Molecular dynamics simulation and docking analysis of NF-κB protein binding with sulindac acid

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Abstract:
It is of interest to document the Molecular Dynamics Simulation and docking analysis of NF-κB target with sulindac sodium in combating COVID-19 for further consideration. Sulindac is a nonsteroidal anti-inflammatory drug (NSAID) of the arylalkanoic acid class that is marketed by Merck under the brand name Clinoril. We show the binding features of sulindac sodium with NF-κB that can be useful in drug repurposing in COVID-19 therapy.

Keywords: NF-κB; molecular dynamics simulation; COVID-19; SARS-CoV2; sulindac sodium

Background:
Sulindac is a non-steroidal anti-inflammatory agent (NSAIA) that can block nuclear factor-κB (NF-κB), a transcription factor (TF) located primarily in the cytoplasm in a complex while in an inactive state. Activation of this TF makes it transition from a latent cytoplasmic form to a nuclear DNA binding state [1] and has been demonstrated to have a role in the prevention of colon cancer [2] by its inhibition [1, 3]. This TF is known to selectively inhibit interferon-gamma-induced expression of the chemokine CXCL9 gene in mouse macrophages. [4] The growth inhibitory and anti-inflammatory properties of sulindac are possibly due to its ability to reduce prostaglandin synthesis by cyclooxygenase inhibition [2]. Furthermore, Yamamoto et al. [2] have demonstrated that aspirin and sodium salicylate inhibited the activity of an IkB kinase β (IKKβ) that is required to activate the NF-κB pathway, which leads to nuclear translocation of NF-κB, where it binds to its cognate DNA and activates transcription of a wide variety of genes involved in host immunity, inflammation, cell proliferation and apoptosis [2, 3]. Various NF-κB inducers are known to be highly variable and include aggregates of bacterial lipopolysaccharides, ionizing radiation, reactive oxygen species (ROS), cytokines, such as tumour necrosis factor alpha (TNF-α) and interleukin 1-beta (IL-1β) and viral DNA and RNA [4]. Upon activation, NF-κB promotes the gene expression of a broad range of cytokines (e.g., IL-1, IL-2, IL-6, IL-12, TNF-α, LT-α, LT-β, and GM-CSF), chemokines (e.g., IL-8, MIP-1, MCP1, RANTES, and eotaxin), adhesion molecules (e.g., ICAM, VCAM, and E-selectin), acute phase proteins (e.g., serum amyloid A: SAA), and inducible effector enzymes (e.g., inducible nitric oxide synthase: iNOS and cyclooxygenase-2: COX-2). The NF ΚB can be considered as a “quick action” primary transcription factor that is able to regulate myriad of cellular responses, including the host’s early innate immune response to infection, and is also associated with chronic inflammatory states, viral infections, septic shock syndrome and multiorgan failure [5, 6] viz-a-viz ascribed as an anti-apoptotic TF [7]. The constitutive activation of NF-κB pathways has been responsible for stimulation of inflammatory diseases, such as multiple sclerosis and rheumatoid arthritis [8, 9]. Furthermore, p38 MAPK-based activation of NF-κB has also been reported [9]. Drug induced suppression of NF-κB has been found to have a strong potential in proapoptotic signal modulation therapy [10].

Recently, a study utilizing molecular dynamics simulations (MDS) - based analysis targeting human angiotensin-converting enzyme 2 (hACE2) as a receptor showed interesting results on development of a lead candidate to be used later as a therapeutic drug against COVID-19 [11]. Though in the current scenario, the spread of COVID 19-based pandemic has slowed down to a great extent, but its resurgence cannot be ruled out. Due to the lack of adequate specific treatments for COVID-19, there is an urgency to develop or repurpose drugs to help end the epidemic completely. The drug discovery process has received a great impetus with the emergence of computer-aided drug discovery (CADD) and has been instrumental over the last decade in exploring protein inhibitors in protein-drug interactions and protein–protein interactions [12]. Since the process involved in the development of a candidate drug into an approved drug is lengthy and expensive, a combination of computational methodologies such as virtual screening, docking, molecular dynamics simulation, and binding free energy evaluation contributes to identifying potential drug candidates from compound libraries [13]. To date, MDS and structure-based virtual screening studies have been carried out to understand the SARS-CoV-2 spike protein functions [14-16], role of repurposed protease inhibitors in blocking SARS-CoV-2 [17], effects of some antiviral drugs on SARS-CoV-2 [18], effect of temperature on the structure of the spike protein [19], SARS-CoV-2-targeted potent drugs that can be effective in controlling COVID-19 [20, 21] as well as many other aspects of biology qualifying to control SARS-CoV-2 progression. However, the rate of occurrence of different pathological human phenotypes and their heterogeneous lethality rates arising from constant mutations of SARS-CoV-2 [22, 23] indicate serious problems that need to be appropriately dealt with. A paradigm shift in the strategy on targeting COVID-19 borne biomolecules-for therapeutic purposes to human-based molecular biomarkers is much necessitated in the current scenario, since the current approaches based on former fail to yield appreciable outcomes. A recent study revealed the importance of the NF-κB pathway in developing therapy regimens for critical COVID-19 patients [24]. Other reports have also substantiated the above fact by arguing that therapeutic benefits of NF-κB inhibitors, including dexamethasone, a synthetic form of glucocorticoid, have increasingly been realized now [17]. Abnormal activation of NF-κB resulting from SARS-CoV-2 infection might be associated with the pathogenic profile of immune cells, cytokine storms and multiorgan defects [23]. Therefore, we currently explored the structural details of NF-κB via
an in silico approach wherein we pinpointed the protein pockets for the binding of SARS-CoV-2 spike proteins. A hypothetical explanation currently generated is that upon entering the human body, the COVID-19 viral protein might be triggering the action of the NF-kB directly or through activation of IKK, the kinase phosphorylating IxBs that otherwise remain bound with NF-kB to keep it in an inactive state in the cytoplasm. This argument is made in the light of the fact that another complex member, IκBα, is known to be phosphorylated by specific kinases at two sites near the N-terminus (Ser-32 and Ser-36), while the phosphorylated protein is then ubiquitinated at Lys-21 and Lys-22, leading to proteosome-mediated degradation [24] and release of NF-kB from the complex. The removal of IκB unmasks the nuclear localization sequence (NLS) of NF-kB and allows its translocation to the nucleus [24]. Thus, IκB is a multifunctional inhibitor of NF-kB that blocks nuclear translocation, DNA binding, and phosphorylation by protein kinase A (PKA). Inhibition of NF-kB blocks the NF-kB-mediated deactivation of immune and inflammatory responses to stimuli, such as cytokines or bacterial/viral infection products [24]. Along with NF-kB, sulindac has been of interest because of its many roles that includes also as a chemopreventive agent for adenomatous colorectal polyps and colon cancer [25]. We show here that sulindac is able to bind specifically to a pocket on NF-kB, a concept that becomes theoretically important for exploring its therapeutic potential [26-29].

Figure 1: Root mean square deviation (RMSD) of the protein and ligand after the initial RMSD values were stabilized. This plot shows RMSD values for the protein on the left Y-axis, whereas for the ligand, these values are indicated on the right Y-axis. The RMSD graph for c-axis shown in blue colour, and for ligand fit on the protein, it is in red colour.

Materials and Methods:
Protein and Ligand Preparation:
Coordinates of the X-ray diffraction-based crystal structure of the IκB-a/NF-kB complex with solvents (PDB ID: 1NFI) at a resolution of 2.70 Å were downloaded from the Protein Data Bank (PDB). Solvent molecules and chains A, B, and F were removed during protein preparation using the Protein Preparation wizard from Schrodinger Maestro (Schrodinger, LLC, and New York, NY, USA). The remaining structures were processed using the Protein Preparation wizard using appropriate methodologies. The SARS-CoV-2 structure possessing a loop (residues 376-381) was missing in the PDB structure and hence was modeled using Schrodinger’s Prime module. Hydrogen atoms were incorporated, and a standard protonate state at pH 7 was used. Bond orders were assigned using the chemical components dictionary (CCD) database. The heteroatom was generated using Epik with a pH of 7 (±2.0), and Prime was used to fill the missing chains and loops. While refining the structure, PROPKA with pH 7.0 was used along with the sample water orientation. While minimizing the structure, a root mean square deviation (RMSD) of 0.30 Å with the OPLS_2005 force field was used. The ligand was selected based on published information regarding its use as a therapeutic molecule for various human diseases. The ligand was prepared using the Ligprep wizard of Schrodinger with the OPLS_2005 force field; the Epik job was submitted with the metal-binding site and included the original state. Thirty-two stereoisomers were assigned per ligand with specified chirality’s. Glide was used to filter the search to locate the ligand in the active-site region of the receptor. The shape and properties of the receptor were represented on a grid to provide a more accurate scoring of the ligand poses. The docked complexes were superimposed on the original crystal structure to calculate the RMSD (Figure 1).

Virtual Screening and Molecular Docking:
The Glide grid generator was used for generating the grid for blind docking. A Schrodinger virtual screening workflow (VSW) was used to score the virtual screening with default parameters employing the Glide program of Schrodinger. Use of HTVS mode allowed the elimination of most of the stereoisomers, and only 10% of the total ligands could be retained that passed the screening. Ligands following QikProp and Lipinski’s rule were filtered for docking. Docking in the VSW was performed with Epik state penalties and further passed through HTVS, SP and XP docking modes. The interactions of the selected ligand and protein docked complexes were analyzed by a pose viewer.

Molecular Dynamics Simulation:
To study the dynamic behavior of the protein complex under simulated physiological conditions, MDSs of the protein-ligand (PL) complex were performed using Desmond, which is available with Schrodinger Maestro (v12.5). The PL complex (9873 atoms) was solvated in a 10 × 10 × 10 Å orthorhombic periodic box built with TIP3P water molecules [30]. The whole system was neutralized by adding an appropriate number of 6 Na+ counterions. This solvated system was energy minimized and position restrained with OPLS_2005 as the force field. Furthermore, 100 ns of MDS was carried out at 1 atm pressure and 300 K temperature, implementing the NPT ensemble with a recording interval of 100 ps, resulting in a total of 1000 read frames. Finally, various parameters of the MDS, such as ligand-binding site analysis, RMSD, root mean square fluctuation (RMSF), PL contacts, secondary structure element (SSE) analysis, etc., were also analyzed to check the stability, compactness, structural fluctuations and PL interactions in the solvated system (Figure 2).
Interaction Analyses
Structural Data:
The PDB was used to extract all 3-D structural information. The X-ray diffraction-based crystal structure of the I-κB-α/NF-κB complex with solvents (PDB ID: 1NFI) at a resolution of 2.70 Å was used for this study. Solvents and chains A, B, and F were removed during protein preparation. The structure of the SARS-CoV-2 spike glycoprotein (closed state) solved using electron microscopy (PDB ID: 6VXX) with a resolution of 2.80 Å was obtained. Furthermore, all water molecules were removed from both structural data sets. The educational version of PyMOL (The PyMOL Molecular Graphics System, v1.2r3pre, Schrödinger, LLC) was used to generate the images derived to understand and analyse the structure and inter chain interaction information.

Molecular Docking:
The ligand sulindac chosen in the current study for the mentioned reasons has its docking score and binding affinity shown in Figure 3. The complex of the drug molecule docked in the pocket of NF-κB was designated S0 (Figure 4A) and was taken as the reference for further interaction analysis.

Pocket Analysis:
The molecule NF-κB was found to having multiple pockets, as calculated using the CASTp 3.0 server [30, 31] Three largest pockets (based on the accessible area) of these were considered for analysis. The residues present in these pockets interacted with the spike protein on exposure changing the area of the pockets significantly due to this interaction and also upon the drug interaction or the combination of the spike protein and the drug together.

Protein-Protein Docking:
Protein-protein docking studies were carried out using ClusPro 2.0 [32] with minor changes following the protocols used in Jorgensen et al. [33]. This job was carried out to understand the interaction of the SARS-CoV-2 spike glycoprotein with NF-κB in the presence and absence of the drug molecule. The best pose of the spike protein binding with the NF-κB-IκB complex was selected based on the cluster size, which is how the models are ranked in ClusPro. S1 (Figure 4B) represents the complex of NF-κB with the SARS-CoV-2 spike glycoprotein, and S2 (Figure 4C) represents the complex of S0 with the SARS-CoV-2 spike glycoprotein.

LigPlot+ v.2.2 [34] was used to find the ligand-protein interaction of complex S0 (Figure 2). PDBsum was used to calculate the interaction between NF-κB and IκB (Figure 5A, 5B), the interaction between the spike protein and NF-κB (Figure 5B) and the interaction between the spike protein and NF-κB docked with drug molecules. PyMOL was used to analyse the structural differences imposed on NF-κB when it interacted with the drug molecule and the spike protein in the presence of the drug molecule.

Prime MM-GBSA (molecular mechanics, the generalized born model and solvent accessibility method) calculations were performed with the docked complex of receptor and sulindac. The implicit solvent model used is VSGB [35] which has an efficient approximation of the solvation free energy with Surface Generalized Born (SGB) over Vacuum and chloroform solvation models. The following calculations were made

Rec Strain = Receptor (from optimized complex) − Receptor
Lig Strain = Ligand (from optimized complex) − Ligand
MMGBSA dG Bind = Complex − Receptor − Ligand
MMGBSA dG Bind (NS) = Complex − Receptor (from optimized complex) − Ligand (from optimized complex) − MMGBSA dG Bind

Rec Strain - Lig Strain:
NS here refers to binding or interaction energy without any involvement of conformational changes occurring in the receptor or ligand. The potential energy of the complex will be reported in kcal/mol.
Results and Discussion:

**MD Simulation Studies:**

The complex RMSD in the initial phase at 1.63 Å went to 4.11 Å while stabilizing the structure for 100 ns of simulation. Initially, at up to 50 ns, the RMSD did not fluctuate much, but after that, the stability fluctuated for 10 ns and stabilized subsequently (**Figure 1**). The RMSF plot analysis displayed minimal fluctuations in the protein structures (**Figure 2**). Furthermore, minimal fluctuations were also observed in the PL complex, and the RMSF plot showed fluctuations in some regions of the protein. The results of the residue interaction analysis after docking are highlighted with specific colours based on the self-explanatory features (**Figure3**). The docking interactions of sulindac (**Figure 4A, 4B**) showed that it docked in the largest pocket of the NF-$\kappa$B-IkB complex that contains most of the residues interacting with the spike protein. It is possible that drug binding to the complex interferes with the above interaction of the spike protein, however, needs to be ascertained with more robust wet laboratory-based experimental designs. The sulindac was found to bind to other pocket than the one occupied by NF-$\kappa$B-IkB complex (**Figure 4C**). Moreover, data shows the importance of NF-$\kappa$B as a suitable drug receptor for COVID-19 for its proven roles as molecular targets in drug discovery-based in silico studies (**Table 1**).

The interaction made by the drug molecule with the residues of the active site cavity of NF-kB, IkB and P50 peptide chains is dynamic in nature and formed, broke and reformed during the simulation duration (**Figure 3**). A brief duration of 20 ns was observed when no interaction occurred between the drug molecule and the NF-$\kappa$B active site residues. Although there was an interaction between the ligand (drug) molecule and residues of NF-kB (chain C) and IkB (chain E) in the first 50 ns, the interaction between the drug and IkB (chain E) disappeared after 50 ns. This loss of interaction strengthened the ligand-NFkB interaction.

**Table 1:** The binding affinities of various drugs with NF-$\kappa$B are shown. This shows that NF-$\kappa$B is a candidate receptor in various in silico drug-based interventions that may have potential for being a therapeutic target.

| Sl. No.| Bioactive compound/ligand                   | Binding pocket                  | Outcomes achieved/remarks                                                                 | References |
|-------|--------------------------------------------|---------------------------------|------------------------------------------------------------------------------------------|------------|
| 1     | Piperine and piperlongumine                | DNA binding sites               | Docking results of the analogues had higher affinity towards NF-$\kappa$B and IL-$\beta$ than natural molecules | [36]       |
| 2     | 71 anti-cancer compounds along with dipeptidyl boronic acid, betulinic acid, and glycyrrhetinic acid inhibitors | Active sites: Gly47, Ala49, Thr21, and Thr1 of the receptor | Docking was at par with the drug, Bortezomib against proteasome target of NF-$\kappa$B pathway | [37]       |
| 3     | Compound series designed based on quinazoline scaffold pharmacophore model | p50 subunit of NF-$\kappa$B      | Compounds can inhibit NF-$\kappa$B function and reduce the proliferation of numerous tumor cell lines especially compound | [38]       |
4. P100 peptide  Binding sites of IkB Kinase α (IκK α)  Highlights the different roles of NIK and IkB in regulating p100 processing and understands the mechanisms that mediate control of p100 processing.

5. Four series of 24 thienopyrimidine/N-methylpicolinamide derivatives substituted with pyrimidine  Active sites of TAK1 kinases  Compound 38 inhibited TAK1 phosphorylation, IκBα degradation, TNF-α-induced IκB kinase phosphorylation, \( \text{IκBα} \) degradation, TNF-α-induced IκBα phosphorylation, reduced the expression of p65 and p50 phosphorylation.

6. Growth arrest specific factor 6 (Cas6)  Autophosphorylation docking site: Tyr867  Merk receptor showed distinct effects for phagocytosis.

7. L-8-dihydroxy-4-methylanthracene-9, 10-dione (DHMA) from Luffa acutangula  DNA-binding region (DBR) of NF-κB  DHMA plays a role in cancer prevention and treatment by having an altered binding affinities against NF-κB and DNA.

8. Natural compounds found in Cupressus pyramidalis and Aegle marmelos extracts  DNA binding sites of proteins  These lead molecules down-regulated the expression of IκB gene and inhibited the activity of IκB-a/p50-50.

9. Phenolic fraction of Citrus max (hopteridine, naringin, naringex, and dexamethasone)  DNA binding region of NF-κB  Docking studies of the phenolic fraction compounds had an inhibitory effect against NF-κB.

10. Diphosphorylated 25-mer p100 peptide model  Four arginine residues Arg285, Arg410, Arg431, Arg521 and Arg544 in hydrophobic B-TrCP  Peptide p100 disrupts the IκB-α/NF-κB signalling pathway.

11. Small molecule, NSC-127102  Phosphorylation of p65 at serine 276  NSC-127102 binds IL-8, VCAM-1 expression and phosphorylation at serine 276.

12. Scopoletin (Pharmacologically active coumarin)  Keap1, NF-κB, HO-1 and p38MAPK are the targets against vancomycin nephrotoxicity  Scopoletin helps in protecting renal injury by interfering with Nrf2/HO-1 and IkBα/p65 NF-κB signalling pathway as well as resorting the antioxidant defense system.

13. Ergosta-7,9(11),22-trien-3β-ol (EK100)  Interferes with Lipopolysaccharide (LPS) docking to the LPS-binding protein (LBP), transferred to the cluster of differentiation 14 (CD14), and bonded to TLR4/myeloid differentiation-2 (MD-2) co-receptors  EK100 equally attenuated NF-κB gene expression and TLR4/NF-κB inflammatory pathway in Drosophila.

Figure 4: X-ray diffraction-based crystal structure of the IκB-α/NF-κB complex with solvents (PDB ID: 1NFI) at resolution of 2.70 Å, showing various chains in colour formats representing different molecules employed in the current study. For example, chain C (shown as a green cartoon model) represents NF-κB-P65, chain D (shown in a cyan cartoon model) represents NF-κB-P50, and chain E (shown in a magenta cartoon model) represents IκBα. (A) Structure of the complex formed after docking of the drug molecule with NF-κB. The highlighted region represents the binding pocket of the drug molecule in the NF-κB molecule. The binding pocket is represented by the labelled residues (G209, D210, E211, D257, R253, R259, S269, E285, etc., of the NFκB molecule and I102, N182, Q183, H184, etc., of the IκBα molecule). (B) Structure of the complex formed after protein-protein docking of the SARS-CoV-2 spike glycoprotein (closed state) (PDB ID: 6VXX, shown in orange colour at a resolution of 2.80 Å) with the NF-κB molecule. The highlighted region represents the binding pocket of the spike protein in the NF-κB molecule. The binding pocket is represented by the labelled residues (G209, D210, E211, D257, R253, R259, S269, E285, etc., of the NFκB molecule and N182, Q183, H184, etc., of the IκBα molecule). This information indicates that the drug molecule binds in the same pocket that the spike protein used to interact with NF-κB, possibly to activate its translocation to the nucleus. (C) Structure of the complex formed after protein-protein docking of the SARS-CoV-2 spike glycoprotein (shown in orange colour) with the NF-κB molecule (already docked with the drug molecule). The highlighted region represents the binding pocket of the spike protein in the NF-κB molecule in the presence of the drug molecule. The drug molecule occupies the same binding pocket on NF-κB that is also used by the spike protein during docking interaction, thus inhibiting the spike protein from interacting with NF-κB at its designated site. In the interim, the spike protein interacts with NF-κB at a different site and hence may not be able to activate it. Currently, these assumptions need validating experimental proof, which is lacking because of the strict restraint on experimentation with live samples from active human COVID-19 cases.
Figure 5: A 3-D structural alignment of the NF-κB-drug complexes (represented in a red cartoon model) and spike-protein-NF-κB-drug complexes (represented in a green cartoon model), with the NF-κB molecule represented in an orange cartoon model. (A) These alignments show the conformational changes imposed in the structure of the NF-κB molecule upon binding of the drug and spike protein. (B) Sequence-based comparison of the interacting residues of the NF-κB molecule with the drug sulindac sodium (yellow coloured bar), IkB molecule (green coloured bar) and spike protein (red coloured bar).

The changes in the 3-D structure of the NF-κB-IkB complex due to the binding of the drug molecule and spike protein in the presence of the drug molecule were compared to that of the unbound complex (Figure 5A). The binding of the drug molecule to the NF-κB-IkB complex led to an increase in the compactness of the complex. The area and volume data of the pockets presented in Figure 5A corroborated this observation. The binding of the spike protein in the presence of the drug molecule further increased the compactness of this complex, possibly due to increased interaction between the chains of the NF-κB-IkB complex. Additionally, we made a comparison of the amino acid (aa) residues of the spike protein, MAP kinase and pLC gamma during their interactions with the residues of the NF-κB molecule to identify if any common pattern existing in aa selection by sulindac during these interactions. However, all three sets of interactions, such as NF-κB-sulindac docked with spike protein, NF-κB-sulindac docked with MAPK and NF-κB-sulindac docked with pLC gamma had different aa residues interacting with the chosen ligand (sulindac)(Figure 6), revealing all these as independent events. In our studies, MM-GBSA studies on docked complexes revealed dG bind values that were reasonably in agreement with ranking based experimental scores, i.e., binding values. We first calculated the MMGBSA dG Bind = Prime Energy (Optimized Complex) − Prime Energy (Optimized Free Receptor) and second, MMGBSA dG Bind(NS) = Prime Energy (Optimized Complex) − Prime Energy (Ligand Geometry From Optimized Complex) − Prime Energy (Receptor Geometry From Optimized Complex). The values were found to be −2.15 kcal/mol for MMGBSA dG bind and −2.71 kcal/mol for MMGBSA dG bind (NS). The atoms contributing to the binding affinities are shown in Figure 7.

Our understanding on the involvement of NF-κB signalling pathway in COVID-19 is limited, but NF-κB inhibitors have been increasingly utilized to gain therapeutic benefits in many human diseases [49, 50]. Abnormal activation of NF-κB in the manifestation of SARS-CoV-2 infection is reported for the pathogenic profile of immune cells, cytokine storms and multi organ defects and cytokine storm alone is capable of triggering excessive inflammatory response to SARS-CoV-2 and is extremely responsible for disease severity [51]. Thus, pharmacological inactivation of the NF-κB, such as proposed currently in the use of sulindac, NF-κB-kinase interaction either directly or by inhibiting the dephosphorylation of its inhibitors, such as IKKα, IKKβ and other associated molecules, strongly represents a potential
therapeutic target to treat the symptoms of COVID-19 in clinical trials. A survey of literature since the emergence of COVID-19 clearly reveals that majority of the drugs formulated to tackle SARS-CoV-2 thus far, have been structured to target virus [52]. However, given the current situation where the virus mutates so rapidly, our conceptual paradigm on targeting the key viral proteins through drug intervention need to get changed towards the host proteins, such as human proteins that hold key portfolio in SARS-CoV-2 – mediated infections. This would possibly take care rapidly, our conceptual paradigm on targeting the key viral proteins through drug intervention need to get changed towards the host proteins, such as human proteins that hold key portfolio in SARS-CoV-2 – mediated infections. This would possibly take care of the various mutants of SARS-CoV-2 that have been difficult to control until now. Furthermore, cytokine storm in COVID-19 [53] and various drugs that were initially used for therapy [54] need to be revisited in light of modern researches based on drug repurposing, which are potentially very effective in treatments of even the anomalies arising post-COVID-19 infection and can thus counteract the many aberrant responses thereof.

Figure 7: MM-GBSA energy based visualization of docked complex highlighting the docked complex of receptor and sulindac sodium are represented with colour highlights on the ligand atoms showing contributions towards binding affinity. Blue represents the receptor and adaptive colors on ligand from Red to Green showcasing higher negative values to least values, respectively. The yellow dashed line represents the hydrogen bond with the receptor.

Author contributions:
S.A.: MDS analysis, Original Draft; P.B.: Docking analysis and Report; J.K.: Docking analysis supervision, Review and Editing; R.P.: MDS analysis confirmation and Report; D.M.: Review and Editing; A.U.: MM-GBSA analysis; V.N.: MM-GBSA report; V.M.: Conceptualization, Revision and Editing, Conclusion and Discussion, Formatting the MS, Overseeing the project and the MS

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