High-Throughput Screening and Quantum Mechanics for Identifying Potent Inhibitors Against Mac1 Domain of SARS-CoV-2 Nsp3

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Abstract—SARS-CoV-2 encodes the Mac1 domain within the large nonstructural protein 3 (Nsp3), which has an ADP-riboseylhydrolase activity conserved in other coronaviruses. The enzymatic activity of Mac1 makes it an essential virulence factor for the pathogenicity of coronavirus (CoV). They have a regulatory role in countering host-mediated antiviral ADP-ribosylation, which is unique part of host response towards viral infections. Mac1 shows highly conserved residues in the binding pocket for the mono and poly ADP-ribose. Therefore, SARS-CoV-2 Mac1 enzyme is considered as an ideal drug target and inhibitors developed against them can possess a broad antiviral activity against CoV. ADP-ribose-1-phosphate bound closed form of Mac1 domain is considered for screening with large database of ZINC. XP docking and QPLD provides strong potential lead compounds, that perfectly fits inside the binding pocket. Quantum mechanical studies expose that, substrate and leads have similar electron donor ability in the head regions, that allocates tight binding inside the substrate-binding pocket. Molecular dynamics study confirms the substrate and new lead molecules presence of electron donor and acceptor makes the interactions tight inside the binding pocket. Overall binding phenomenon shows both substrate and lead molecules are well-adopt to bind with similar binding mode inside the closed form of Mac1.

Index Terms—COVID-19, Mac1, Macro X, Molecular Dynamics, Nsp3, Quantum Mechanics, SARS-CoV-2

1 INTRODUCTION

The Coronavirus Disease 2019 (COVID-19) pandemic outbreak, an infectious disease caused by a novel Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2), is highly catastrophic and has a high impact on the global economy [1]. This coronavirus swiftly spread through communities and reached all continents except Antarctica in an incredibly limited time [2]. The current confirmed case numbers are more than five million with a mortality rate of 2.6 percent (according to https://ourworldindata.org/mortality-risk-covid). SARS-CoV-2 is an enveloped virus from family Coronaviridae and genus beta-coronavirus, comprising a large positive-strand single-strand RNA (+ssRNA) genome (~30 kb), which encodes four structural proteins (spike, envelope, membrane, and nucleocapsid protein) that are components of the virus particle, several nonstructural (Nsp) proteins mostly with enzymatic activities and accessory proteins [3]. The ORF1a and b encoded two large polypeptides which were later cleaved by viral proteases into 16 functional proteins (Nsp1-16), where Nsp11 alone is a short peptide. Nsp3 is the largest multidomain protein (~200kDa) in coronaviruses and is notable because of the presence of a key enzyme, papain-like cysteine protease (PLpro), which is essential for viral replication and a target protein for drug discovery [4], [5]. The Nsp3 is found to be significantly different in two SARS-CoVs in comparison with other Nsp3 [6], [7]. Computational studies have shown that coronaviral Nsp3 may comprise 10-16 domains [8]. SARS-CoV-2 Nsp3 also includes multiple domains, an N-terminal ubiquitin-like (Ubl) domain followed by a highly variable and conserved macrodomain that binds to ADP-ribose [9], hereafter called as Mac1, previously called as Macro X domain because of its unknown structure and function, and multiple downstream SARS unique macrodomains, namely Mac2 and Mac3 (previously called as SUD-N and SUD-M) that binds to G-quadruplex, and a domain preceding Ubl2 and PLpro (DPUP) domain (previously called as SUD-C) that binds to ssRNA [10], [11]. The macrodomains are followed by the Ubl2 domain, PLpro, nucleic acid binding domain (NAB) and several other domains in the C-terminus which also includes transmembrane regions [12], [13], [14]. The macrodomains (~130-190 AAs) can be either stand-alone or a part of a multi-domain protein [15]. These domains are also widely present in viruses, bacteria, archaea, and several eukaryotes. They recognize ADP-ribose or its derivative or post-translational modification covalently bound to proteins, thereby playing a varied role in cellular processes (e.g., DNA repair, signal transduction, immune response) in all kingdoms of life [16]. The yeast macrodomain (Foa1) is found to bind ADP-ribose-1′-phosphate, which is a by-product of tRNA splicing [17], [18], [19]. In viruses, they play a role in virulence and suppressing the host innate immune response and...
have shown potential roles in viral replication and pathogenesis [20], [21]. The macrodomains are found in three viral families, such as Coronaviridae, Togaviridae, and Hepeviridae, which are evolved to counteract the host defense mechanism [11], [22]. The coronaviral Mac1 domain specifically binds to ADP-ribose-1'-phosphate and has dephosphorylation or ADP-ribosylhydrolase activity, demonstrating their possible function in over-turning host antiviral ADP-ribosylation [23], [24]. Recent studies have shown that the SARS-CoV-2 Nsp3 Mac1 binds to ADP-ribose with a higher binding affinity compared to SARS-CoV in vitro and catalyzes the hydrolysis of ADP-ribose-1'-phosphate. Even though the Mac1 domain is found to be dispensable for viral RNA replication, several studies are available showing its regulatory role in counteracting host innate immunity [9], [11], [25]. Therefore, because of the emerging novel roles of macrodomains, they are considered as a potential therapeutic target for drug discovery in the viral world.

2 MATERIALS AND METHODS

2.1 Sequence and Structure: Comparative Analysis

Pairwise protein structural comparison was performed between the Mac1 domain of SARS-CoV-2 (PDB ID: 6WEY) and SARS-CoV (PDB ID: 2FAV) using TM-align server [26], [27]. The structure-based sequence alignment of SARS-CoVs Mac1 domain sequences was prepared using ENDscript server 2.0 [28]. Secondary structural elements are displayed above the sequence and the residues are highlighted based on the residue conservation (red for identical and yellow for similar), with the lower bar indicating the solvent accessibility and hydrophathy scale. All structure and interaction figures were rendered using PyMOL v2.3.5 [29].

2.2 Protein Preparation

The multi-domain Nsp3 is an essential component of the replication/transcription complex and from this the Mac1 domain structure is retrieved from the protein data bank (PDB ID: 6WEY). From the available Mac1 domains of SARS-CoV-2, the 6WEY has the ultra-high resolution of 0.95 Å and thus the apo structure is chosen for the study (Fig. 1a) along with the ADP-ribose from the PDB ID: 6YWL. The substrate co-ordinate, namely adenosine-5'-diphosphoribose is taken from the X-ray structure of SARS-CoV-2 Mac1 domain in complex with ADP-ribose (Fig. 1c, PDB ID: 6YWL) and merged with high resolution structure using the Maestro (Schrödinger). After the complex formation, the correctness of the structure is obtained using the protein preparation wizard [30]. Here, the partial charges of the atoms that are filled along with bond orders, missing atoms and side chains are refined. Sidechain amino acids

Fig. 1. Structure and sequence alignment of Mac1 domain from SARS-CoVs (a) Structural alignment of the Mac1 domain from experimental structures of SARS-CoV-2 (colored as the rainbow) with SARS-CoV (colored as gray) (b) Structure-based sequence alignment of Mac1 domain from SARS-CoVs highlighting identical residues in red and similar residues in yellow (c) Surface representation of closed ADP-ribose bound Mac1 domain structure from SARS-CoV-2 (PDB ID: 6YWL) and zoomed-in view of the binding site displaying crucial residues for ADP binding and catalytic activity.
Ast, Gln, and His angles are flipped to influence the H-bond formation and generate tautomers/ions states [31]. The protein-substrate structure is applied with OPLS-3e force field (FF) for optimizing the intramolecular hydrogen bonds using the optimization method and minimized till the root-mean-square deviation (RMSD) levels reaches the 0.30 Å [32].

### 2.3 ZINC Database Preparation for Screening

The small molecule library of 230 million compounds is downloaded from ZINC database (https://zinc.docking.org/) [33]. The whole library is prepared by maintaining the pH values around 7.0 ± 2.0 for maintaining the ligand ionization state and ligands stereoisomer conformations are checked by LigPrep (Schrodinger) [34]. By applying the OPLS3e FF with all combinations applied, up to three conformations per ligand are generated.

### 2.4 Glide - Grid Generation

The prepared complex of Mac1-ADP-ribose complex is taken for grid generation using the Schrodinger’s Glide–Grid generation method [35]. The ADP-ribose molecule is manually picked, and the grid box is customized around it. For softening the receptor non-polar regions, the vDW scaling factor is set to 1.0 kcal/mol and partial charge for atoms cut-off is set to default 0.25. The region around 2 Å from the ADP-ribose is set on focus for the molecular docking approach and the amino acids playing interactions with ADP-ribose are fixed as H-bond constrains [36]. Finally, the positional grid is generated around the ADP-ribose, where the new ligands will be replaced and from the best compounds will be reported using the molecular docking method.

### 2.5 Virtual Screening

The prepared ligand from the LigPrep and grid file from the receptor grid generation is input for the Schrodinger’s Virtual Screening Workflow (VSW) [37]. Initial filtering is done by applying the Qikprop filter to pass the compounds having strong ADME profiles. Lipinski filter is avoided, as the substrate binding pocket is moderately bigger in size and able to hold large size molecules. Reactive functional group molecules are filtered out and from these filters, unnecessary compounds are filtered out and subjected to molecular docking approach. Initial level screening is performed using High-Throughput Virtual Screening (HTVS), to check for the apt compound that suits with ADP-ribose binding pocket. The top 10 percent of the successive compounds from HTVS is subject to pass to the standard precision (SP) docking method, and from that top 5 percent compounds will be examined for its interactions with Extra Precision (XP) docking method [38]. Pose validations will be done for the final compounds which are the best compounds from the glide docking score and Prime MM/GBSA ranking [39].

### 2.6 Quantum Polarized Ligand Docking

For obtaining the accurate interactions of the successive lead compounds, the redocking methodology using the Schrödinger’s Quantum Polarized Ligand Docking (QPLD) is employed [40]. In this, the QSite and Jaguar enabled with Quantum Mechanics (QM) will be applied for the ligand and binding site residues. For the other regions of the proteins, the Molecular Mechanics (MM) will be applied using the Impact. QM region is assigned through DFT with the B3LYP using 6-31G** basis set, and “Ultraline” SCF accuracy level (iacc = 1, iacscf = 2) [41]. Through this, the pseudo charges of each protein-ligand atom will be refined and get accurate charges. QPLD methodology initially intakes normal Glide docking with XP; those poses will be subjected to charge refinement using the QM and MM methods and refined charges are employed for redocking method using the XP/QPLD [42], [43].

### 2.7 Density-Functional Theory (DFT) Calculations

The electronic charge transfer effects of drug like molecules that play insights in pharmacological effects are calculated using the DFT analysis [44]. For DFT analysis, the ADP-ribose substrate and the top five compounds are imported with jaguar workflow panel. Functional set is assigned to B3LYP with 6-31G**+** basis set. For calculating the multipole moments, the Coupled Perturbed Kohn-Sham (CPKS) equations are applied, and for elements solution, the Poisson-Boltzmann Finite (PBF) is applied. Among the QM properties, calculating the molecular orbitals (HOMO/LUMO/Band gap), electron density and molecular electrostatic potential (MESP) are marked for output [45], [46].

### 2.8 Molecular Dynamics Simulations

Molecular Dynamics (MD) simulations are performed for the ADP-ribose substrate bound with the Mac1 domain, and also for the top five lead compounds bound with the Mac1 domain using the GROMACS 5.1.4 (GROningen MAchine for Chemical Simulations, http://gromacs.org) [47], [48]. All the six protein-ligand complexes are simulated for the timescale of 50 ns for understanding the stability of ligand inside the protein binding pocket. For that, the complex files are solvated with TIP3P water model, periodic boundary set at 1.0 nm and systems prepared using OPLS-AA force field. Ligand charge adoption for the system is performed using the external PRODRG server (http://prodrg1.dyndns.org/) and the whole system is neutralized by adding suitable (Na+/Cl-) ions in required concentrations [49], [50]. Energy minimization step is performed for 1000 steps by applying steepest descent algorithm with tolerance of 10 kJ/mol/nm to avoid the steric clashes. Reference pressure is set to 1.0 bar using Parrinello-Rahman along with periodic boundary conditions with cut-offs for Lennard-Jones and reference temperature is set to 300 K using Berendsen thermostat. Successful minimized systems are subjected to NVT and NPT ensembles for initiating the MD simulation step for 50 ns of timescale [51].

### 3 Results and Discussion

#### 3.1 Sequence and Structural Analysis of Mac1

The sequence comparison is done between the SARS-CoV-2 and SARS-CoV Mac1 domain. The Mac1 domain of SARS-CoV-2 has the sequence length of 172 amino acids, and the SARS-CoV has the sequence length of 182 amino acids. Both the SARS-CoV-2 and SARS-CoV share the sequence similarity of 81.7 percent (143/172) and identity of 70.9 percent (124/172) as shown in Fig. 1b. Even though the difference
between these sequences hold ~18 percent, the structural alignment between these two proteins shows similar structural and functional phenomenon with the RMSD of 1.2Å. The three-dimensional structure of Mac1 comprises a central six-stranded mixed beta sheet surrounded by helices that adopt a highly conserved globular three-layered α/β/α sandwich fold with minor variations. The overall fold is quite similar to the human non-histone domain of macroH2A-like fold, classifying them as a member of MacroD-like family [16]. ADP-ribose binding cleft seems to undergo conformational changes upon binding or transition between apo and complex state. In SARS-CoV, key residues in loop 1 and 2 found between β2-α2 and β5-α5 respectively, contribute to ADP-ribose binding. Mutational studies on Asp23, Asn38, Asn41, His46, Gly131, Gly47+Gly48 and Phe133 from SARS-CoV have shown that they disrupt their catalytic function or virulence in vivo [11], [18], [52]. Protein topology network shows that SARS-CoV-2 Mac1 is arranged by 2 sheets, 1 beta alpha beta unit, 2 beta hairpins, 1 psi loop, 2 beta bulges, 7 strands, 9 helices, 6 helix-helix interacs, 13 beta turns and 2 gamma turns, while the SARS-CoV is arranged as 2 sheets, 1 beta alpha beta unit, 2 beta hairpins, 1 psi loop, 2 beta bulges, 7 strands, 9 helices, 7 helix-helix interacs, 11 beta turns and 2 gamma turns.

3.2 Molecular Interactions of ADP-Ribose With Mac1

The structure alignment provided in Fig. 1a shows the both proteins from the SARS-CoV-2 and SARS-CoV are structurally identical. Mac1 domain shows two distinct features in apo and holo forms by an open and closed binding pocket mechanism, respectively. Before the ADP-ribose binding, the protein substrate binding site is showing the open pocket, and while the ADP-ribose bound within the pocket, the loops come closer to each other and form the closed binding pocket. In SARS-CoV-2, the ADP-ribose bound to the substrate binding site (closed form) as shown in Fig. 1c, and the amino acids Asp226, Ile227, Asn244, Val253, Ser332 and Phe336 plays the role of holding the ADP-ribose through hydrogen bonding interactions. Phe360 also contributes to the binding of ADP-ribose through II-II interactions as shown in Figure S1, which can be found on the Computer Society Digital Library at http://doi.ieeecomputersociety.org/10.1109/TCBB.2020.3037136. In this the Asp226 and Asn244 plays the vital role in the catalytic reaction mechanism, and these two residues are highlighted in Fig. 1c.

3.3 Virtual Screening of Mac1 Domain

The binding form of ADP-ribose with Mac1 domain shows that the substrate binding pocket is holding the large regions of positive electron density with the core amino acid Asp226, Ile227, Asn244, Val253, Ser332, Phe336 and Phe360. Among these, Asp226 playing the role in holding ADP is negatively charged, where the other key residue Asn244 is a polar and non-charged amino acid (Fig. 1). Apart from these two amino acids, Ile227 and Val253 are non-polar categorized as aliphatic amino acid due to the availability of aliphatic side chain functional group. The other amino acid Ser332 is polar with non-charged amino acid, while Phe336 and Phe360 are the only aromatic amino acid in this pocket. Integrating the ligand and structure-based approach can determine a strong and potent compound that suitably bound towards these large regions of positive electron density pocket. Pointing the ADP-ribose focused ligands that retrieve few compounds through HTVS, SP and XP docking. Those compounds with docking score of < -10 kcal/mol and the binding energy of < -50 kcal/mol provides few compounds, that exactly matched the substrate binding pocket. For enhancing the accuracy of the docked complex, the redocking methodology using the Quantum Polarized Ligand Docking (QPLD) is utilized. This is to attain the select of correct and accurate binding poses by implementation of QM/MM docking. QPLD derived partial charges are playing the vital role in terms of enhancing the accuracy and provides the possible binding pose that is close to the experimental structures. From this screening, the compounds namely ZINC08765069, ZINC08792474, ZINC0887336, ZINC08879971 and ZINC00897592 arise as strong lead candidates, that actively bind with ADP-ribose binding pocket of Mac1 domain (Fig. 2), and all these compounds have the docking score of < -10 kcal/mol and binding energy of < -50 kcal/mol as shown in Table 1.

Here, in this table, the scoring values are shown to hold the difference between the XP and QPLD docking. This is due to changes in atomic charges in the protein-ligand complex of XP docking is considered as pseudo charges, and the hybrid QM/MM incorporated in QPLD to replace the accurate charges that enhance the scoring and interaction pattern of the QPLD poses. These accurate charges derived protein ligand interactions are necessary, as the charge of each atom is replaced and thus, the interaction phenomenon of positive charge tend to negative charged atoms, while that of negative charge tend towards the positively charged atoms. The role of partial charges between the positive and negative charges tends to alter the hydrogen bonding in XP and QPLD and provides the slight variations in the binding pose. The RMSD variations of XP docked ligand and QPLD docked ligand are provided in Table 1, which shows the variations that occurred due to the replacement of atomic charges. From this, the role of QPLD in this screening has obtained higher benefit by the charge modulations, and explains that the XP docking provides pose conformations in binding pocket, and adding up the QPLD redocking method provides the change in angle conformations that bound more accurately inside the binding pocket.

3.4 HOMO/LUMO and MESP Calculations

The DFT calculations are performed to understand the key level atomic properties of the molecular structure and here, the QM techniques are applied to understand the charge transfer ability of atoms for the substrate and new lead compounds through HOMO/LUMO calculations. The exact conformations obtained from the QPLD docking pose are subject to optimize with B3LYP/6-31G**++ level of theory, and by this the angles and bond lengths of the substrate and ligand optimized geometries are accommodated in realistic position. For understanding the solvation effect, geometry optimization, the base sets are performed with the 6-31G**++ and PBF solvation. In the substrate and lead molecules, the electron donors are accessed by HOMO, the electron acceptors are accessed by LUMO, and the gap between the HOMO and LUMO is
called as band gap energy that elucidates the electron transfer swift between both energy levels. The representation of HOMO/LUMO and MESP is provided in Fig. 3 for the substrate and ligand by showing its quantum mechanical features of HOMO/LUMO and MESP for the ADP-ribose (a) and top lead compounds (b-f). When analyzing the structural features of substrate with new lead molecules, except the molecule ZINC00897592 all the other compounds show similar posture and while dissecting the electron donor region of the substrate and new leads, all the ligands show electron donor tendency in the head region, which is shown in Fig. 3 (HOMO). In terms of electron acceptor tendency except the compound ZINC08879336, all the other compounds show the LUMO regions in central part, and the ZINC08879336 shows the LUMO in the tail part. In terms of MESP, as shown in Fig. 3, red color surface indicates the most negative potential region, and the blue surface indicates the most positive potential region. This shows that, the substrate MESP surface is matched in the head and tail portions for all the new compounds except the ZINC00897592, and also the difference in central part is due to the presence of

| Compounds       | Docking Score (XP) kcal/mol | Docking Score (QPLD) kcal/mol | MM/GBSA energy (XP) kcal/mol | MM/GBSA Energy (QPLD) kcal/mol | RMSD (Å) |
|-----------------|-----------------------------|-------------------------------|-------------------------------|-------------------------------|----------|
| ADP-ribose      | −7.21                       | −7.49                         | −42.38                        | −51.31                        | 0.058    |
| ZINC08765069    | −13.38                      | −12.86                        | −71.62                        | −70.33                        | 0.106    |
| ZINC08792474    | −12.89                      | −12.20                        | −86.35                        | −85.55                        | 0.092    |
| ZINC08879336    | −12.27                      | −12.83                        | −77.90                        | −81.37                        | 0.068    |
| ZINC08879971    | −11.82                      | −11.88                        | −68.52                        | −70.30                        | 0.061    |
| ZINC00897592    | −11.17                      | −11.55                        | −55.23                        | −58.69                        | 0.023    |

Fig. 2. Molecular docking interactions of ADP-ribose (a) and top lead compounds (ZINC08765069 (b), ZINC08792474 (c), ZINC08879336 (d), ZINC08879971 (e), ZINC00897592 (f)) with the Mac1 domain of SARS-CoV-2 using the Quantum Polarized Ligand Docking.

TABLE 1

Scoring Values of Substrate Binding and New Ligand Binding With Mac1 Domain of SARS-CoV-2
phosphate atom in the substrates and those are more are less replaced by the nitrogen atoms in the ligands except ZINC00897592.

3.5 Molecular Dynamics Simulations

The protein-ligand complex for the substrate and ligand bound complex is analyzed for the molecular dynamics simulations for the timescale of 50 ns. It is mandatory to compare the protein-ligand bonds for the substrate and new lead molecules, and their exhibiting kinetic energy for their binding in the dynamic environment. Among the interactions, hydrogen bonding is core important to observe in dynamic environment, especially the amino acid contribution to hold the ligand in the aqueous environment, and so the average hydrogen bonds between the protein and ligands are shown in Fig. 4a. The average hydrogen bonds for the ADP-ribose show 2.1, while the new lead compounds show 2 and from that the compound ZINC00897592 shows the higher average number of hydrogen bonds by 2.9. This is due to the compound ZINC00897592 allocation in binding pocket, which is comparatively smaller and able to hold with aromatic amino acids. The stableness of protein-ligand simulations is calculated using the RMSD, and RMSF fluctuations, provided in the supplementary information. The RMSD (Figure S2 available online) and RMSF (Figure S3 available online) favor the protein-ligand stability in the MD simulations on the dynamics aqueous state environment. In terms of kinetic energy, almost all the substrate and new lead molecules have the tendency to poseur 80,000 kJ/mol and from this, only the compound ZINC08765069 can surpass the kinetic energy with the score of 120,000 kJ/mol. Through this hydrogen bond analysis and kinetic energy analysis, we assume that the new compounds are having similar binding with ADP-ribose binding mechanism, and in dynamic environment, it can actively participate in tight binding with the Mac1 domain of SARS-CoV-2. As discussed earlier, the binding pocket has the unique feature by showing the open binding pocket in the apo form, and closed binding pocket in the holo form. The new lead compounds show similar HOMO/LUMO and MESP surfaces along with the substrate molecules shown in the DFT calculations. By that, the new leads are checked for its common binding mode with the substrate binding. The comparison is shown in Fig. 5, indicating that the new lead compounds have the tendency to bound in the similar binding pose with the substrate (green). Also, the new lead molecules binding did not alter the closed gate of the binding pocket, and this may be due to the new compounds having nitrogen atoms in the middle part that does not allow the binding pocket to open and remains stable in the MD simulations. We also assume that; these substrate binding can show open and closed binding pocket in the long-range timescale in the MD simulations.

Fig. 3. DFT calculations for the substrate and ligand by showing its quantum mechanical features of HOMO/LUMO and MESP for the ADP-ribose (a) and top lead compounds (ZINC08765069 (b), ZINC08792474 (c), ZINC08879336 (d), ZINC08879971 (e), ZINC00897592 (f)).
In summary of this work, we come up with deep understanding of ADP-ribose substrate binding with Mac1 domain and based on the structural insights, we found few strong potential lead molecules that can regulate the SARS-CoV-2 pathogenesis. Through this work, we report five new lead molecules from the ZINC database namely ZINC08765069, ZINC08792474, ZINC08879336, ZINC08879971 and ZINC00 897592. These compounds are the outcome of strong filtering criteria of docking score of < -10 kcal/mol and binding energy of < -50 kcal/mol. The interacting amino acids and the ligand orientations are making accurate by applying the QM charges towards the ligand binding environment. The QM-based DFT calculations strongly says that the electron donor atoms are functioning in head part of the substrate and new lead molecules, while the electron acceptor is located in the middle part of the substrate and new lead molecules, except ZINC08879971 molecule. This unique feature of the presence of electron donor and acceptor in the head and middle part of the ligands is making these ligands to adopt well inside the binding pocket with the similar binding mode. Interestingly, in substrate-binding and lead molecule binding does not affect the closed binding pocket of the Mac1. This substrate and lead molecule binding is validated using the molecular dynamics in aqueous environment, showing that the substrate and lead molecules are stable by their tendency of forming two or more average hydrogen bonds between the protein-ligand complex involving crucial residues Asp226 and Asn244 which are essential for ADP-ribose binding and catalytic activity in SARS-CoV Mac1 domain. This complex structure also verified with kinetic energy shows all the complex shows lower 80000 kJ/mol that indicates, strong and stable binding of substrate and new lead molecules. We also report that, the closed pocket of Mac1, may have open and closed gated mechanism in their enzyme reaction active states. Through this work, we recommend that, these five lead compounds on further experimental validations along with strong international effort can make these compounds as strong inhibitors against the SARS-CoV-2 Mac1 domain ADP-ribosylhydrolase activity.

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