecular interactions is determined mainly by noncovalent forces like Coulombic electrostatic forces, hydrogen bonds, London's dispersive forces, and van der Waals forces. In addition, the structure of biomolecules is a function of forces among the constituent structural elements, as well as environmental forces. Binding affinity is inversely proportional to free Gibbs energy. To find the specificity of hPR for different steroid derivatives, monomers and dimers of hPR were studied separately because it had been found in vivo that there is a difference in the binding pattern of



Figure 2 Progesterone (A), cholesterol (B), testosterone (C), testolactone (D), estradiol (E), estrone (F), norethindrone (G), exemestane (H), and norgesterl (I) docked onto the human progesterone receptor A (hPR A, green) in its lowest energy-docked conformation. Ligand molecules are shown in red.

Figure 3 Progesterone (A), cholesterol (B), testosterone (C), testolactone (D), estradiol (E), estrone (F), norethindrone (G), exemestane (H), and norgesterol (I) docked onto the human progesterone receptor A (hPR B green) in its lowest energy-docked conformation. Ligand molecules are shown in red.



Figure 4 Progesterone (A), cholesterol (B), testosterone (C), testolactone (D), estradiol (E), estrone (F), norethindrone (G), exemestane (H), and norgestrel (I) docked onto the human progesterone receptor A (hPR AB green) in its lowest energy-docked conformation. Ligand molecules are shown in red.

ligands to hPR A and B. They adopt distinct conformations in cells, as well as having different signaling responses.^{18–20,33–35} Earlier studies of elephant and hamster uterine progesterone receptors reported that there are many natural and synthetic steroid derivatives which are nonendogenous ligands and have higher relative binding affinity with progesterone receptors

as compared with progesterone.^{36,37} Similarly, progesterone is the endogenous ligand for hPR, but in this in silico study, where there was one-to-one correspondence between hPRs and ligands, as well as their flexible docking, we found that estradiol has more affinity with hPR A than progesterone, and estrone had less affinity with hPR A as compared with progesterone,

Table 4 Amino acid residues involved	n binding of estradiol and estrone with human	progesterone receptors A, B, and AB
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Human progesterone receptors	hPR-A	hPR-A	hPR-B	hPR-B	hPR-AB	hPR-AB
Estrogens	Esterone	Estradiol	Esterone	Estradiol	Esterone	Estradiol
Amino acids constituting	GLU A 18	GLU A 18	LEU B 38	LEU B 38	LYS B 155	HIS A 204
the binding sites	PRO A 19	PRO A 19	LEU B 41	ASN B 42	HIS B 204	ASP A 205*
of estrogens	VAL A 21	ASP A 20	ASN B 42	LEU B 44	ILE B 208	LYS A 208
	ILE A 22	VAL A 21	LEU B 44	GLN B 48	PRO B 243	LEU A 244
	GLN A 48	ILE A 22	GLN B 48**	TRP B 78*	VAL B 248	LYS B208
	SER A 51	GLN A 48	TRP B 78	MET B 79	LYS B 249	ILU B 243
	VAL A 52	SER A 51	MET B 79	MET B 82	LEU B 250	LEU B 244
	LEU A 81	VAL A 52	MET B 82	VAL B 83		MET B 247
	MET A 82	LEU A 81	VAL B 83	LEU B 86		
	TRP A 88	MET A 82	LEU B 88	ARG B 89*		
	ARG A 89	LEU A 86	ARG B 89*	PHE B 101		
	HIS A 93*	TRP A 88	PHE B 101	LEU B 104		
	PHEA 141	ARG A 89	LEU B 120	PHE B 117		
	LYS A 145	HIS A 93*	MET B 124	LEU B 120		
		PHEA 141	LEU B 210	MET B 124		
		LYS A 145	TUR B 213	LEU B 210		
			CYS B 214	TYR B 213		
			PHE B 228	CYS B 214		
			MET B 232	THR B 217		
				MET B 232		

Notes: *Amino acids in the binding sites involved in the formation of a single H-bond with estrogens; **Amino acids in the binding sites involved in the formation of double H-bonds with estrogens.

Abbreviation: hPR, human progesterone receptor.

whereas cholesterol has a higher Gibbs energy value compared with progesterone. In the case of hPR B, estrone, estradiol, and cholesterol molecules had higher affinity with hPR B than did progesterone. In general, the binding energy for the ligands with hPR A was considerably higher than that of hPR B. This supports the view that, in the presence of hPR A and hPR B monomers in a cellular system, hPR A can exert an inhibitory effect on the transactivator behavior of hPR B.^{16,17} Our finding is also in agreement with earlier studies showing that hPR A-rich tumors are more aggressive.²⁵ The higher binding affinity of ligands like estradiol and estrone for hPR A than for hPR B is supported by the previous study which showed that cumulative exposure of estradiols to hPR is the most common risk factor for the development of breast cancer.

Prolonged lifetime exposure to estrogens, especially estrone and estradiol, has long been linked to the promotion and progression of breast cancer because of their physiological action on the mammary gland.^{23,24} Another in vitro study also reports significantly increased stimulation of the progesterone receptor by estrone,³⁸ although there is a lot of evidence suggesting that binding of ligands with hPRs and their transcriptional activity is also a function of phosphorylation of C-terminal amino acid residues as well as corresponding conformational changes in hPRs.³⁹⁻⁴¹ Due to the limitations of this in silico molecular modeling study, we did not consider different conformational changes of hPRs due to their differential phosphorylation inside the cells, and further crystallographic studies are required to confirm our findings. Our results may contribute an insight into the risk of breast cancer due to nonendogenous ligands, especially estrogens that bind with hPRs, although the higher binding energy of cholesterol with hPR AB cannot be explained by this study. However, it can be predicted that, in the presence of a deficiency of progesterone or an abundance of cholesterol, formation of a cholesterol-hPR AB complex may be favored at equilibrium, and may be another risk factor for breast cancer. This observation is in agreement with earlier studies reporting that increased total cholesterol levels are associated with development of breast cancer which is resistant to certain drugs, eg, tamoxifen.42-44

Conclusion

The absence of higher binding affinity of hPRs with progesterone in comparison with estradiol and estrone in a cellular or animal model can be explained by ligand receptor interaction kinetics and also by study of the physiological fate of such ligands in animal and cellular systems. This study may enhance our understanding of the specificity in binding of steroid derivatives to hPR. Our findings advocate that estradiol and estrone binds with hPR A and hPR B monomers with higher affinity in comparison to progesterone. Such interactions cause aggressive tumor development. Hence, our study may be helpful in understanding why the hPR A:hPR B ratio and prolonged exposure of nonendogenous ligands, particularly estrogens, to hPRs in the presence and absence of progesterone, is a risk factor for breast cancer.

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Disclosure

The authors report no conflicts of interest in this work.

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