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Computational studies reveal Fluorine based quinolines to be potent inhibitors for proteins involved in SARS-CoV-2 assembly

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ABSTRACT

World is witnessing one of the worst pandemics of this century caused by SARS-CoV-2 virus which has affected millions of individuals. Despite rapid efforts to develop vaccines and drugs for COVID-19, the disease is still not under control. Chloroquine (CQ) and Hydroxychloroquine (HCQ) are two very promising inhibitors which have shown positive effect in combating the disease in preliminary experimental studies, but their use was reduced due to severe side-effects. Here, we performed a theoretical investigation of the same by studying the binding of the molecules with SARS-COV-2 Spike protein, the complex formed by Spike and ACE2 human receptor and a human serine protease TMPRSS2 which aids in cleavage of the Spike protein to initiate the viral activation in the body. Both the molecules had shown very good docking energies in the range of -6kcal/mol. Subsequently, we did a high throughput screening for other potential quinoline candidates which could be used as inhibitors. From the large pool of ligand candidates, we shortlisted the top three ligands (binding energy -8kcal/mol). We tested the stability of the docked complexes by running Molecular Dynamics (MD) simulations where we observed the stability of the quinoline analogues with the Spike-ACE2 and TMPRSS2 nevertheless the quinolines were not stable with the Spike protein alone. Thus, although the inhibitors bond very well with the protein molecules their intrinsic binding affinity depends on the protein dynamics. Moreover, the quinolines were stable when bound to electronegative pockets of Spike-ACE2 or TMPRSS2 but not with Viral Spike protein. We also observed that a Fluoride based compound: 3-[3-(Trifluoromethyl)phenyl]quinoline helps the inhibitor to bind with both Spike-ACE2 and TMPRSS2 with equal probability. The molecular details presented in this study would be very useful for developing quinoline based drugs for COVID-19 treatment.

1. Introduction

In the year 2019, we saw the emergence of a highly contagious disease known as COVID-19. The disease was caused by the novel betacoronavirus known as SARS-CoV-2 [1]. The virus is believed to have originated primarily from animal sources which then subsequently spread across the world through human-human contact [2]. This novel form of coronavirus is closely related to SARS-CoV; however, it is much more infectious and mutation prone compared to its predecessors [3]. It has claimed millions of lives across the globe and severely affected several others.

The SARS-CoV-2 is a single-stranded RNA-enveloped virus, encoding 9860 amino acids [4]. The genes express structural and nonstructural proteins. They encode structural proteins such as Spike (S), Envelope (E), Membrane (M) and Nucleocapsid (N) and nonstructural proteins, such as 3-chymotrypsin-like protease, papain-like protease, and RNA-dependent RNA polymerase [5]. SARS-CoV-2 uses the human Angiotensin-converting enzyme 2 (ACE2) receptor for cell entry, in synergy with the host’s Transmembrane protease, serine 2 (TMPRSS2) [6].

The Spike is a transmembrane protein with an intracellular domain and an extracellular domain. The trimeric Spike protein can be further divided into S1 and S2 subunits. While, S1 subunit contains the receptor-binding domain (RBD) which interacts with the receptor, the S2 subunit facilitates virus-cell fusion and entry [7,8]. SARS-CoV-2 and SARS-CoV, both require an Angiotensin-converting enzyme 2 (ACE2) receptor for entering the host cells [9]. Transmembrane serine protease 2 (TMPRSS2) is a surface protein expressed in humans. Endothelial cells...
primarily express it throughout the respiratory and digestive tracts. It is believed to be involved in certain pathologies, however, the exact biological function of TMPRSS2 is unknown [6,10,11]. The viral Spike glycoprotein is proteolytically cleaved by TMPRSS2, induces virus-cell membrane fusion at the cell surface and represents one of the essential host factors for initiating SARS-CoV-2 pathogenicity [12,13]. This process is analogous to cell entry and viral activation seen in other coronaviruses, including SARS-CoV, MERS-CoV as well as influenza viruses such as influenza H1N1. The lack of TMPRSS2 in the airways decreases the severity of lung infection by SARS-CoV and MERS-CoV [14,15]. The furin cleavage site is found in SARS-CoV-2 unlike SARS-CoV-1 or any other known sarbecovirus. As mutations at this site decrease viral infectivity it has been established that by acquiring the furin cleavage site the virus has benefitted itself. However, in other mutations in the site of furin cleavage, the infectivity was increased to a great extent [16-20].

Several pharmaceutical companies and research organizations are actively working towards standardizing a treatment strategy for COVID-19. In the later part of 2020, many vaccines have been proposed for COVID-19. However, most of the drugs administered for treatment are based on drug repurposing and those that are used for treatment of flu like diseases [21-23]. HCQ and CQ (CQ) are two well-known drugs that have been used since the early 1900s for the treatment of malaria, rheumatoid arthritis and systemic lupus erythematosus [24,25]. During the 2002 SARS epidemic, various research groups worked on using CQ and HCQ for treatment. Both HCQ and CQ were found to successfully prevent viral replication inside cells [26,27]. It is believed that CQ might prevent the binding of SARS Spike protein and its ACE2 receptor by inhibiting glycosylation [28]. Similarly, HCQ also prevents SARS binding to ACE2 receptors by restricting its interaction with gangliosides [29,30]. Physiologically, HCQ and CQ were found to have increased the pH of the intracellular compartments and degraded the cellular components [26]. Very recently, it has been observed that HCQ was more potent in inhibiting SARS-CoV-2 than SARS-COV and even more effective than CQ [31]. However, both are equally effective in inhibiting virus replication [32]. Several clinical trials, although with small sample size, had found the inhibitory potency of the drug [33]. The preliminary evidence of the drug’s effectiveness had even resulted in the worldwide shortage of HCQ and CQ based drugs as a means to prevent COVID-19 [34-36]. Unfortunately, due to lack of sufficient trials and significant and unexpected side-effects, the study has not progressed much [37-41].

Here, we implement an in-silico approach combining high throughput docking and computer simulations to propose new HCQ and CQ based analogues. The interactions were mainly studied with Spike, Spike-ACE2 and TMPRSS2 which prove as proteins vital for the establishment of viral contact and entry into the host cell [12,42-46]. We began by docking the CQ and HCQ to the proteins to locate the favorable binding sites [47,48]. Subsequently, we virtually screened a large number of ligands with the quinoline backbone and performed high throughput docking. Based on the docking results, molecular Dynamics (MD) simulations were performed on the best docked complexes. However, the results show a difference in the steadiness of the inhibitors with the proteins. It was found that the Spike-ACE2 complex most favorably binds to quinoline based inhibitors followed by TMPRSS2.

2. Materials and methods

The crystal structures of the proteins were obtained from the Protein Data Bank (PDB IDs: 6W41, 6M17, 2OQ5). The associated protein human antibody CR3302 in 6W41 and B0AT1 complex from 6M17 were manually removed before docking. The spike-ace2 receptor used is in complex with the Receptor Binding Domain (RBD), which comprises binding hotspots of the spike protein to ACE2. Again, since the RBD can fold independently with respect to the rest of the spike protein we considered the complex with the RBD rather than the whole spike protein as reported in earlier papers [49-51].

2.1 Molecular docking

Molecular docking studies of HCQ and CQ respectively with the three aforementioned proteins were performed. A library consisting of about 10,000 quinoline analogues was created using the PubChem chemical database. The molecules downloaded were filtered using a Lipinski Rule of Five filter, where approximately 7000 molecules were shortlisted. The protein structures were opened in the BIOVIA Discovery Studio

Fig. 1. (a) Structure of Receptor binding domain of Spike protein (shown in red), (b) Spike-ACE2 complex where spike protein is shown in red, ACE2 protein in cyan (c) TMPRSS2 protein.
Table 1
Docking energies of CQ and HCQ with (a) Spike, (b) Spike-ACE2 and (c) TMPRSS2.

|            | HCQ     | CQ     |
|------------|---------|--------|
| TMPRSS     | -6.1    | -6.4   |
| Spike-ACE  | -6.1    | -5.8   |
| Spike      | -6.4    | -5.9   |

visualizer to remove heteroatoms and co-crystallized ligands. The protein structure and the ligands were loaded in PyRx and the ligands were minimized using Universal Force Field to their lowest energy configuration. The binding of these ligands was performed on the active sites of the protein. The interactions of top 10 ligands obtained by High Throughput Screening were studied in the BIOVIA Discovery Studio visualizer.

2.3. MD Simulations

Molecular Dynamics (MD) Simulations were initiated with the top 3 minimum energy configurations by using GROMACS (version 2020.4) MD simulation software package [56]. Charmm36-jul2020 force field was used to get the parameters for protein [57] which was immersed in a water box containing TIP3P water model [58]. The topology and parameters for ligand molecules were generated using the CHARMM General Force Field Server [59]. The docked complexes were placed inside a cubical box with TIP3P water molecules and neutralizing ions. The box dimensions were adjusted in such a way that the box edge was at least 1 nm away from the protein-ligand complex. After initial minimization using initially steepest descent and later conjugate gradient algorithms, the system was well-equilibrated in an isochoric-isothermal ensemble at 300 K. This was followed by equilibration at an isothermal-isobaric ensemble. Positional restraints were added to the ligands during the minimization and equilibration. The time step was kept 2 fs. Electrostatic and Van der Waals interaction cut off were kept at 1.2 nm. The long-range electrostatic interactions were treated by using Particle-Mesh Ewald sum and SHAKE was used to constrain all bonds involving hydrogen atoms. After equilibration, the structures were subjected to MD simulation to generate data for analyses.

3. Results and discussion

The crystal structures of Spike protein’s receptor binding domain, Spike-ACE2 complex and TMPRSS2 (Fig. 1) were obtained from the protein data bank (PDB IDs: 6W41, 6M17, 2Q5Q) [60]. Heteroatoms, such as co-crystallized ligands and water molecules were removed. The associated protein human antibody CR3302 in 6W41 and B0AT1 complex from 6M17 were manually removed before docking. The potential binding sites of the inhibitors were chosen from the previous studies. We began by docking HCQ and CQ to the proteins to ascertain their binding affinities and ligand binding sites. Secondly, we virtually screened 7000 quinoline based small molecule inhibitors to find out the best binding ligands. Finally, we performed Molecular Dynamics (MD) simulations to get more insights on protein-ligand complexes.

3.1. Repurposing HCQ and CQ shows promising results

HCQ (HCQ) and CQ (CQ) have shown potential to inhibit the replication of SARS and SARS-CoV-2 viruses in vitro [31,32,61]. To begin with we wanted to study the binding characteristics of HCQ and CQ with the viral Spike protein, the viral Spike and human ACE2 receptor complex and the human protease TMPRSS2 to gain a better understanding of the nature of binding of the drug and proteins.

The large Spike protein has more than thousand residues and is also a transmembrane protein. We selected the receptor binding domain of the Spike protein and docked HCQ and CQ. The active sites for spike protein were obtained from literature review [62,63]. HCQ and CQ, despite having similar chemical structures, it was observed that there were significant differences in their binding energies. The binding energies were -6.4 kcal/mol with HCQ compared to CQ with -5.9 kcal/mol (Table 1). The 0.5 kcal/mol difference in energy can be attributed due to additional interactions of HCQ with the protein including two hydrogen bonding interactions of the ligand with SER469 and ILE472 in the receptor binding domain of Spike protein (Fig. 2). Similarly, we docked the Spike-ACE2 receptor complex with HCQ and CQ. The active sites for docking were taken from literature [64,65]. The complex comprises the RBD of Spike protein and the interacting region of the ACE2 receptor. Spike-ACE2 protein complex was selected because it plays a vital role in the entry of virus, the glycoprotein attaches to the ACE receptor which fuses the virus into the cell membrane [65].

We wanted to check if HCQ and CQ bind to the complex with similar affinities. Upon docking it was observed that the inhibitors docked at the interface of the proteins. Here also the HCQ binding energy (-6.1 kcal/mol) was higher than CQ (-5.8 kcal/mol) (Table 1). Although comparative binding could be observed for both the ligands, presence of unfavorable interactions reduced the overall binding energy (Fig. 2).
TMPRSS2 is a very significant protein due to its undeniable role in the pathophysiology of SARS-CoV-2 [11,15]. The protease aids the virus in host cell entry and replication. Active sites of the protein were deduced from the earlier literature [44,66]. Surprisingly, the binding energies of HCQ and CQ to TMPRSS2 were found to be very similar (-6.1 kcal/mol and 6 kcal/mol respectively) (Table 1). Nevertheless, we observed more numbers of residues interacting with the ligand in the presence of the polar groups in HCQ (Fig. 2). The binding pockets of all the proteins are shown in Fig. 3 for clarity.

From the above calculations, it is inferred that the HCQ and CQ both could bind well with all the three proteins which have been considered for our study. Furthermore, we could garner a threshold value of the binding energy and the probable binding sites in the proteins. Coadministration of HCQ with antibiotics lead to side effects like QT prolongation, hence administration of HCQ has not been carried out in clinical studies for COVID-19 [67,68]. Thus, we did an extensive search on the alternate quinoline analogues which could prove as alternate candidates for drug designing.
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binding affinity of these ligands in the docked pockets over a period of time under physiological conditions. In order to understand this, we performed MD simulations of the proteins with the docked ligands and studied the changes in interaction over a period of time.

3.3. Protein-ligand binding dynamics

The first three quinoline based inhibitors which showed the highest docking score were subjected to Molecular Dynamics (MD) simulations in order to check the complex stability in nanoseconds time scale and to observe any change in the protein dynamics. After minimization and equilibration, all the protein-ligand complexes were subjected to MD simulations in an NPT ensemble. The ligand parameters were obtained from CGENFF web server. Gromacs 2020.4 and CHARMM36 force field was used for all the simulations (details in materials and methods).

To verify if the systems are stable in an aqueous environment with neutralizing ions, the Root Mean Square Deviation (RMSD) from the initial docked conformation was calculated (Fig. 5). From the RMSDs, it can be seen that Spike ligand complexes had comparable values indicating that all the systems had similar conformation. However, the Spike-ACE2 ligand systems show dramatic RMSD plots. The protein when bound to the third ligand shows the least RMSD value. A major conformational change was observed to have taken place in the Spike-ACE2 ligand 1 complex, Spike-ACE2-ligand 2 also had a higher RMSD value. This indicates that the complex is more stable when bound to the third ligand. RMSF of ACE2 shows similarity of fluctuations in the three complexes (Figure S1). The Spike-ACE2 interactions in wild-type and inhibitor bound complexes. were preserved in all the three complexes (Table S9). In the TMPRSS2 complex, where the ligand binding site is at the central protease pocket of the protein, no remarkable changes were observed.

Fig. 4. The top 3 ligands with the highest binding energies with (a-c) Spike protein (d-f) Spike-ACE2 protein complex (g-i) TMPRSS2. The color depicts different interactions; Green for H-bonding, blue for halogen, pink and orange for different interactions involving pi bonds.

Fig. 5. RMSD of (a) Spike, (b) Spike-ACE2 and (c) TMPRSS2 with the three ligands are shown. Ligands 1, 2 and 3 are colored in black, red and green colors respectively. RMSD of the Spike-ACE2 ligand complex shows the existence of conformational changes.
conformational changes in the protein could be deciphered from the RMSD values.

After checking that the proteins are stable, we looked for the ligand occupancy in the binding pockets for all the systems (Table 3). Surprisingly we found that the inhibitors quickly left the binding pocket for all the three spike proteins and had less than 50% occupancy time. To check if this was an artefact, we simulated another docked complex but it also exhibited the same phenomena (data not shown). However, this peculiarity of ligand getting displaced from the binding pocket was not observed in the Spike-ACE2 complexes; in two out of the three systems, the ligand bound very well in the Spike-ACE-2 complex. Similarly, in TMPRSS2 also we observed that two out of three systems displayed stability. Comparison of the ligand binding sites in the initial and final states show that all the three quinoline analogues lost interaction with the Spike protein (data not shown). Thus, despite very high docking scores of ligands with the Spike protein, i.e., >–8 Kcal/mol, the binding affinity of the ligand is greatly affected by the dynamics of the binding site and the solvent environment.

We then checked for the binding of inhibitors to Spike-ACE2 protein.

It has been earlier reported that CQ and HCQ bind very well to the Spike-ACE2 complex [71,72]. The docked structures had shown very good binding energies and a consistent hydrogen bond with GLU 471. Thus, we wanted to verify if the structures are stable after simulating it for 20 ns. Out of the three ligands under study, we found that the third ligand bound very well, the second was displaced from the binding pocket but the first ligand had completely left the pocket. Fig. 6 shows that the third ligand has more interactions with the Spike-ACE2 complex when compared to the first two. The binding is facilitated by a strong hydrogen bond with GLU37 and halogen bond formed with GLY496, apart from that LYS353 and TYR505 have hydrophobic interactions with the inhibitor molecule. However, the second inhibitor is displaced from its original position losing in the process many important interactions that were there initially present. LEU73 hydrophobically interacts with the third inhibitor, however, in longer time runs the possibility of displacement of the inhibitor persists. TMPRSS2 is a relatively small protein compared to Spike or ACE2 receptors. When we compared the binding of the ligands only two ligands remained bound to TMPRSS2. Both ligands were observed to have consistent hydrogen bonds with GLN192 during the 20ns of simulation time frame (Fig. 7). Surprisingly, ligand two with salt bridge and large number of hydrogen bonding interactions lost its durability in the binding pocket. But considering an electronegative oxygen atom in the ligand structure, the possibility of repulsive interaction in the binding pocket cannot be ruled out.

One of the major reasons for the greater and effective binding of the fluorinated analogue to the protein receptor is due to the involvement of halogen bonding interactions which are seen in the fluorinated analogues, which involve the Fluorine of the ligand and the amide-carbonyl

|      | Ligand1 | Ligand2 | Ligand3 |
|------|---------|---------|---------|
| Spike| 42.35   | 11.5    | 28.15   |
| Spike-ACE2| 71.7 | 100     | 100     |
| TMPRSS2| 100    | 69.7    | 100     |

Fig. 6. Starting (a, c) and end (b, d) conformation of SPIKE-ACE2 complex with Ligands 2 (a, b) and 3 (c, d). Protein residues are shown in cyan and ligands are shown in orange color.
group of the protein residues, apart from this there are ring stacking interactions more in the fluorinated analogues as compared to HCQ/CQ. Electronegative GLU37 in Spike-ACE2 helps in the strong binding of quinoline. Similarly, TMPRSS2 inhibitors interacting with GLN192 help its stability. This result matches very well with our initial docked complexes of CQ and HCQ shown in Table 1. Spike-ACE2 interacts with HCQ in the docked complex via a hydrogen bond with GLU37. Similarly, both CQ and HCQ lie very close to GLN192 in docked complexes and could hydrogen bond in the long run. Additionally, one of the inhibitors docked with the proteins was the same, it was found to be quite stable in SPIKE-ACE2 and TMPRSS2 binding pockets.

The vacuum electrostatics of the docking pockets in the final conformation of Spike-ACE2 and TMPRSS2 show that the inhibitor binding pocket is negatively charged (Fig. 8). Results show that in Spike-ACE2 ligand 2 complex, the ligand is displaced from the docked site due to protein dynamics, the electronegative environment that was seen for the Spike-ACE2 ligand 3 and TMPRSS2 ligand 1 and 3 is different from Spike-ACE2 ligand 2 complex. Therefore, quinolines with electropositive groups would interact with Spike-ACE2 and TMPRSS2 proteins more favorably. The above observations from computational studies would prove very beneficial for fashioning quinoline structure-based drug designing studies for COVID-19.

4. Conclusion

The COVID-19 pandemic has affected crores of individuals all around the world. People are being vaccinated, however, recurrence of disease even after vaccination has been reported. New mutants of SARS-CoV-2 are also observed. In such a scenario, alternative therapeutic strategies by repurposing drugs have also been studied. CQ and HCQ used for treatment of malaria, rheumatoid arthritis and systemic lupus erythematosus were found to successfully prevent viral replication inside cells. However, the drastic side effects, such as QT prolongation, caused by the inhibitors, restricted its application for further treatment of COVID-19. Here, we implement an in-silico approach to propose novel quinoline based inhibitors which could be used as an alternative treatment strategy for COVID-19.

We studied the binding of HCQ and CQ with viral Spike, viral Spike-human ACE2 complex and human TMPRSS2 protease. All of these proteins play a role in initiation and assembly of the virus. Apart from ascertaining the binding site, we also got the threshold value of inhibitor binding energies. Later, we did an extensive search on the alternate quinoline analogues which could prove as promising candidates for drug designing. By our high throughput screening studies carried out for nearly 10000 quinoline based ligands, we were able to identify the best candidates, who also fulfill the ADME characteristics. The docking scores of the inhibitors were found in the range of 8kcal/mol. In order to understand the effect of solvent and protein dynamics in physiological conditions, we performed Molecular Dynamics (MD) simulations of the proteins with the docked ligands and studied the changes in interaction over a period of time.

In MD simulations, we observed that the protein dynamics and effect of solvation does not permit long term stability of ligands with the viral spike protein. None of the ligands remained in the binding pockets with the Spike protein alone. On the contrary, when the quinoline inhibitors bound to Spike-ACE2 protein complex and TMPRSS2 protease, the occupancy of the inhibitors was observed to be for a longer period. It has been observed through many studies that addition of fluorine to ligands
has a significant effect not only on the binding of the ligand but also on the physicochemical properties [73]. Firstly, the metabolic stability of the molecules changes significantly, and therefore inclusion of fluorine atoms lends metabolic stability by blocking the labile site [74,75]. Secondly, it is found that fluorine substitution leads to a slight enhancement of binding affinity due to an increased lipophilicity of the molecule and one of the most major effects of fluorine is the way it indirectly influences the acidity/basicity of the ligand. The study by van Niel et al. describes how fluorinated compounds result in drastic improvement in the pharmacokinetic properties of the ligands [76]. In one of the studies by Olsen et al. they showed how the monofluorinated analogue of a thrombin inhibitor binds five times stronger than its non-fluorinated parent compound [77]. The CF3 group containing fluoro-pharmaceuticals is the second largest Fluorine based FDA approved drug and along with mono fluorinated moieties constitute as much as 86% of all FDA approved Fluorine inhibitors. Moreover, fluoroquinolones like ciprofloxacin, norfloxacin, and levofloxacin, are widely used fluoro-pharmaceuticals for treatment. Comparative docking scores are shown in Table S10 [78]. The macromolecular electrostatics shows that the binding pocket is partially negatively charged in nature and would favor the binding of quinoline inhibitors with electropositive groups. From our studies, we found that quinoline based inhibitors prefer binding to TMPRSS2 human protease or viral spike bound to human ACE2 receptors. The inhibitor binding pocket is largely electronegative in nature. The lead compound, 3-[3-(Trifluoromethyl)phenyl]quinoline bound favorably with both Spike-ACE2 complex and TMPRSS2 which could be considered as a candidate for future drug designing studies.

Author contribution

NS, AT and JG designed research; NS, AT, JG and SLR performed research; NS, AT, JG and SLR analyzed data; NS, AT, JG and SLR wrote the paper.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.jfluchem.2021.109865.

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