In silico study: Assessment of the inhibition of cyclo-oxygenase 2 by ibuprofen by validating molecular docking and cardiovascular effects reported during the COVID 19 pandemic

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Abstract – Introduction The Covid 19 pandemic has put the cardiovascular risk incurred when using nonsteroidal anti-inflammatory drugs at the heart of the discussion. Based on the information currently available, WHO does not recommend the use of ibuprofen. The objective is to evaluate the inhibition of cyclo-oxygenase 2 by ibuprofen by validating molecular docking.

Method The crystallographic structure of ibuprofen bound to cyclooxygenase-2 was obtained from the Protein Data Bank (PDB) at a resolution <3.00 Å.

The receiver was visualized using Discovery Studio Visualizer version 2.5.5. It was efficiently prepared using AutoDock / Vina software.

The 3D structure of Ligand (Ibuprofen) was downloaded from the Drugbank database (https://www.drugbank.ca/): Accession number DB01050

Results Molecular docking was chosen as the first-line discrimination of the ibuprofen-COX2 interaction for the in silico study of putative competitors. The complex formed by Ibuprofen-COX 2 from the experimental model gives a docking score (Affinity: -7.3 (kcal / mol) with a mean square deviation of (RMSD = 23.884).

Conclusion The evaluation of the inhibition of cyclo-oxygenase 2 by ibuprofen was validated by molecular docking. Cardiovascular effects already reported in patients treated with traditional non-steroidal anti-inflammatory drugs and coxibs have been observed in patients with COVID 19. Molecular docking becomes an essential step in drug discovery to explore other drug targets.

Keywords: Cyclooxygenase-2, Ibuprofen, Molecular docking, Covid 19

1. Introduction

The use of ibuprofen during COVID 19 infection has been prohibited. Severe forms reported in young patients who had no health problems after using nonsteroidal anti-inflammatory drugs (Day 2020)

Several previous studies have shown a complicated course with an increased incidence of empyema, lung cavitation, and prolonged intensive care unit stay when nonsteroidal anti-inflammatory drugs (NSAIDs) were used in patients with pneumonia (Voiriot and al. 2019).

Cardiovascular side effects associated with NSAIDs have been previously reported including hypertension, heart failure and kidney failure (Karsh 2006).

Coxibs were developed as an alternative solution to the gastrointestinal toxicity of NSAIDs. Unfortunately, an increased risk of cardiovascular episodes has been observed with all coxibs and, where data were available, with traditional NSAIDs such as diclofenac and ibuprofen (Gislason and al. 2006). These risks have led to a re-evaluation of these traditional therapies.

This is why we can no longer underestimate the risks of cyclo-oxygenase inhibitors, whether traditional NSAIDs or coxibs. Moreover, the risks are not limited to long-term use; clinically serious adverse cardiovascular episodes have been reported during the first days of treatment (Karsh 2006)

In view of the scientific debates led by researchers around the world on the report, the objective is to evaluate the inhibition of cyclo-oxygenase 2 by ibuprofen through a molecular docking validation.
2. Materials and methods

To locate the sites of interaction, four different categories of COX inhibitors have been documented by Smith et al., (Smith and al. 2011). Several works have determined the crystal structures of COX-1 and COX-2 in complex with inhibitors and substrates (Kurumbail and al. 1996), (Vecchio and al. 2011). Selinsky et al determined the crystal structure of IBP bound to COX-1. (Selinsky and al. 2001).

However, the analgesic and anti-inflammatory effects of ibuprofen (IBP) arise from inhibition of COX-2 rather than COX-1 (Laneuville and al. 1994). In order to compare the mode of binding of IBP to COX-2 versus COX-1, and to reveal a possible mechanism of selective substrate inhibition mediated by IBP, Orlando et al determined the crystal structure murine (mu) COX-2 complexed with Ibuprofen (IBP) (Orlando and al. 2015). The coordinates of the X-ray crystals of COX-2 in complex with ibuprofen (IBP formula: C13 H18 O2) (PDB ID: 4PH9) in Table 1, 2, 3, were extracted from the RCSB database (http://www.rcsb.org/pdb). This is selected for modeling studies.

Table 1: Macromolecules in Ibuprofen (IBP) Bound to Cyclooxygenase-2 Complex

| Entity ID : 1 | Molecule | Chains | Sequence Length | Organism | Details |
|--------------|----------|--------|----------------|----------|---------|
|              | Prostaglandin G / H synthase 2 | A, B | 551 | Mus musculus | Mutation(s): 0 Gene Noms: Ptgs2, Cox-2, Cox2, Pghs-b, Tis10 EC: 1.14.99.1 |

Table 2: Oligosaccharides in Ibuprofen in (IBP) bound to cyclooxygenase-2 complex

| Entity ID : 2 | Molecule | Chains | Sequence Length | Glycosylation |
|--------------|----------|--------|----------------|---------------|
|              | 2-acétamido-2-désoxy-bêta-D-glucopyranose- (1-4) -2-acétamido-2 désoxy-bêta-D-glucopyranose | C, D | 2 | N-glycosylation |

| Entity ID : 3 | Molecule | Chains | Sequence Length | Glycosylation |
|--------------|----------|--------|----------------|---------------|
|              | alpha-D-mannopyranose- (1-4) -2-acétamido-2-désoxy-bêta-D-glucopyranose- (1-4) -2-acétamido-2-désoxy-bêta-D-glucopyranose | E | 3 | N-glycosylation |
| Ligands          | ID   | Chains | Name / Formula / InChI Clé                          | Diagramme 2D               |
|------------------|------|--------|----------------------------------------------------|----------------------------|
| HEM              | A, B |        | PROTOPORPHYRINE IX CONAIN FE                      |                             |
|                  |      |        | C_{34} H_{32} Fe N_{4} O_{4}                      | KABFMIBPWCXCRK-RGGAHWMASA-L|
| BOG              | A, B |        | octyl bêta-D-glucopyranoside                       |                             |
|                  |      |        | C_{14} H_{28} O_{6}                                | HEGSGKPQLMEBJL-RKQHYHRCSA-N|
| NAG              | A, B |        | 2-acétamido-2-désoxy-bêta-D-glucopyranose          |                             |
|                  |      |        | C_{10} H_{15} NO_{6}                               | OVRNDRQMDRJTHS-FMDGEEDCSA-N|
| IBP              | A, B |        | IBUPROFEN                                         |                             |
|                  |      |        | C_{13} H_{18} O_{2}                                | HEFNNWSXXWATRW-JTQLQIEISA-N|
2. 1- Protein preparation

The X-ray crystallographic structure of ibuprofen bound to cyclooxygenase-2 (4PH9) (from Mus musculus) was obtained from the Protein Data Bank (PDB) (http://www.rcsb.org) at resolution <3.00 Å (resolution: 1.81 Å, free R-value: 0.197, working R-value: 0.160, observed R-value: 0.162).

This file is not directly used by Autodock 1.5.6 tools, it was first viewed using Discovery Studio Visualizer version 2.5.5. Discovery Studio is a biological molecular design solutions software package for chemists and computational biologists. (Discovery Studio 2.5 (CDOCKER Dock, Dassault Systemes BIOVIA, USA) Discovery Studio facilitates the examination of the properties of large and small molecules

Water molecules, ligands and other heteroatoms have been removed from protein molecules. These crystallographic structures were preserved without any treatment for molecular anchoring. The protein PDB files were efficiently prepared using AutoDock / Vina (Molecular Graphics Lab, The Scripps Research Institute Plugin, La Jolla, CA, USA) and saved as pdbqt files. Polar hydrogens and Kollman chrages have been added.

2- 2- Preparation of the ligand

The 3D structure of Ligand (Ibuprofen) was downloaded from the Drugbak database (https://www.drugbank.ca/): Accession number DB01050. Torsional connections have been verified. Ligand was saved in the same anchor folder in pdbqt format. The charges were then added.

2 3- Molecular docking protocol

Docking calculations were performed using standard AutoDock Vina defaults. The active site was placed in a 40 × 50 × 40 Å cubic box at the geometric center of the selected flexible residue set, which has 0.375 Å as the grid point spacing

The values of the root mean square deviation (RMSD) between the mooring and the initial poses were calculated. The resulting best poses were ranked based on Vina scores (kcal / mol), assessed by free binding energies (S score, kcal / mol) and binding interactions between the ligand atom and the active site residues.
3. **Result**

Molecular docking was chosen as the first-line discrimination of the ibuprofen-COX2 interaction as a structural discrimination procedure based on the in silico prediction of putative competitors (Liu and al. 2017). The structural differences of each molecule also influence its interaction with COX2 (figure 1).

![Image 1: Structure of Ibuprofen and other drugs with anti-inflammatory action (By DrugBank)]

Each COX isoform is a structural homodimer which functions as a heterodimer. According to Mitchener et al., one subunit, containing the required heme prosthetic group, acts as a catalytic site, while the other serves as an allosteric site (Mitchener and al. 2015). Previous evidence suggests that inhibitors may act at one or both sites, depending on the structure and concentration of the inhibitor (Dong and al. 2013). Regardless of the site, Picot et al., identified an open area called a "lobby" through which binding requires that a small molecule must first enter (Picot and al. 1994).

As shown in Figure 2, ibuprofen was identified in the pocket of the COX2 enzyme binding site.

![Image 2: Ibuprofen in the pocket of the COX2 enzyme binding site (with Discovery Studio 2.5 software).]
The active site was placed in a cubic box at the geometric center of the selected flexible residue set at 0.375 Å as the grid point spacing.

The coordinates of the Grid Box (number of points X, Y, Z dimensions and spacing) for the Ibuprofen-COX2 model (shown in Figure 3) were noted.

![Figure 3: Grid box adjusted to the Ibuprofen-COX2 model](image)

A validation docking of the crystal structure of murine (mu) COX-2 in complex with Ibuprofen (IBP) (Orlando and al. 2015). (PDB ID: 4PH9) (http: //www.rcsb.org/pdb). Was performed on Autodock vina after visualization in discovery studio and processing of the ligand and receptor.

![Figure 4a](image)  ![Figure 4b](image)

**Figure 4:** COX2 pocket (4a: Binding Ibuprofen, 4b Free)

The complex formed by Ibuprofene-COX 2 from the experimental model of Orlando et al has a very low energy value and gives a docking score (Affinity: -7.3 (kcal / mol) with a mean square deviation of (RMSD = 23.884 ).

The interactions between COX2-Ibuprofen have been illustrated in Figures 5a and 5b.
4. Discussion:

Furse et al studied molecular dynamics simulations of COX-1 and COX-2 enzymes with its human substrate (arachidonic acid). These simulations were compared to reference simulations of arachidonate in solution to explore the effect of the enzyme on the conformation and positioning of the substrate in the active site. The simulations suggest that the substrate has greater conformational freedom in the COX-2 active site compared to COX-1, which is consistent with the greater volume of COX-2 active site observed in crystal structures on x-rays (Furse and al. 2006).

Cyclooxygenases are functional homodimers. In our work, the active site was identified in the A chain of cox2 in the cox2 structure linked to ibuprofen. Our simulations only involved a single monomer, like some work (Cukier and al. 2002), or others that focused on an even smaller fragment of the protein, like the membrane binding domain (Nina and al. 2000) or the membrane and helix binding domain comprising the active site of cyclooxygenase (García-Nieto and al. 2000).

In addition, previous studies have indicated that many different NSAIDs (including IBP) bind tightly to a single monomer of the COX-2 dimer and allosterically inhibit oxygenation of the substrate into the partner monomer (Dong and al. 2011), (Duggan and al. 2011).

The results of our docking confirm those found by Orlando et al on the experimental level. Orlando et al identified ibuprofen in an area of the enzyme between the opening of the substrate channel and the top of the active site (Orlando and al. 2015).

Looking at gastrointestinal outcomes as the primary endpoint, preliminary evidence has shown the link between COX inhibition and the risk of cardiovascular disease. The VioXX trial in patients with rheumatoid arthritis treated with Rofecoxib were five times more likely to have a myocardial infarction than patients on non-selective anti-inflammatory inhibitor Naproxen (Bombardier and al. 2000).

Following the results of the Adenomatous PolyP Revention On Vioxx (APPROVe) study, rofecoxib was withdrawn from the market in 2004 (Baron and al. 2008).

A meta-analysis of 280 trials of nonsteroidal anti-inflammatory drugs versus placebo showed that high dose naproxen was associated with a risk, albeit a low one, of vascular events (Bhala and al. 2013). Furthermore, the risk of cardiovascular disease was not equal for all COX-2 inhibitors. Questions also remained about the dose and duration of use of nonsteroidal anti-inflammatory drugs associated with a high risk of cardiovascular disease.

5. Conclusion

Our docking validated the experimental results known in current practice concerning the inhibition of cyclooxygenase by ibuprofen. Molecular docking becomes an essential step in drug discovery so that new exoges ligands can be identified.
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