

Title page

Predicting and repurposing of drug and drug like compounds for inhibition of the covid-19 and its cytokine storm by computational methods

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Abstract

Background: The SARS-CoV-2 virus is a new highly pathogen virus that can suppress cell immune response (Poly ADPR process, and interferon releasing in infected cell) by viral macrodomain and create fatal pneumonia. Cytokine storm phenomenon which can create consequence of SARS COV-2 by increase of the performance of human TRPM2, is a dangerous condition for human and its body organ such as kidney, heart, and liver. it seems like prevention of beginning of these events is necessary. **Material and methods:** in this study drug and drug like database was used for inhibition of the viral macrodomain and human TRPM2 that can initiate of the viral cell cycle infection and human cytokine storm respectively. Ligand base drug designed, pharmacophore modeling, docking and molecular dynamic simulation was performed **Results:** among up to billion compounds, 20 compounds were screened by infinisee program. Then 7 compounds were selected by pharmacophore modeling. between 7 compounds, 5 compound was removed because of high similarity to ADPR and possibility of toxic effect for human RNA and DNA polymerase. One compound was losartan and selected for docking and molecular dynamic simulation. **Conclusion:** losartan earned a proper dock score and binding affinity to create the complexes with TRPM2 and macrodomain. Molecular dynamic simulation has shown losartan had an adequate binding free energy for human trpm2 and viral macrodomain. The inhibitory effect of losartan on these protein has shown and it could interfere in several points (PARP, PARG- macrodomain and TRPM2) and decreases oxidative stress, apoptosis, cytokine storm in COVID-19.

Key words: Viral Macrodomain; COVID-19; TRPM2; losartan; Bioinformatics

Introduction

At the end of 2019, the emergence of the novel coronavirus infection in Wuhan, China, it has rapidly spread across China and many other countries. The International Committee on Taxonomy of Viruses has been named SARS-CoV-2 and the disease it causes has been named coronavirus disease 2019 (COVID-19). The SARS-CoV-2 virus is a new pathogen that is highly contagious, which can spread quickly [1]. This virus must be considered capable of causing enormous health, economic and societal impacts with their setting The SARS-CoV-2 virus like SARS-CoV and MERS-CoV, is a beta-coronavirus which all of them have originated from bats. On 30 January 2020, the World Health Organization (WHO) declared that the SARS-CoV-2 outbreak constituted a Public Health Emergency of International Concern [2]. The outbreak has rapidly evolved, and more than 180 countries worldwide have reported laboratory-confirmed cases of SARS-CoV-2 [3].

Coronaviruses (COVs) are positive-sense enveloped RNA viruses which are larger than the other RNA viruses, encoding 16 non-structural proteins (nsps). They have different proteins that are important for preventing and blocking the innate immune response, including enzymes such as an endoribonuclease (nsp15), an O-methyl transferase (nsp16), and an macrodomain with ADP-ribosyl hydrolase (nsp3) [4]. The macrodomain of nsp3 harbors several conserved domains in which some enzymatic activities have been attributed. COVs macrodomain has a close functional association with poly (ADP-ribose) polymerases (PARPs), which is known as the regulator of apoptosis in eukaryotic cells. PARPs regulate important cellular processes such as cell-cycle progression, cell division, host-virus interaction, DNA damage repair, genome maintenance, transcription regulation, protein degradation, ageing, and cell death. PolyADP-ribosylation can be removed by the action of two divergent enzymes, poly (ADP-ribose) glycol hydrolase (PARG) and ADP-ribosyl hydrolase 3 (ARH3) [5,6]. PARG is a catabolic enzyme that cleaves

ADP-ribose (ADPR) polymers that synthesized by members of the PARPs family of enzymes. The Viral macrodomains involve in cellular signaling consist of; ADP-ribose-generating or-consuming pathways and roles in determining RNA synthesis, replication, and virulence. This protein hydrolase ADPR polymer from ADP-ribosylated proteins. ADPR-binding to the CoVs nsp3 macrodomain that is necessary for the initiation of virus replication, whereas the hydrolase activity facilitates to amplify the replication complex. Therefore, macrodomains can remove ADP-ribose polymer from modified proteins [7].

The antiviral properties of PARPs are regulated by interferon (IFN), and under evolutionary selection, it has been given that primary function of viral macrodomain is counter the host response viral infection. Accordingly, macrodomain by removing ADP-ribosylation from modified host proteins, prevent host responses to viral infection[8].

On the other hand, PARG activity of MD leads to increase the level of free ADPR in the cell, which activates the Transient Receptor Potential 2 (TRPM2). TRPM2, a subtype of TRP-melastatin subfamily (TRPM), is a multifunctional Ca^{2+} permeable and nonselective cation channel. Exposure to hydrogen peroxide (H_2O_2), tumor necrosis factor α (TNF α), reactive oxygen and nitrogen species is generally accepted to promote channel activation through the generation of ADPR, an agonist for TRPM2 (9). It performs as an important Ca^{2+} signaling regulator in a variety of cells, contributing to cellular functions including cytokine production, insulin release, cell motility, oxidative stress and cell death. TRPM2 activation cause to induce Ca^{2+} influx, and providing positive feedback for channel activation, led in necrosis and cell death (9). The Severe inflammatory response, cytokine storms, and necrosis in the patient's lungs with COVID-19, can explain the activation of TRPM2 signaling pathway, which can result the action of macrodomain or stress conditions that lead to accumulation of ADPR. Therefore, both macrodomain and TRPM2 inhibitors blocked calcium influx through TRPM2, inhibiting the

cytokine storms, tissue necrosis, protecting cells from plasma membrane damage and cell death [9,10].

Losartan is an angiotensin-converting enzyme (ACE) inhibitors that used to treat hypertension. ACE inhibitors are used for a similar indication but associated with a cough [11]. Here, we showed that Losartan is a functional ligand for inhibition of the TRPM2 and macrodomain and has a high affinity to them. The comparative data have shown Losartan is effective agent for inhibition of the TRPM2 and macrodomain.

Methods

Protein and ligand preparation

High-resolution X-ray diffraction SARS CoV-2 of macrodomain binding ADP ribose phosphate and human TRPM2 protein structures were downloaded from protein data bank (<https://www.rcsb.org/pdb>). The exported files included 6W02 consists of two COVID-19 macrodomain molecules complexes with ADP ribose phosphate as the main ligand. Because macrodomain has only one monomer, we elected, chain A of the PDB structure, and other additional parts were omitted. Another exported file, 6PUR contains four TRPM2 molecules complexes with ADP ribose phosphate. Among the four molecules, chain A was selected, and extra parts were removed. In chain A, sequence between residue 1234 to 1534 was selected because it forms binding site of TRPM2, and this sequence which has been located in the cytoplasmic part involved in the cell cascade [12,13].

All hydrogen atoms were added by the Discovery Studio software 2.5 (DS, Accelrys Inc, San Diego) and the proteins were prepared for the docking and virtual screening. Energy minimization was performed through a simulation module of Discovery Studio by conjugate gradient method and CHARMM force field [12,13]. This process was done with energy gradient

of $0.1 \text{ cal}\text{\AA}^{-1}$. Ligands' 3D structures were prepared and minimized by the Marvin Suite software [14].

Chemical library

The procedure of this study shown the Figure 1. The virtual screening was done to recognition of the hit compounds from the GalaXi_2019-10, KnowledgeSpace_2019-05, and REALspace_2019-12 databases in biosolveit (<https://www.biosolveit.de/>) with infinisee software [15], for finding similar compounds to ADP ribose phosphate. The all current databases collections include up to 2.9×10^{14} compounds, which consist of every compound on the world, but in this study on the base of the necessity and urgency of investigation about coronavirus treatment was more focused on the GalaXi_2019-10 database that has drug-like and all drug library (other databases is not removed).

Primary virtual screening was performed over all databases by the infiniSee software (ligand base, similarity 30-100), resulted in to obtain 20 compounds. This collection was imported to the ligandScout software and converted to ligandScout library, then, on the base of the ADPR pharmacophore model (two hydrogen donors and six hydrogen acceptors) (figure2) secondary virtual screening was done. After that, seven compounds were obtained by high pharmacophore-fit score. Figure 2 Pharmacophore model of molecules that are similar to ADPR and pharmacophore of ADPR

Five of seven obtained compounds, were a structural analog of ADPR or other nucleotides, so these compounds were removed due to the potential toxic effects on human DNA and RNA polymerase. One compound was sulphonamide which was removed due to not being FDA-approved drug. The final compound, i.e. Losartan was finalized for the molecular docking step.

Molecular docking

In this step, we used SeeSAR 9.2, AutoDock Vina module of ligandScout 4.3, and Molegro Virtual Docker v6.0 to investigate the conformational of Losartan in interaction with macrodomain and TRPM2 proteins [16,17]. At first SeeSAR 9.2 was used for a primary docking of Losartan. The applied parameters were used the genetic algorithm. The population size, selection pressure, and number of run were 100, 1.2 and 100, respectively. Conformations that were obtained in this step were used for final docking with Molegro Virtual Docker and AutoDock Vina, and the best affinity score was estimated between our load conformations [18,19]. The population size and the number of run in this step were 500 and 200, respectively. Profiles of interactions between Losartan and proteins were obtained via Discovery Studio Visualizer and LIGPLOT software [20].

Molecular dynamic simulation

The molecular dynamic simulation was performed by AMBER18 package [21], and LEAP module of AMBER18 that was used for adding hydrogen to atoms. The ligand charges were generated by AM1-BCC method of Antechamber module. The topology of macromolecules was created by AMBER99SB-ILDN force field and tleap module of AMBER tools [22,23]. The generalized AMBER force field (GAFF2) was used for the ligand topology building with Antechamber module of AMBER18. Then the complexes were solvated in a truncated octahedral box of TIP3P. The water molecules with a buffer size of 10 Å, were neutralized by Na⁺ or Cl⁻ ions for each complexes. The default protonation of Amber18 was used for titratable residues in protein. The energy minimization and MD simulations were used for each system with the sander module. Ten thousands cycles of minimization were performed with the steepest descent; and conjugate gradients for all parts of the systems were done to remove the bad steric interactions and get better minimum energy. Then, the position restraints at the constant volume

(NVT) for 100 ps, with a restraint force of 10 kcal/mol at temperature of 100 K, and then at constant pressure (NPT) for 100 ps, with a restraint force of 1 kcal/mol at temperature of 300 K, were performed. Afterward, an equilibration step at NPT was performed for reaching to equilibrium in density for 100 ps, while the restraint force was removed. This method allows the water to equilibrate around the complexes. In the position restraints and equilibration steps, Langevin dynamics were used to control the temperature. In final step, 20 ns (nanosecond) MD simulation was used for each protein-ligand complex. For analyses of MD simulations, CPPTRAJ of AMBER18 was used [24].

Binding energy calculation

An attractive and interesting method in Computer-Aided Drug Design (CADD) is Molecular Mechanics Poisson Boltzmann/ Surface Area (MM-PBSA). The binding energies ($\Delta G_{\text{binding}} = G_{\text{complex}} - G_{\text{protein}} - G_{\text{ligand}}$) in this study were calculated with the MM-PBSA method by Amber18 [25]. The accuracy of docking was calculated through obtaining the binding free energy over the molecular dynamics simulations. The last five ns of MD simulation trajectories were used for calculation of binding free energy [24].

After calculations of Gibbs energy changes (ΔG) and entropy changes (ΔS) values by the amber18 package, the enthalpy change (ΔH) was calculated ($\Delta G = \Delta H - T\Delta S$).

If the enthalpy change (ΔH) was negative numbers, the reactions between ligands and proteins would be exothermic.

Results

The virtual screening for macrodomain and TRPM2

After the primary virtual screening on the database by the infiniSee software, a collection containing of 20 ligands was obtained table1. Then, seven compounds were screened after

pharmacophore modeling by ligandScout with pharmacophore-fit score 86.16 - 85.56 (table2 and fig2). Losartan (with pharmacophore-fit score 85.26) was between of them, and other compounds because were nucleotides or similar to nucleotides (such as guanosine ribose phosphate and other nucleotides or their analog) were removed, Losartan was selected as a hit compound.

The docking results

ADP ribose and Losartan in the macrodomain

The docking score ADP ribose phosphate and losartan in the Molegro Virtual Docker software for for macrodomain was calculated -207 and -197, respectively (table3). The similar scores were also obtained using SeeSAR and Autodock Vina software tools. Losartan formed six hydrogen bonds with four amino acids, Phe132, Ser128, Gly180, and Ile131 by the distance between 1.83 and 2.44 Å. Besides, 20 hydrophobic contacts were exhibited with Asp22, Ile23, Phe156, Ala154, Val155, Lue164, Pro136, Pro125, Gly48, Gly51, Gly47, Gly46, Asn40, Ala39, His45, Leu127, Gly133 and Ile131 among the range 3.40-3.90 Å. The highest hydrophobic interaction belonged to Ile131 with three hydrophobic interactions (figure 3 and table4).

ADP ribose and Losartan in the TRPM2

It was shown that ADP ribose phosphate, and losartan docking scores in the Molegro software were -149 and -157, respectively (table3). These ligands had a similar arrangement of score in both proteins in the Seesar and Autodock vina software tools. Losartan was exhibited four hydrogen binds with Asn1487, Arg1433 and Ser1382 residues within the range of 1.8 –2.8 Å. A total of 13 hydrophobic bonds were found with Pro1388, Val1377, Leu1379, Ser1382, Ala1386, His1348, and Tyr1349 in the range of 3.1-3.9 Å. The highest hydrophobic interactions were observed in Ala1386 with three contacts (figure 4 and table 4).

Molecular dynamic simulation

For checking the stability of the protein-ligand complexes, 20 ns molecular dynamic simulations were used to all complexes. Figure 5 exhibits the RMSD value for every complex of trpm2 and macrodomain structures on the simulation time. Assessment of average last five ns of RMSD plots showed that they are relatively stable backbone trajectories of TRPM2-Lsr (average 3.44Å, fig5B) and macrodomain-Lsr (average 2.16, fig5A) as compared to TRPM2-ADPR (average 4.2Å, fig5B), and macrodomain-ADPR (average 1.58Å, fig5A) complexes. In addition, it can be inferred that creating losartan complex with these proteins is led to the stability of protein and it is a useful ligand for inhibition of them.

Figure 6 shows RMSF profile of these complexes. The residues of macrodomain have lower fluctuation in complex with ADPR relative to Losartan, but there are some peaks in the plot that are amino acids in the active site and had similar stability and same fluctuation as compared to macrodomain-ADPR fig6A, 6B, and. So it has concluded Losartan can contest with ADPR for binding macrodomain and create resemblance stability

In the case of TRPM2 protein, the Losartan induces less fluctuation or more stability relative to the ADPR inhibitor. (Figure 6B). These results are compatible with RMSD results (figure 5). In other words, despite of the more stability of the ADPR than Losartan, Losartan can compete to ADPR for creating a complex with macrodomain protein, and in the case of TRPM2, Losartan could be a proper inhibitor for TRPM2. The free binding energy and thermodynamics parameters of all ligands in complexes have been mentioned in table 5, it has shown that ADPR has more binding affinity for macrodomain, and Losartan has more binding affinity for TRPM2 protein. The binding between proteins and ligands were exothermic. These results also are similar to RMSD and RMSF results.

Discussion

The worldwide spread of the coronavirus became a severe and significant threat to human health. World health organization have announced that the coronavirus becomes a pandemic disease[2,3]. As yet, there is not any precise treatment for coronavirus. In some articles have suggested that some of the protease inhibitors, remdisivir, and chloroquine are effective on the

COVID-19, but have not shown an efficient and potent therapeutic effect on this virus. This limitation and threat made us work on some drug databases that are FDA approve and some drug-like databases. In the primary virtual screening by infinisee, the new bank had created by some molecules with relative similarity to ADPR. Next, pharmacophore modeling was shown seven molecules that have similar pharmacophore. Between them, because five compounds were analog of nucleotide, they were omitted. Nucleotides analog have toxicity for human cells and bodies, so were removed.

Criteria for selection of the ligand was six terms consist of:

- 1- visibility
- 2- FDA approves existed drugs
- 3- Availability
- 4- cost-benefit effective compounds
- 5- the relative similarity to ADPR but not capability to binding to other nucleotide-binding enzymes (on the other hand compound have selective toxicity for other enzymes, and their effect had been proved)

Above criteria and capability creating complex with macrodomain and TRPM2 (that assessed by docking, and MD simulations), Losartan was chosen as an effective ligand for these proteins. Results that were obtained from docking and molecular dynamic simulations emphasized that Losartan is a functional ligand for inhibition of the TRPM2, macrodomain and has a high affinity to them. Thermodynamic parameters revealed that losartan complexes most likely created spontaneously. Losartan had a decent dock score and binding affinity for creating complexes with TRPM2 and macrodomain. Besides, it can contest with ADPR for binding to these proteins and could disable the performance of the proteins. So TRPM2 and macrodomain became inactive and other signaling pathways that depend on them may be deactivated.

Xiao Luo and *et al.* worked on the TRPM2 inhibitors by two novel ligands that were synthesized as ADPR analogs [26]. Xiao Luo's work was done without attention to the possible toxicity of these compounds for every enzyme (enzymes that have a high affinity for binding to ADPR, such as DNA-polymerase or RNA-polymerase). Xiao lue and *et al.* had focused only on the synthesis of these compounds. That study conceded ADMET investigation and laboratory assays for these ligands in the future studies for other scientists.

Our study against the Xiao Lue *et al* study, was done with Precise focused on the possible toxicity, and utilization of the FDA approved drugs and drug-like database. The bioinformatic calculation was performed to increase the accuracy of screening methods and obtaining the best drug to inhibit the both proteins.

John G. Starkus and *et al.* studied scalaradial (natural product compounds) as an inhibitor of the TRPM2 with the experimental assay, but they have not any computational approaches [27]. Scalaradial is not an available compound and, more importantly, is not as an FDA approved drug. Then there is an urgent necessity for drug and inhibition of the TRPM2 and macrodomain for treatment of any person affected by COVID-19 [28].

Our knowledge Until now, there was not any study for the inhibition of the macrodomain and its function. This study is for TRPM2 and macrodomain inhibition by a bioinformatics approach with the utilization of the drug and drug-like database. Particularly as now, there is no study for inhibition any coronavirus's macrodomain (that has high affinity to bind ADPR). It seems the activity of MD protein of COVID19 runs a lethal cycle between PARP, MD and TRPM2 Zeinalian, M., *et al* explained this cycle and suggested that PARP, MD (NSP3 of virus) and TRPM2 are critical therapeutic targets in COVID19 [29].

In this study, it has been focused on the coronavirus pathogenicity and mechanisms that how the human cell can combat them. It was concluded that viral macrodomain has the essential roles for

the initiation of the cell infection and the virus-cell cycle. Macrodomain with poly ADPR degradation and increase of the free ADPR's levels can provide activation of the TRPM2 and prevent of the produce interferon-gamma so that cell would be susceptible to COVID-19 infection; therefore, it has been suggested as the target protein for efficient therapeutic approaches against the coronavirus infection.

Some different studies have shown that ACE inhibitors such as Losartan could have a protective effect on some tissue injury due to the systemic inflammatory diseases. Losartan has anti-aggregative platelet, anti-diabetic, and prevent end organ injuries and fibrosis. Also in the disease that affected with chronic kidney disorder, and blood pressure, prescription losartan led to decrease from kidney damage. In another study has shown that Losartan was able to prevent damage to the ischemic and oxidative stress and proliferation of tubules cells and penetration of the macrophage from parenchyma kidneys [30]. On the other hand, losartan could also reduce indirectly oxidative stress through inhibition of TRPM2. It leads to prevent cytokine storm, apoptosis, and cell necrosis. Losartan can also directly bind to TRPM2, and creates an inhibitory complex with it [31].

ADP-ribosylation alters the structure and functions of the protein and could be involved in several processes including genetic stability, DNA damage repair, chromatin remodeling, cellular stress response, and virus infection. PARG is a hydrolytic enzyme that commonly cleave the α (1''-2') *O*-glycosidic linkages in PolyADP-ribose chains. This enzyme is believed to have both exo- and endo-glyco hydrolase activities, acting at terminal and internal sites of PolyADP-ribose chains, respectively. Thus, in addition to free ADPR, PARG generates the short chains of PolyADP-ribose which as a powerful cell death signa. TRPM2 activation by short chains of PolyADP-ribose or free ADPR results in calcium influx which make the cells become susceptible to cell death; whereas, inhibition of TRPM2 function enhances the cell survival [32].

Conclusion

It seems Losartan can inhibit the viral macrodomain and TRPM2 so that it can interrupt the cell cycle of the virus and apoptosis, necrosis, fibrillation, and cytokine storm consequence of the COVID-19 infection. Therefore, Losartan, with preventing the induction of overexpression of the inflammatory cytokine, may decrease the severity of the disease and time of hospitalization. Presumably, Losartan with inhibition of these proteins activate the production of the interferon-gamma and intracellular defense against the COVID-19 virus, so enhancement resistance of person to the coronavirus.

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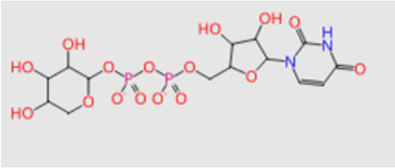
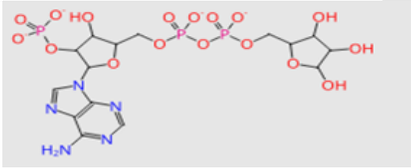
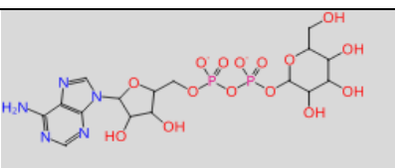
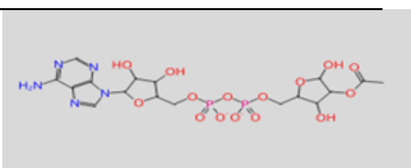
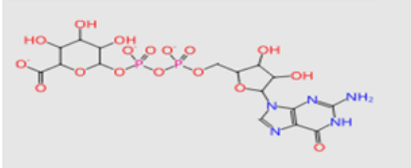
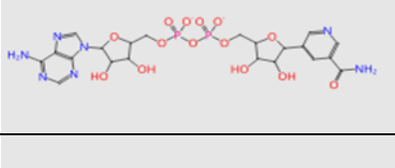
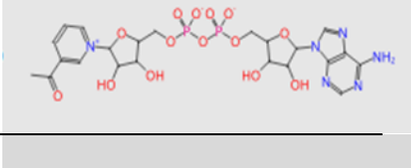
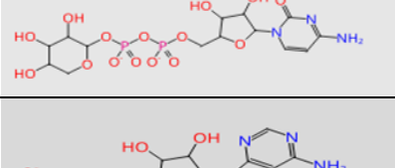
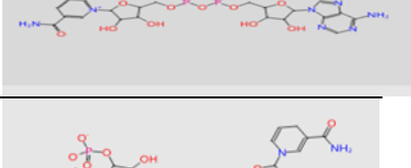
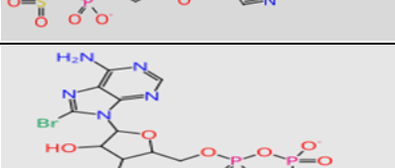
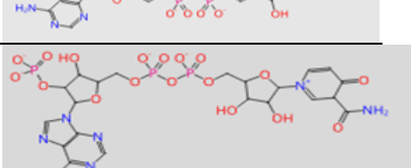
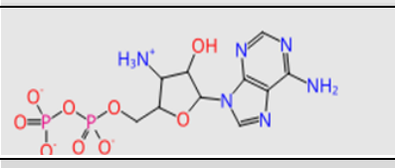
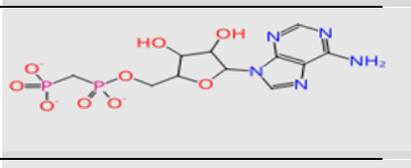
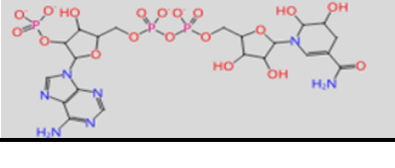
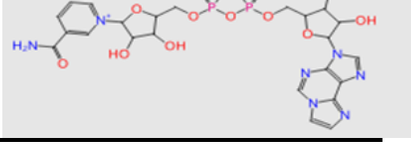
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Table 1. The molecular structures of 20 screened compounds by infiniSee program for coronavirus macrodomain and TRPM2

Compound's Names	Compound's structures	Compound's Names	Compound's structures
UDP-alpha-D-xylose		2'-Monophosphoadenosine-5'-diphosphoribose	
Adenosine-5'-Monophosphate Glucopyranosyl-Monophosphate Ester		Adenosine 5'-(trihydrogen diphosphate), P ¹ ->5-ester with beta-D-ribofuranose 2'-acetate	
Adenosine-5'-Monophosphate MethylGlucopyranosyl-Monophosphate Ester		GDP-alpha-D-mannuronic acid	
5-beta-D-ribofuranosylnicotinamide adenine dinucleotide		AZD-1236	
Cytidine-5'-diphospho-beta-D-xylose		Deamido-Nad	
Adenosine 5'-phosphosulfate		NADPH	
8-Bromoadenosine-5'-Diphosphate		4-Oxo-nicotinamide-adenine dinucleotide phosphate	
3'-Deoxy 3'-Amino Adenosine-5'-Diphosphate		Adenosine 5'-methylenediphosphate	
5,6-dihydroxy-NADP		Etheno-NAD	

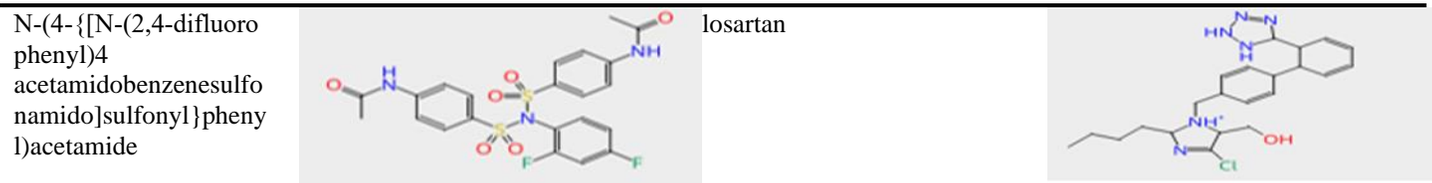


Table 2. six molecules that were removed after ligandScout virtual screening and Losartan selected for docking

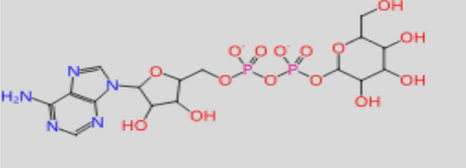
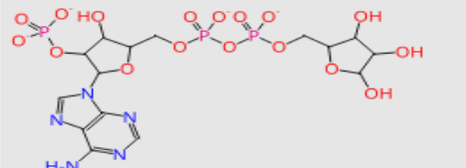
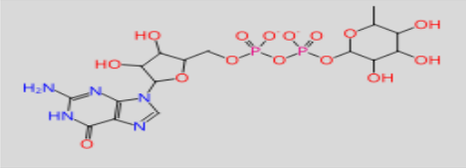
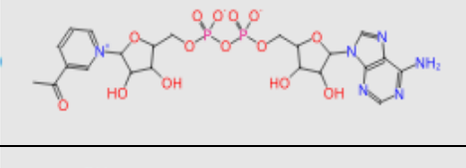
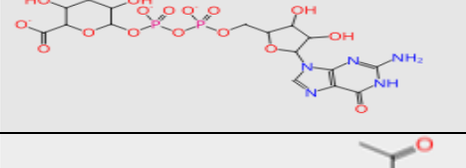
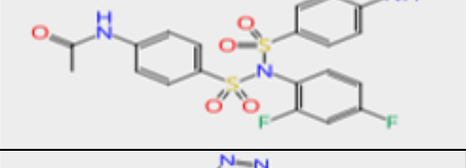
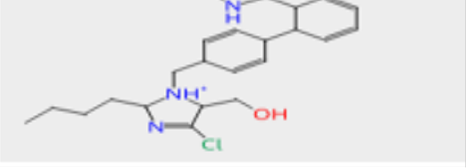
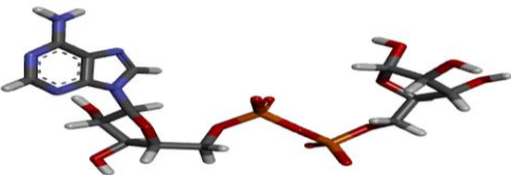
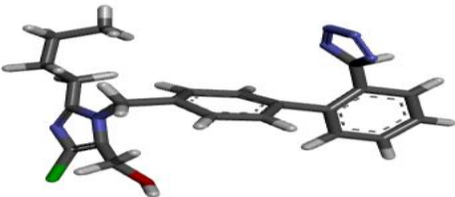
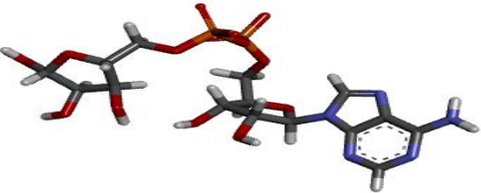
Numbers	Compound's Names	Compound's Names
1	Adenosine-5'-Monophosphate Glucopyranosyl-Monophosphate Ester	
2	2'-Monophosphoadenosine-5'-diphosphoribose	
3	Adenosine-5'-Monophosphate Methyl Glucopyranosyl-Monophosphate Ester	
4	AZD-1236	
5	GDP-alpha-D-mannuronic acid	
6	N-(4-([N-(2,4-difluorophenyl)4-acetamidobenzenesulfonamido]sulfonyl)phenyl)acetamide	
7	Losartan	

Table 3. The affinity and binding score of the ligand-protein complexes.

Complex	Structures	MolDock Score	AutoDock Vina kcal/mol
ADPR complex with macrodomain		-207	-22.50
losartan complex with macrodomain		-197	-20.30
ADPR complex with TRPM2		-149	-20.60
losartan complex with TRPM2		-157	-17.30

Docking score in Molegro Virtual Docker and docking affinity in the AutoDock Vina software tools. Docking score and binding affinity of the Losartan is very close to ADPR, even higher than the ADPR.

Table 4. Hydrogen bonding and hydrophobic interactions of the Complexes and distance of the

complex	Hydrogen binds		Hydrophobic interaction	
	Amino acids	Distance (Å)	Amino acids	Distance (Å)
Macrodomain complex with Losartan	Phe132, Ser128, Gly180, Ile131	1.83-2.44	Asp22, Ile23, Phe156, Ala154, Val155, Lue164, Pro136, Pro125, Gly48, Gly51, Gly47, Gly46, asn40, Ala39, His45, leu127, Gly133	3.40-3.90
TRPM2 complex with Losartan	Asn1487, Arg1433, Ser1382	1.8 – 2.8	Pro1388, Val1377, Leu1379, Ser1382, Ala1386, His1348, Tyr1349	3.1-3.9

interacting atoms between ligand and protein.

Table 5. Thermodynamic parameters of binding ligands to Macrodomain and TRPM2.

complex	$\Delta G_{MM-PBSA}$ (kcal/mol)	ΔS (kcal/mol)	ΔH (kcal/mol)
Macrodomain-ADPR	-90.87	-13.33	-4063.21
Macrodomain-Losartan	-26.51	-13.09	-3927.33
TRPM2-ADPR	-16.13	-13.10	-3919.93
TRPM2- Losartan	-39.99	-13.34	-4015.31

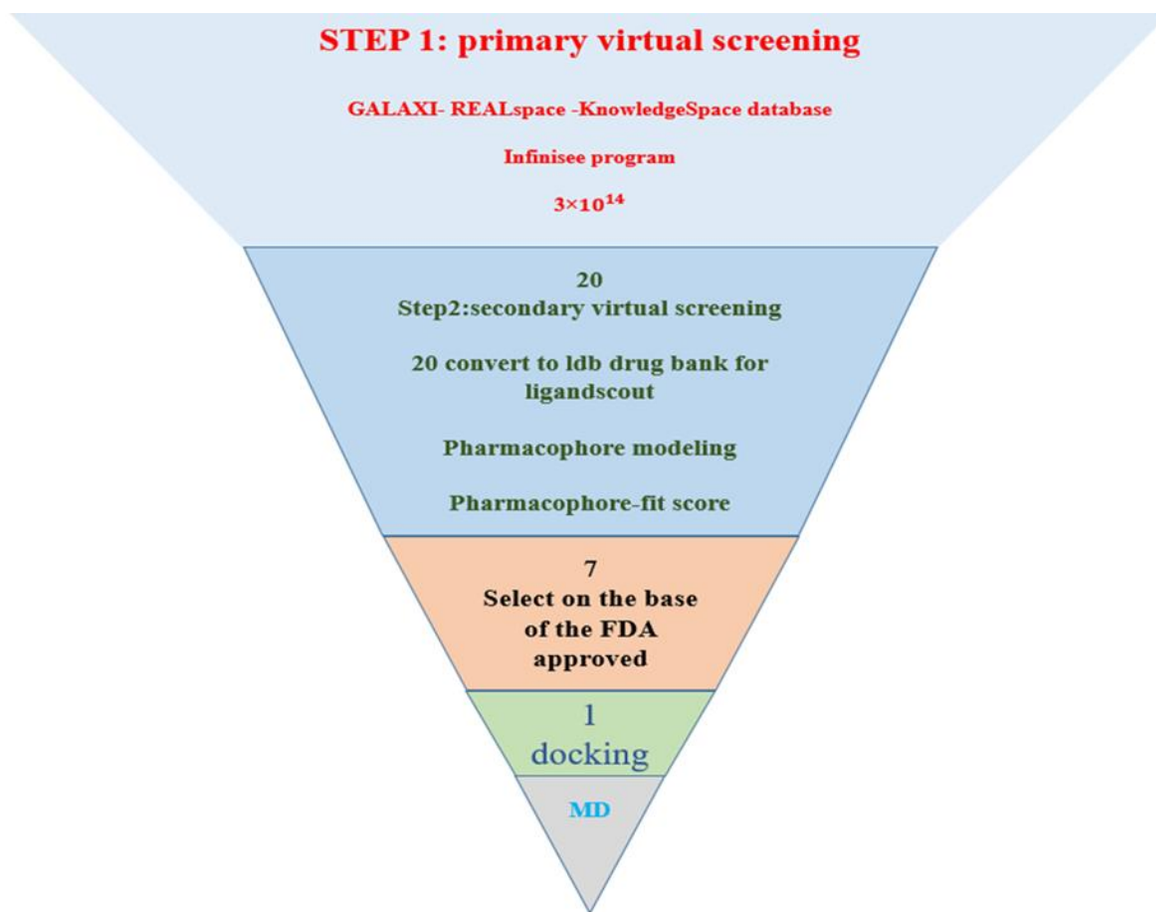


Figure 1. The procedure of this study. MD; Molecular dynamic simulation

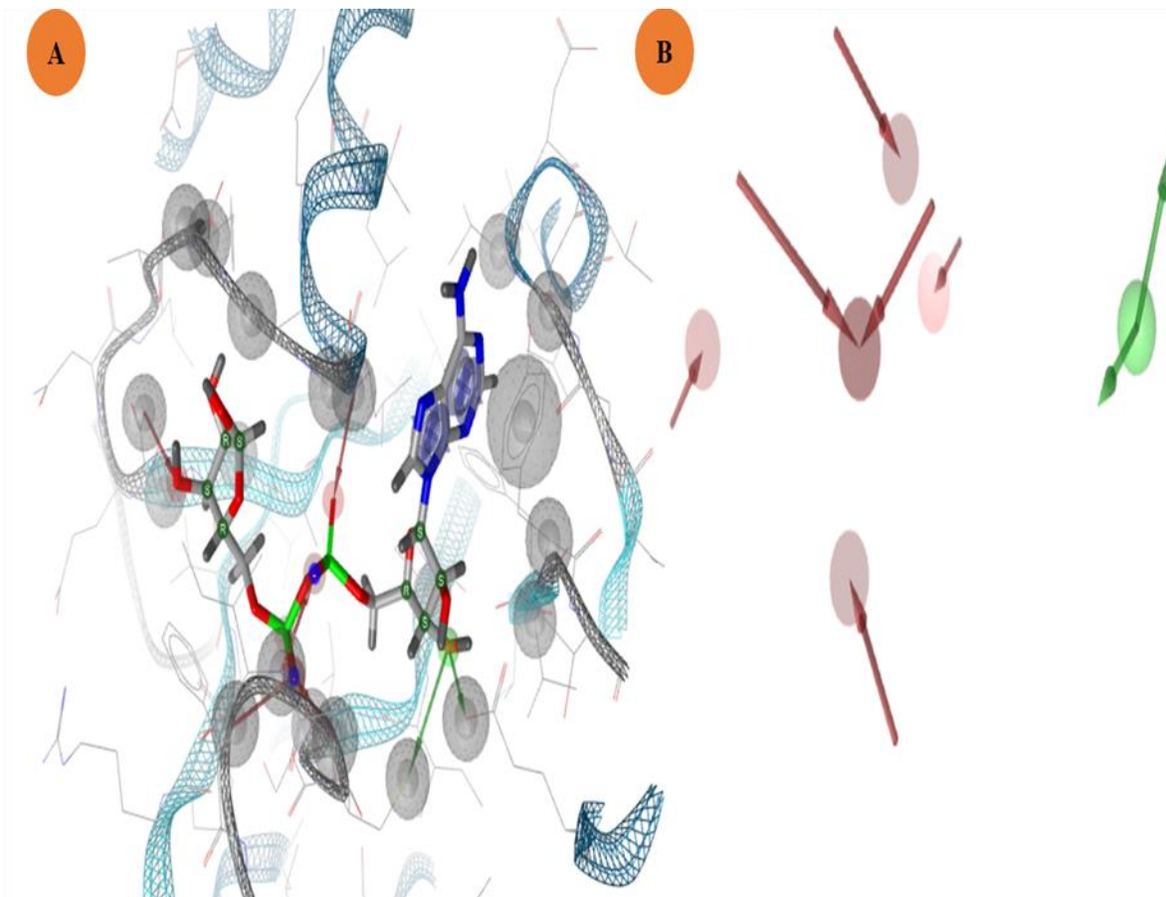


Figure 2. Pharmacophore model of ADPR. A-pharmacophore of the ADPR and its interaction with active site's amino acids macrodomain. B- pharmacophore model of ADPR which consist of two Hydrogen bond donor and six hydrogen bond acceptor

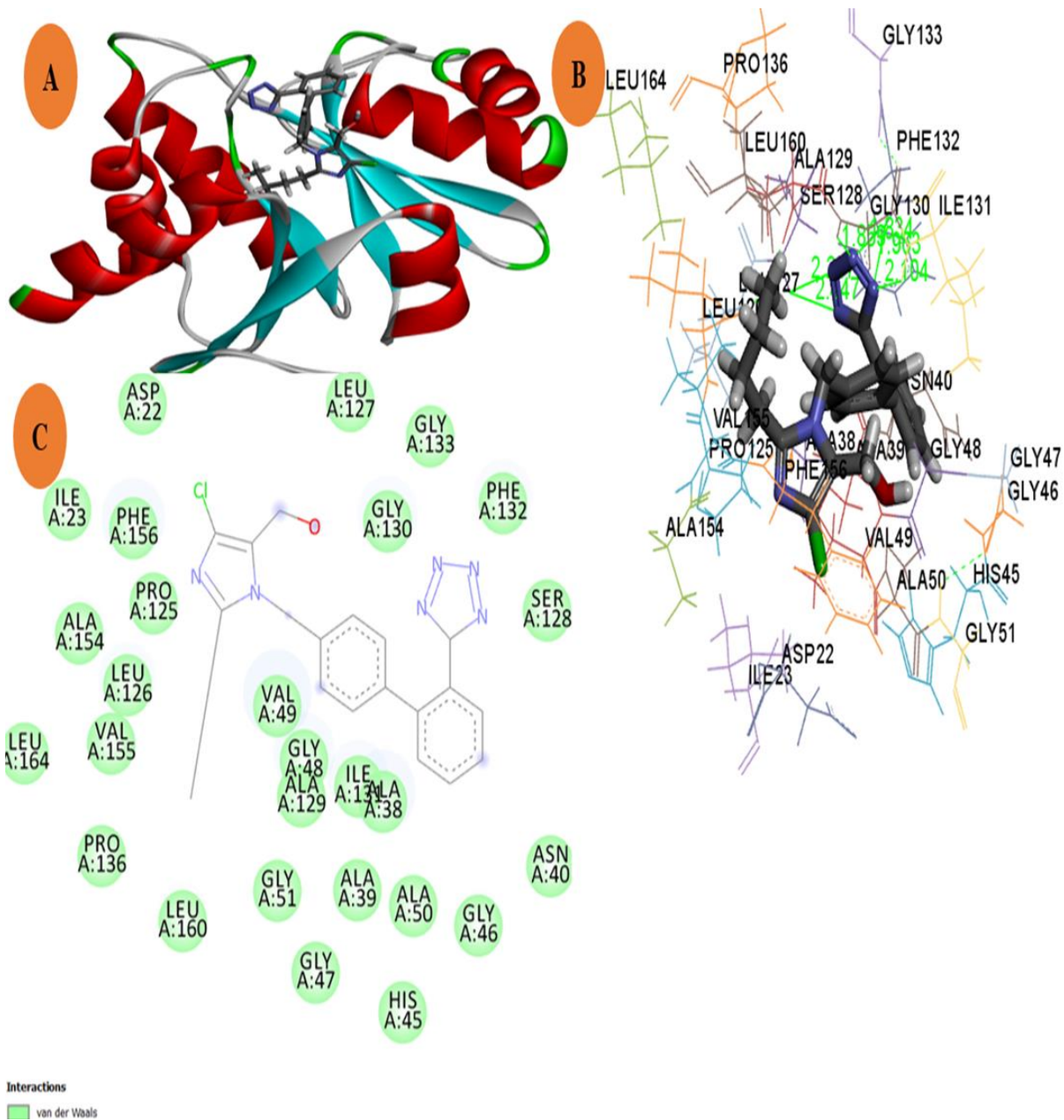


Figure 3. The predicted complex Macrodmain- Losartan binding mode. (A) The position of the Losartan in the active site of the macrodomain. (B) Hydrogen bond of Losartan with macrodomain in the active site (green line). (C) Van der Waals interaction between amino acids of active site and Losartan.

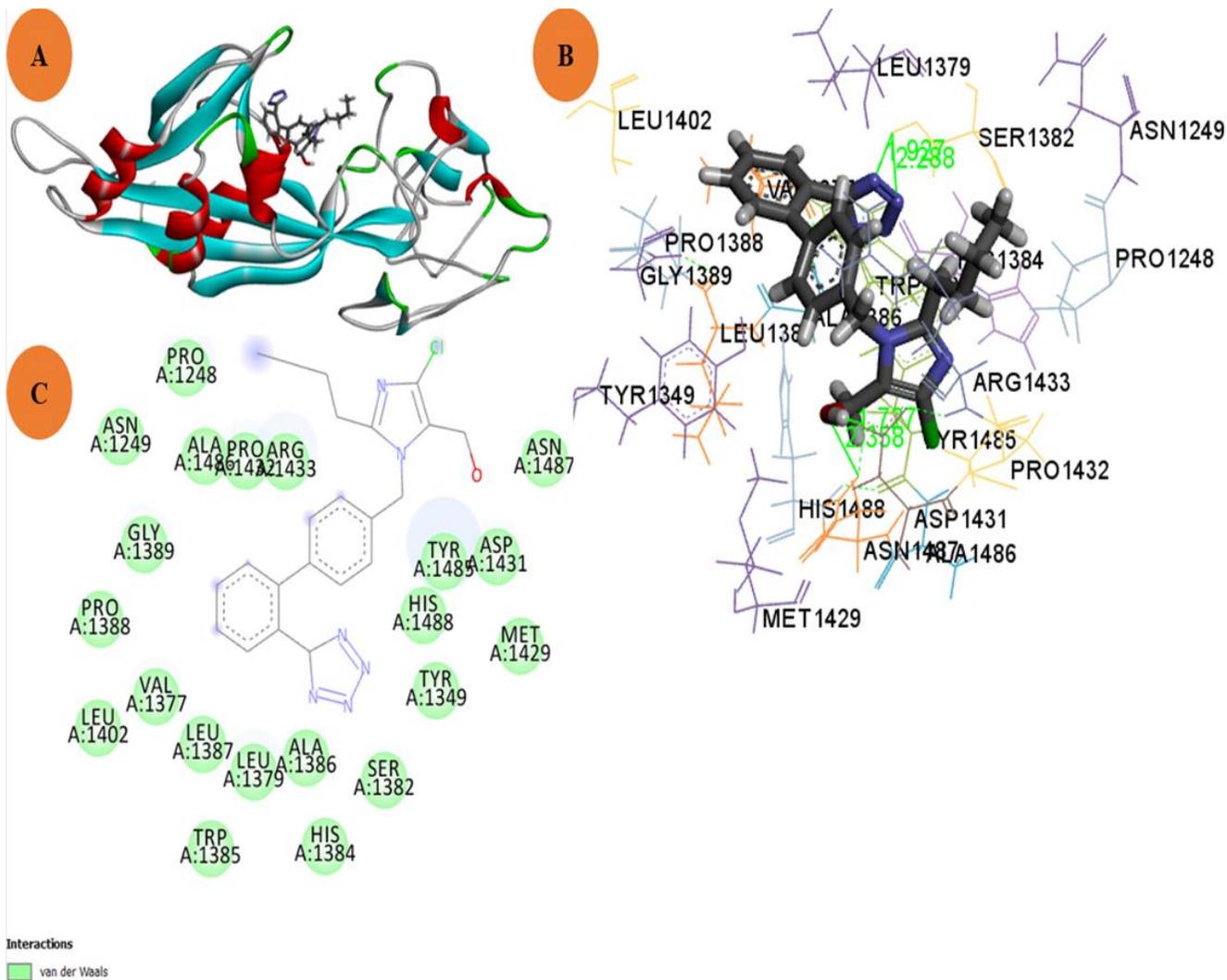


Figure 4. The predicted complex TRPM2-Lsr binding mode. (A) The position of the Losartan in the TRPM2'S active site. (B) Hydrogen bond of Losartan with TRPM2 in the active site (green line). (C) Van der Waals interaction between amino acids of the active site and Losartan.

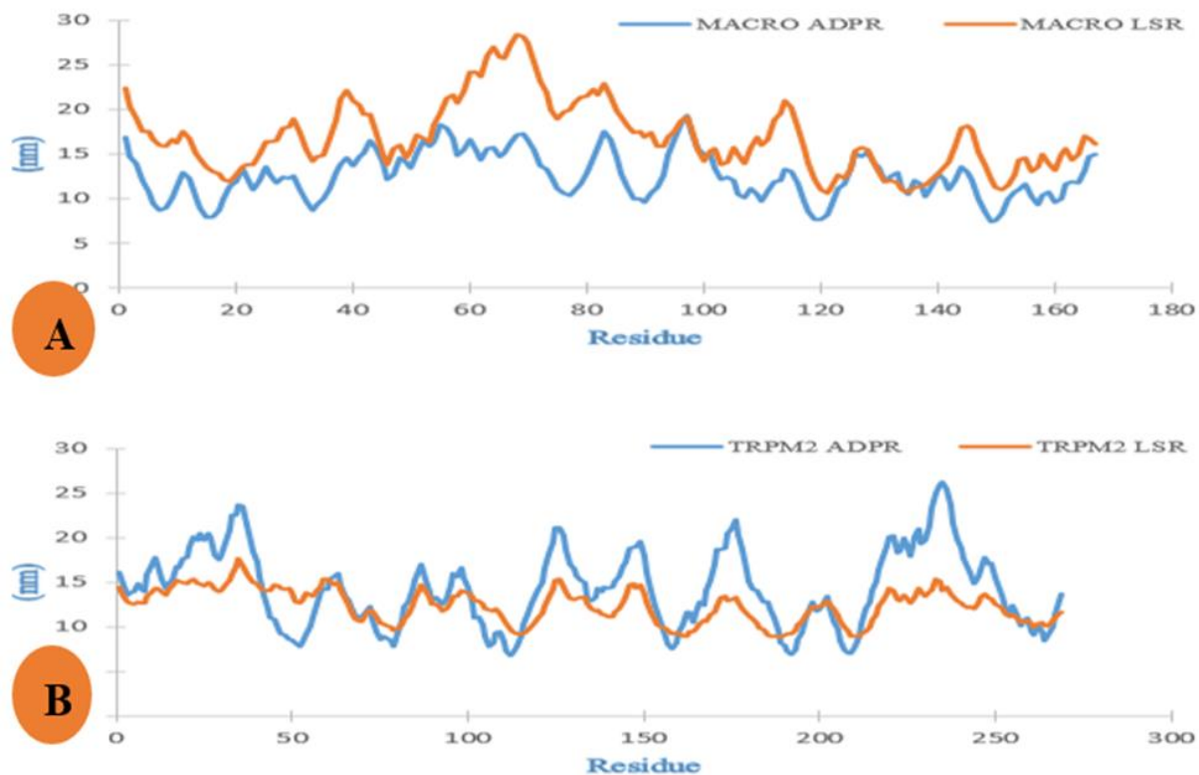


Figure 5. RMSD analyses of viral macrodomain (A) and TRPM2 (B) in complex with ADPR and Losartan.

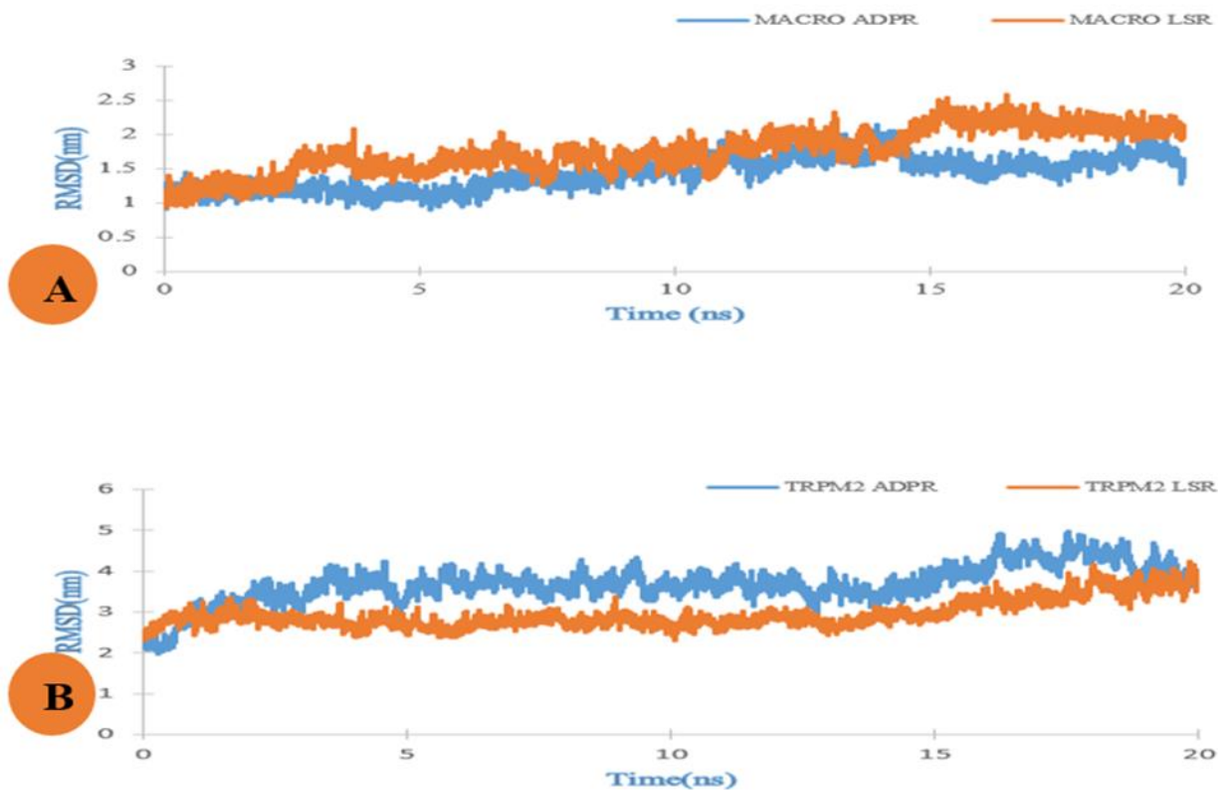


Figure 6. RMSF analyses of macrodomain (A) and TRPM2 (B) in complex with ADPR and LSR ligands.

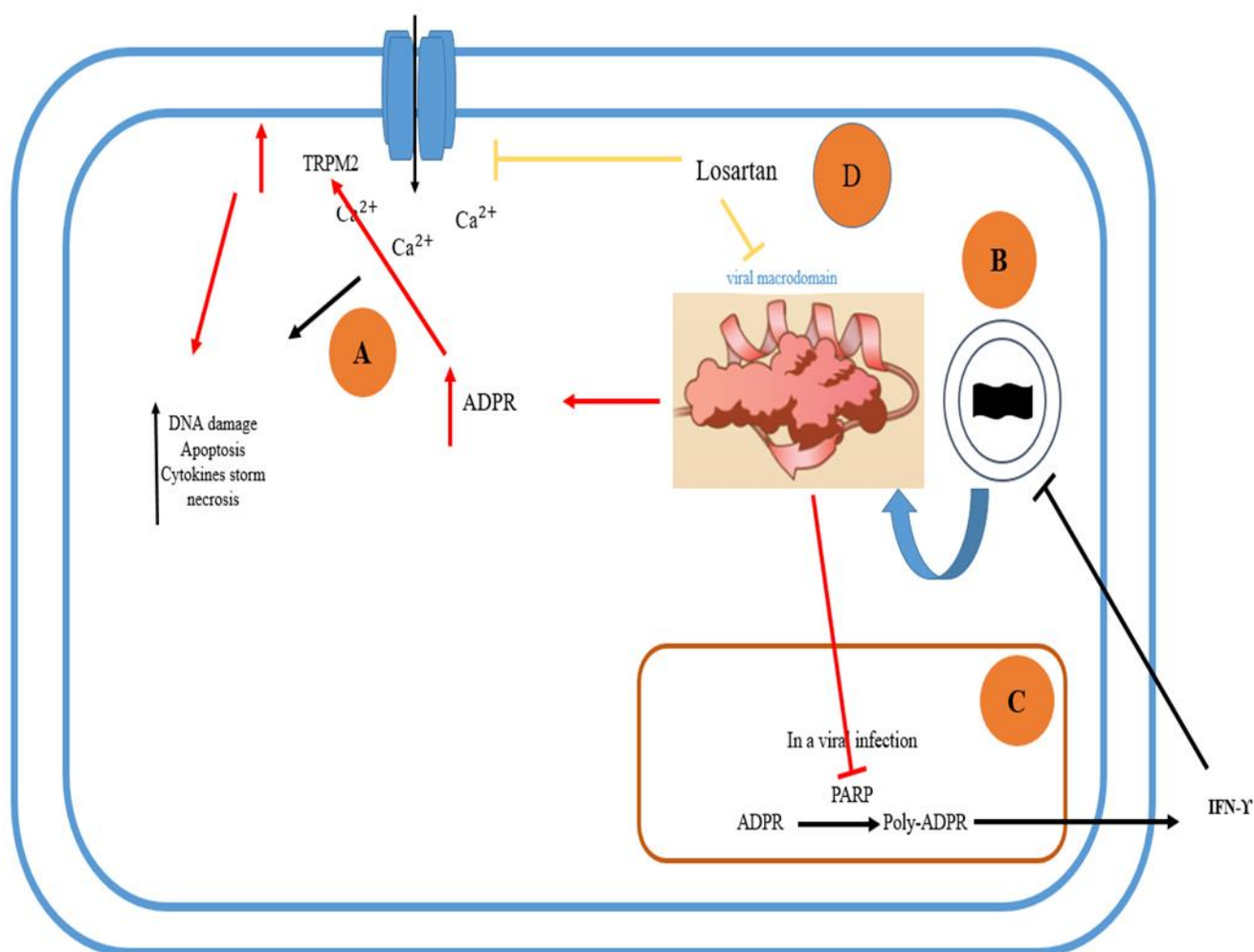


Figure 7. **A-Black arrows** are the cell mechanism in the normal situation (health people): TRPM2 can activate running of the cell death, and apoptosis signaling cascade. This protein is active in the high level of the ADPR in the cell. **C**-in a viral infection PARP enzyme active in the viral infection. PARP creates poly-ADPR and ADP-ribosylate different types of the protein in the cell. ADP-ribosylation of the proteins is the initiation for the Interferon gamma (IFN γ) signaling cascade, so IFN γ produce and release from the cell. The IFN γ can activate the human cellular immune response against the virus, and increase the resistance of the human cell in confronting with covid-19 virus. **B**- after the entering to the cell, virus RNA convert to the viral nsps proteins. One of these nsps is the macrodomain which degrade poly-ADPR from the protein, so prevent the IFN γ signaling in the cell, and increase ADPR volume in the cell. **A-Red arrows** show that increasing of ADPR in the cell activate the TRPM2 and its signaling cascade consequence of its activating, so cytokine storm, and apoptosis signaling cascade will initiate. **D**-losartan can bind to the viral macrodomain, and human TRPM2 and prevent their function, therefore cytokine storm will halt, and human cellular immune response will increase