A Molecular Docking of New 9β-Halogenated Prostaglandin Analogs with an Ester Group at C-6 Atom of the α-Side Chain

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Abstract: Prostaglandins with cytoprotective activity were studied for a long time and a few PGE1 and PGE2 stable analogs were promoted as drugs: arbaprostil, enprostil, misoprostol and riptostol. Nocloprost, a 9β-chlorine prostaglandin analog, has been also promoted as a cytoprotective drug; the success with this compound stimulated the research, and many 9β- or 11β-substituted prostaglandins were synthesized and studied for their biological activity. In the same direction we previously synthesized new 9β-halogenated prostaglandins having also an ester group at the carbon atom 6. These compounds were now used in a molecular docking study to predict their potential cytoprotective (anti-ulcer) activity. The study has been done with CLC Drug Discovery Workbench 2.4. software and an oxidoreductase enzyme receptor, chosen from the Protein Data Bank, ID: 4KEW. Two recognized drugs, omeprazole (co-crystallized with the enzyme) and nocloprost were used as standard in the study. The 9β-halogenated prostaglandin analogs were finally docked. Nocloprost and all 9β-halogenated compounds had docking score greater than that of omeprazole. The majority of the 9β-halogenated analogs have a docking score even greater than that of nocloprost, indicating that these compounds could have potential cytoprotective activity. Correlations between docking score and substituents on the prostaglandin skeleton have been done.

Keywords: 9β-halogenated prostaglandins; molecular docking; nocloprost; omeprazole; oxidoreductase enzyme receptor 4KEW; docking score; cytoprotective (anti-ulcer) activity.

1. Introduction
Prostaglandins with cytoprotective activity were studied for a long time and a few PGE1 and PGE2 stable analogues were promoted as drugs: arbaprostil [1], enprostil [2], misoprostol [3] and riptostol [4,5]. The compounds had secondary effects, mainly: diarrea, stimulation of uterine contraction, abdominal pain. So, other modified prostaglandin compounds, halogenated with chlorine, fluorine and bromine at 9 (9α or 9β) or 11 (11α or 11β) position of the cyclopentane ring of prostaglandins and also at the ω-side chain were synthesized [6-9] and their biological activity was determined. ‘Nocloprost” exhibits cytoprotective (anti-ulcer) activity [10] but other compounds, ZK-118182 [11], ZK-110841 [12, 13] and flunoprost have antitrombotic activity, 13,14-dihydro-ZK-110841 (AL-6556) [14] (Figure 1) reduces intraocular pressure, etc.

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Figure 1. Biological active 9-halogenated prostaglandin analogues
9-Halogenated prostaglandins mimic the 9-keto group and acts at the receptors of PGE₂ (Nocloprost is in this group of prostaglandin analogs) and 11-halogenated prostaglandins have great affinity for receptors of PGD₂, like for example compound ZK 110841. In the same direction we previously synthesized 9β-halogetaned prostaglandins and the results were published in a paper [15] and a patent request [16]. Brossing the literature, a molecular docking study of the 9-halogenated prostaglandins had not been found for predict their cytoprotective (anti-ulcer) activity.

2. Materials and methods

The docking studies have been realized according to the docking protocol [18]: importation of the protein/enzyme receptor, preparation of the protein receptor, setup the binding site and setup the binding pocket, introduction of the ligands in the “Molecule Project”, docking the ligands in the active binding site, extraction of the docked ligands in to a “Molecule Table”, calculation of the molecular properties of the ligands, screening the docking results. The validation of the method and of the docking parameters acquired from the molecular docking studies, have been carried out by redocking the co-crystallized docked in the active binding site of the protein receptor. The docking score and the hydrogen bonds established with the amino acids from the group of interaction are used to predict the binding modes, the binding affinities and the orientation of the docked compounds in the active site of the protein/enzyme receptor. The molecular properties of the small molecules, such as parameters of the Lipinski’s rule of five: the molecular weight, number of hydrogen bond donors, number of hydrogen bond and Log P (octanol-water partition coefficient), have been calculated using the “Calculate Molecular Properties” tool [22].

3. Results and discussions

The paper was taken in the study a molecular docking to predict the cytoprotective (anti-ulcer) activity of the 9β-halogenated prostaglandins of type 2 and 3. The compounds were obtained by opening the 8-lactone group of compounds 1 with diols (n = 0 to 4), or 2-butyne-1,4-diol, catalyzed by toluenesulfonic acid (Scheme 1), with the formation of an ester having an alcohol group at C-1 (prostaglandin numbering) spaced to carboxylic group by two to six methylene group or a 2-butyne group. The compounds have not only a 9β-halogen (Cl, Br and F) to act at the PGE₂ receptors, but also an ester group at the carbon atom 6. Such an ester group was presented in the literature and the PGE₂ type compounds showed cytoprotective activity [17] and the synthesized compounds presented in Scheme 1, having both modifications in the molecule, are waiting to have also cytoprotective activity.

![Scheme 1. Synthesis of 9β-halogenated prostaglandin analogues of type 2 and 3](image-url)
To put in evidence their predicted cytoprotective activity, we have done a molecular docking study, using CLC Drug Discovery Workbench 2.4. software [18]. It is to be mentioned that in the literature we didn’t found a similar molecular docking for Nocloprost or for other 9 or 11-halogenated prostaglandins.

The affinity of a compound to an identified protein or enzyme target is consider a relevant parameter in the process for development of a new drug. The prediction of the mode of binding of the ligand (generally, compounds in study) to the target (protein/enzyme) by molecular simulation could allow the restricting of the organic synthesis to the most promising chemical compounds.

We chose the 9β-halogenated prostaglandin analogues 2a-2l and 3a-3d, presented in Scheme 1, for the molecular docking study. The computational molecular simulation was performed to determine the affinity and orientation of the compounds and their mode of binding to an oxidoreductase enzyme receptor, chosen from the Protein Data Bank, ID: 4KEW. (www.rcsb.org) [19]. In the study we used as standard two recognized drugs, omeprazole (co-crystallized with the enzyme) and nocloprost (F2α prostaglandin analogue with a 9β-chlorine atom instead of 9α-OH group, Figure 1).

As usual, the binding site and binding pockets, used in the molecular docking of the ligands, were well established, and the search was carried out inside the binding site volume (Figure S1a, green sphere). The protein receptor, ID: 4KEW was loaded from Protein Data Bank, water molecules were removed, omeprazole co-crystallized was extracted, the binding site and also the binding pocket (which play an important role in orientation during molecular docking) were defined. The co-crystallized omeprazole was reposed in the protein pocket, docking validation and hydrogen bonds between co-crystallized omeprazole and amino acid residues of receptor were done. The prostaglandin analog drug nocloprost, used as standard in the study, then has been docked and the results are presented in Table 1 (entry 2); docking pose of the interactions between nocloprost and the amino acid residues are presented in Figure 2.

Table 1. Docking score and the molecular properties of ligands: omeprazole, nocloprost, 9β-halogenated prostaglandin analogs: 2a-2l and 3a-3d, calculated with CLC Drug Discovery Workbench 2.4 software

| Compound (Ligand) | Score | RMSD* | Atoms No. | Weight [Daltons] | Flexible bonds | Lipinski violation | HD | HA | Log P |
|-------------------|-------|-------|-----------|-----------------|----------------|-------------------|-----|-----|-------|
| Omeprazole (co-crystallized) | -58.11 | 0.06 | 41 | 328.41 | 5 | 0 | 0 | 5 | 3.28 |
| Nocloprost | -71.35 | 1.32 | 64 | 400.98 | 12 | 1 | 3 | 4 | 5.38 |
| 2a | -66.93 | 1.02 | 49 | 417.78 | 10 | 0 | 2 | 6 | 2.35 |
| 2b | -73.00 | 0.83 | 52 | 431.31 | 11 | 0 | 2 | 6 | 2.71 |
| 2c | -70.67 | 1.18 | 52 | 475.76 | 11 | 0 | 2 | 6 | 2.88 |
| 2d | -74.92 | 0.78 | 52 | 414.81 | 11 | 0 | 2 | 6 | 2.46 |
| 2e | -74.67 | 0.85 | 55 | 445.33 | 12 | 0 | 2 | 5 | 3.26 |
| 2f | -74.92 | 0.43 | 58 | 478.89 | 13 | 0 | 2 | 6 | 3.32 |
| 2g | -97.73 | 1.67 | 55 | 489.74 | 12 | 0 | 2 | 6 | 3.23 |
| 2h | -73.63 | 0.92 | 55 | 428.88 | 12 | 0 | 2 | 6 | 2.82 |
| 2i | -77.79 | 1.60 | 55 | 459.16 | 12 | 0 | 2 | 6 | 3.42 |
| 2j | -71.73 | 1.28 | 61 | 473.39 | 14 | 0 | 2 | 6 | 2.78 |
| 2k | -75.03 | 0.78 | 57 | 441.33 | 12 | 0 | 3 | 6 | 2.72 |
| 2l | -77.37 | 1.25 | 60 | 480.90 | 13 | 0 | 3 | 6 | 2.97 |
| 3a | -75.09 | 1.42 | 51 | 441.30 | 11 | 0 | 2 | 6 | 2.41 |
| 3b | -82.27 | 0.65 | 54 | 474.83 | 12 | 0 | 2 | 6 | 2.66 |
| 3c | -77.47 | 0.41 | 52 | 443.32 | 11 | 0 | 3 | 6 | 2.06 |
| 3d | -84.51 | 1.28 | 56 | 476.87 | 12 | 0 | 3 | 6 | 2.32 |

* RMSD: root-mean-square deviation; RMSD should be <2.

The docking score (PLANTPLP score) is a function described in Korb et al. [20]. “For a strong binding, the score has a negative value; for weak or non-existing binding, the score has a less negative or even positive value” [21].
The 9β-halogenated prostaglandin analogs 2a-2l and 3a-3d, presented in Scheme 1, were finally docked and the results of the calculated properties (flexible bonds, Lipinski violations, the number of hydrogen bond donors, the number of hydrogen bond acceptors and log P) are presented in Table 1. The calculated parameters can predict if a molecule possesses properties that might turn it into an active drug, according to the Lipinski’s rule of five [22,23]: 1) the number of hydrogen donors (N-H and OH) to be < 5, 2) the number of acceptors hydrogen < 10, 3) molecular weight < 500 Da, 4) the octanol-water partition coefficient [23] (log P) < 5. The number of violations of the Lipinski rules give a guidance to evaluate drug likeness for a molecule (if a molecule has chemical and physical properties to become a likely active orally active drug) [20,23]; omeprazole is taken as an example of a drug that confirms to this five’s Lipinski rules. According to the data presented in Table 1, all 9β-halogenated compounds comply with the Lipinski rules (Lipinski violation is 0), and nocloprost drug have one violation [20,23]. In Table 1, the docking score (and RMSD, which is < 2) is also presented. All 9β-halogenated analogs and nocloprost also have docking scores greater than that of omeprazole (-58.11, RMSD 0.06). The majority of the 9β-halogenated analogs have a docking score greater than that of 9β-chlorine nocloprost prostaglandin recognized drug (-71.25, RMSD 1.32), used as standard in the study, with the exception of the compounds 2a (-66.93, RMSD 1.01), 2c (-70.67, RMSD 0.83), 2g (-67.73, RMSD 1.67) and 2j (-71.13, RMSD 1.36) (See also Fig. 3). Basing on the docking score, the study shows that the cytoprotective (anti-ulcer) activity of the compounds 2b, 2d-2f, 2h-2i, 2k-2l, 3a-3d is greater than that of nocloprost. A few other observations should be mention:

- 3(CF3-phenoxo) are more active than 3-chloro-phenoyo substituted compounds (2f (-74.92) > 2e (-74.67); 2l (-77.57) > 2k (-75.03); 3b (-82.27) > 3a (-75.09); 3d (-84.91) > 3c (-77.47)).
- the length of the alkyl spacer between the 6-ester group and the hydroxyl group influence the anti-ulcer activity, in the order: 2a (n = 0; -66.93) < 2b (n = 1; -73.00) < 2l (n = 2; -74.67) > 2i (n = 3; -72.29) > 2j (n = 4; -71.13), most active being the compound 2l, with a similar length of eight atoms (with the oxygen group of the ester) to that of known in the α-side chain of natural prostaglandins.
- by reduction of 15-keto group to 15α-OH group, the docking score increase: -74.67 docking score of 2e increase to -75.03 for 2k and -74.92 docking score for 2f increase to -77.57 for 2l; the same is for 2-buten synthon of compounds 3a-3d: -75.09 for 3a increase to -82.27 for 3b and -77.47 for 3c increase to –84.91 for 3d.
- the influence of the 9β-halogen on the docking score is clear for keto-compounds: F (2d, -74.92) > Cl (2b, -73.00) > Br (2c, -70.67); for the 15-OH allylic alcohols, the chlorine substituted analog has a docking score (2e, -74.67) greater than that of the fluorine analog (2h, -73.63) and the bromine analog has the smaller value for docking score (2g, -67.73).
- The compounds with a 2-buten scaffold (3a-3d) in α-side chain had a docking score greater than that with a linear chain (2a-2l).
- the compound 3d had the best score (-84.91), followed by compound 3b (-82.27), compounds with a 2-butyn synthon in α-side chain; for compounds with normal α-side chain, the best score is for the compound 2l (-77.57).

An expressive presentation of the docking score is presented in Figure 3:

![Figure 3](image)

**Figure 3.** Docking score of the 9β-halogenated prostaglandin compounds 2a-2l and 3a-3d by comparison with the docking score of two cytoprotective (anti-ulcer) recognized drugs: omeprazole and no cloprost.

Besides the parameters mentioned in Table 1, group interaction, hydrogen bonds of ligands with amino acid residues were determined and hydrogen bond length was calculated. These are presented in the Table 2.

The docking poses of the ligands with the best score interacting with the amino acid residues of the protein are presented as follows: 3d in Figure 4, 3b in Figure 5, 3c in Figure 6 and 2l in Figure 7.

![Figure 4](image)

**Figure 4.** a) Hydrogen bonds between the residues of the ALA 330, TYR 51 and ARG 47 and the compound 3d; b) Docking pose of the compound 3d interacting with amino acid residues in the binding site.
**Figure 5.** a) Hydrogen bonds between the residues of the TYR 51, ALA 74, GLN 73, LEU 437 and GLU 352 and the compound 3b; b) Docking pose of the compound 3b interacting with amino acid residues in the binding site.
Figure 6. a) Hydrogen bonds between the residues of the ALA 330, ALA 74, TYR 51, THR 49, GLU 352 and the compound 3c; b) Docking pose of the compound 3c interacting with amino acid residues in the binding site.

Figure 7. a) Hydrogen bonds between the residues of the ALA 330, GLN 73 and GLU 352 and the compound 2l; b) Docking pose of the compound 2l interacting with amino acid residues in the binding site.

In fact, majority of 9β-halogenated ligands were found to have the same orientation with that of co-crystallized omeprazole and nocloprost, as can be observed in the Table 2. The compounds 2b, 2d, 2g, 2h and 2i adopted a different orientation than that of the co-crystallized and nocloprost.

It was observed that the best orientation is shown by the following 9β-halogenated compounds:

- **3d**, who reveals the best docking score -84.91 (RMSD 1.28 Å) and shows the occurrence of 5 hydrogen bonds with ALA 330 (2.987 and 3.280 Å), TYR 51 (2.656 Å) and with ARG 47 (3.053 and 3.280 Å) (Figure 4a);

- **3b**, who reveals the docking score -82.27 (RMSD 0.63 Å) and shows the occurrence of 5 hydrogen bonds with TYR 51 (3.140 Å), ALA 74 (2.837 Å), GLN 73 (2.999Å), LEU 437 (2.783 Å) and with GLU 352 (2.907 Å) (Figure 5a);
-3c, who reveals the docking score -77.47 (RMSD 0.41 Å) and shows the occurrence of 6 hydrogen bonds with ALA 330 (3.047 and 3.167 Å), ALA 74 (3.246 Å), TYR 51 (3.059 Å), THR 49 (3.319 Å) and with GLU 352 (3.194 Å) (Figure 6a);

-2I, who reveals the docking score -77.57 (RMSD 1.25 Å) and shows the occurrence of 5 hydrogen bonds with ALA 330 (3.113, 3.237 and 2.972 Å), GLN 73 (2.985 Å) and with GLU 352 (2.795 Å) (Figure 7a).

Table 2. The docking score and list of docking interactions between the ligand molecules and oxidoreductase enzyme receptor ID: 4KEV using CLC Drug Discovery Workbench Software.

| Compound (Ligand) | Score   | RMSD* | Group interaction | Hydrogen bond | Bond Length (Å) |
|-------------------|---------|-------|-------------------|---------------|-----------------|
| Omeprazole (co-crystallized) | -58.11  | 0.06  | PHE 42, LEU 20, THR 49, TYR 51, LEU 29, LEU 88, PRO 25, VAL 26, GLN 73, MET 354, SER 72, ALA 330, PRO 329, LEU 75, ALA 74, LEU 437, GLU 435, LEU 181, VAL 78, THR 438, THR 88, ALA 328, PHE 87, PHE 82 | -O sp² – (O2) - O sp³ – TYR 51 - N sp¹ (NE1) - N sp² – ALA 330 | 2.946 3.180 |
| Noxaprostone | -71.25  | 1.32  | PHE 81, PHE 82, VAL 78, LEU 181, LEU 75, THR 438, GLU 435, ILE 263, PHE 87, ALA 264, GLU 267, THR 268, ALA 328, THR 327, PRO 329, PHE 331, ALA 330, VAL 26, MET 354, LEU 29, TYR 51, ALA 74, GLN 73, LEU 188, LEU 20, SER 72 | -O sp² – (O1) - O sp³ – ALA 330 - O sp² – (O4) – N sp³ – ALA 74 - O sp³ (O4) - N sp³ – GLN 73 | 3.079 3.023 3.204 |
| 2a | -66.93  | 1.02  | PHE 87, CYS 400, LEU 73, SER 72, PHE 331, MET 354, TYR 51, ALA 330, ALA 74, ALA 264, GLY 265, ILE 263, GLN 73, VAL 26, PRO 329, THR 327, PRO 329, LEU 78, MET 354, THR 438, THR 327, GLU 435, LEU 20, VAL 26, PRO 25, LEU 188, PRO 329 | -O sp² – (O3) - O sp³ – TYR 51 | 2.880 |
| 2b | -73.00  | 0.83  | THR 88, PHE 87, PHE 82, VAL 78, LEU 75, SER 72, PRO 332, ALA 330, PRO 331, LEU 188, ALA 330, MET 354, THR 436, ALA 328, LEU 437, ALA 181, LEU 264, ILE 263, GLU 267, THR 268 | -O sp² – (O6) - O sp³ – SER 72 | 2.995 |
| 2c | -70.67  | 1.18  | GLY 265, THR 268, ALA 264, ILE 263, PHE 82, THR 438, LEU 437, ALA 74, GLN 73, VAL 26, LEU 29, PHE 42, ARG 47, LEU 188, LEU 20, VAL 78, THR 49, TYR 51, MET 354, PRO 329, ALA 330, PHE 331, PRO 329, LEU 75, SER 72 | -O sp² – (O1) - N sp³ – ALA 330 - O sp³ – (O6) – O sp³ – TYR 51 - O sp³ (O3) – O sp³ – TYR 51 - O sp² (O2) – N sp³ – ALA 74 | 3.005 3.255 2.932 3.320 |
| 2d | - 74.92 | 0.78  | LEU 20, LEU 188, ARG 47, PHE 42, THR 49, TYR 51, GLU 352, MET 354, PHE 331, ALA 330, SER 72, GLN 73, ALA 74, VAL 26, LEU 75, PRO 329, PHE 87, ALA 330, THR 327, PHE 82, VAL 78, LEU 437, THR 438, PHE 82, ALA 264, THR 260, THR 268, ILE 263, GLU 267, LEU 181, THR 436 | -O sp² – (O3) - N sp³ – ALA 264 - O sp² – (O1) - O sp³ – LEU 437 | 3.226 2.863 |
| 2e | -74.67  | 0.83  | PHE 81, PHE 82, VAL 78, LEU 181, PHE 87, ILE 263, ALA 264, GLU 267, THR 268, THR 327, ALA 328, THR 438, LEU 437, THR 327, PHE 329, PHE 331, VAL 26, ALA 330, MET 354, LEU 29, TYR 49, LEU 20, SER 72, ALA 74, LEU 75, GLN 73, LEU 188 | -O sp² – (O6) – O sp³ – SER 72 - O sp² – (O3) – O sp³ – TYR 51 | 2.994 2.979 |
| 2f | -74.92  | 0.43  | GLU 435, MET 185, PRO 25, LEU 20, LEU 29, VAL 26, LEU 188, LEU 181, GLU 267, THR 268, THR 263, ALA 264, PHE 82, PHE 87, LEU 75, SER 72, PHE 331, MET 354, TYR 51, LEU 356, GLN 73, ALA 330, ALA 328, ALA 74, PRO 329, THR 327, THR 438, LEU 437, PHE 42 | -O sp² – (O1) – O sp³ – SER 72 - O sp² – (O3) – O sp³ – TYR 51 | 3.285 2.610 |
| 2g | -67.73  | 1.67  | PRO 25, LEU 188, MET 185, GLU 435, THR 436, LEU 181, VAL 26, GLU 437, ALA 74, VAL 78, THR 438, GLU 267, ILE 263, THR 327, THR 268, ALA 264, PHE 82, PRO 329, ALA 328, ALA 330, THR 88, PHE 87, PHE 331, SER 332, MET 354, SER 72, MET 354, SER 72 | -O sp² – (O6) – O sp³ – SER 72 | 2.815 |
| 2h | -73.63  | 0.92  | ARG 47, LEU 188, GLN 73, GLU 352, PHE 42, THR 49, LEU 20, SER 72, ALA 74, LEU 75, VAL 78, PHE 82, LEU 181, PHE 87, THR 260, ILE 263, ALA 264, LEU 437, THR 436, THR 438, SER 332, MET 354, TYR 51, VAL 26, ALA 330, PRO 329, ALA 328, THR 327, THR 268, PHE 331 | -O sp² – (O3) – N sp³ – ALA 264 - O sp² – (O1) – O sp³ – LEU 437 | 3.110 3.081 |
| 2i | -72.29  | 1.60  | GLY 265, GLU 267, ILE 263, THR 268, ALA 264, THR 327, ALA 328, PHE 87, PRO 329, PHE 331, ALA 330, PHE 82, THR 438, VAL 78, LEU 437, GLU 435, MET 185, PRO 25, LEU 188, LEU 29, PHE 42, PHE 87, GLN 73, VAL 26, ALA 74, SER 72, MET 354, TYR 51, LEU 75 | -O sp² – (O1) – O sp³ – SER 72 - O sp² – (O5) – N sp³ – THR 438 | 3.259 3.179 |
### 4. Conclusions

Prostaglandin analogues substituted with a 9β-halogen and with an ester group with a diol at the C-6 carbon atom, previously synthesized, were used in a molecular docking study to determine their potential cytoprotective (anti-ulcer) activity. The study has been done with CLC Drug Discovery Workbench 2.4. software and an oxidoreductase enzyme receptor, chosen from the Protein Data Bank, ID: 4KEW. (www.rcsb.org). In the study we used as standard two recognized drugs, omeprazole (co-crystallized with the enzyme) and noclopast. The 9β-halogenated prostaglandin analogs 2a-2l and 3a-3d were finally docked. Noclopast and all 9β-halogenated compounds had docking score greater than that of omeprazole. The majority of 9β-halogenated analogs have a docking score greater than that of noclopast, with the exception of the compounds 2a, 2c, 2g and 2j. The compounds 2b, 2d-2f, 2h-2i, 2k-2l, 3a-3d had a greater docking score than that of noclopast; this indicate that the compounds could have potential cytoprotective (anti-ulcer) activity. A few correlations between docking score and substituents on the prostaglandin skeleton have been done.

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