ABSTRACT

Background: HCQ has been found to provide the immunomodulation that helps to abate the possible cytokine storm response caused by the SARS COVID-19 virus and subsequent failures of major organs. It is also being used historically in treating Malaria and Rheumatoid arthritis and is known to be useful in cancer treatment. However, it shows a half-life of up to 70 days and has severe adverse effects including gastric disorders and retinopathy.

Hypothesis: Therefore, we propose to explore some of its metabolites/intermediates namely Desethyl hydroxychloroquine (DHCQ), Desethyl chloroquine (DCQ), and Bis-desethyl chloroquine (BDCQ) as alternate drug systems with minimized toxicity.

Results and Discussion: The structural and spectral studies of these lead compounds to be unraveled using computational modeling and the evaluation of ADME parameters such as absorption, distribution, metabolism, and excretion would be performed using the SwissADME tool to assess the drug-likeness and medicinal chemistry friendliness. The future scope of the work involves the docking studies of these compounds to assess the therapeutic compliance for the treatment of viral infections and rheumatoid arthritis.

Key words: Hydroxychloroquine metabolites, COVID-19 drugs, Computational, ADME studies, Rheumatoid arthritis, Immunomodulation.

INTRODUCTION

HCQ, a known antimalarial drug has assumed importance during the current pandemic COVID-19, due to its potential use in severe cases of infections. The current treatment protocol currently involves the management of symptoms and other antiviral support. HCQ, derived from CQ, has been found to provide immunomodulation which helps to abate the possible cytokine storm response caused by the virus and subsequent failures of major organs. Both CQ and HCQ reportedly hinder the binding of viruses onto cell membranes by altering the surface pH and further block the viral multiplication. Though CQ has been initially chosen as the antiviral drug for Covid, due to its availability and cost-effectiveness, an overdose of CQ can cause severe poisoning even leading to morbidity. On the other hand, HCQ, derived from CQ, has an equivalent drug efficacy but with a reduced toxicity impact. A recent comparative study on antiviral potency showed that HCQ provides a better antiviral effect than CQ in treating COVID-19. HCQ has also been explored recently in cancer treatment to inhibit auto phagocytosis or the self-devouring of body cells. Another major application of HCQ is for treating rheumatoid arthritis; however, it suffers from the major side effect of induced retinopathy and GI disorders.

Both CQ and HCQ are 4-aminoquinoline derivatives with a planar aromatic core and a slightly basic side chain which helps in the intracellular lysosomal accumulation of drugs. These molecules possess chiral centers and hence are present as enantiomers. The S (+) enantiomer binds more strongly to the plasma protein as compared to the R (-) counterpart. Upon oral administration,
HCQ undergoes CYP450 assisted metabolism to produce major metabolites namely DHCQ, DCQ, BDCQ. Among these DHCQ has been identified as the major metabolite even with a therapeutic value greater than HCQ itself while BDCQ has been found to cause the listed side effects.

With the novel Coronavirus undergoing rapid genetic mutations, new treatment options and drug combinations are needed to be explored. Potential compounds with possible antiviral properties need to be identified and qualified to keep in mind the possibility of future pandemics and also to reduce the cost of treatment and hospitalizations. Computational studies often provide testable evaluations of probable drug systems.

In the present study the three metabolites of HCQ namely DHCQ, DCQ, BDCQ are selected for the in silico structural and spectral studies to understand the structure-activity relationships. These compounds are also subjected to Absorption, Distribution, Metabolism, and Excretion (ADME studies) computationally to assess the drug likeness and medicinal chemistry friendliness.

**MATERIALS AND METHODS**

**DFT Calculations**

The ab-initio computational modeling for the optimization and spectral studies of the molecules were performed using the Gaussian09 software package. The graphical interface GaussView 6.0 was used for creating the input structures and then interpreting the output files.

**ADME Studies**

The ADME parameters were analyzed using the web tool SwissADME. The physiochemical properties of the molecules were evaluated by using the bio-availability radar and the BOILED- Egg plots available with the program.

**RESULTS**

**DFT STUDIES**

**Structural Optimization**

The optimized structures were computed with Gaussian 09 using the Density Functional Theory approach and B3LYP/6-311 basis sets. The optimized structures of DHCQ, DCQ and BDCQ corresponding to the global minimum energies were appended below with atom labeling (Figure 1). The global minimum energies were computed at -1322.58, -1247.3 and -1168.75 a.u for DHCQ, DCQ, and BDCQ respectively while the corresponding dipole moments were calculated at 6.66, 6.41 and 6.55 D. Similarly, the polarizabilities of the metabolites were computed at 206.9, 204.23, and 188.39 Bohr for DHCQ, DCQ, and BDCQ respectively. Both dipole moments and polarizabilities are useful indicators for molecular docking investigations.

The three-dimensional molecular geometry and the corresponding geometrical parameters derived computationally reveal useful information regarding the drug-receptor interactions. The geometrical parameters such as bond lengths and bond angles of the metabolites are computed and presented below in Table 1. The computed parameters are then juxtaposed against the experimentally derived data corresponding to the parent molecule CQ for comparison.

**Frontier Orbital Analysis**

The frontier molecular orbitals namely Highest Occupied Molecular Orbitals (HOMO) and Lowest Occupied Molecular Orbitals (LUMO) determine the extent of interaction of drug systems with receptors and other molecules. HOMO and LUMO energies describe the electron releasing and electron-withdrawing capabilities respectively and depict the thermodynamical stability and chemical reactivity of the systems. These are useful descriptors in gauging the propensity of HOMO to LUMO electron transfer and hence the stability of the systems. The detailed HOM- LUMO profiles of DHCQ, DCQ, and BDCQ and global reactivity descriptors such as electronic chemical potential, absolute electro negativity, global softness, global hardness, and electrophilicity are presented in Table 2. According to Koopman’s theorem, the ionization potential (I), and electron affinity (A) are calculated as the negative values of the energy of the highest occupied molecular orbital and the lowest unoccupied molecular orbital respectively.
The absolute electronegativity ($\chi$) is calculated using the equation $\chi = \frac{1 + A}{2}$ and the electronic chemical potential ($\mu$) is computed as $\mu = -\chi$. The value of global hardness ($\eta$) and softness ($S$) were obtained using the equations $\eta = \frac{ELUMO - EHOMO}{2}$ and $S = \frac{1}{2\eta}$ respectively. The electrophilicity ($\omega$) of the compounds were calculated using $\omega = \frac{\mu}{2\eta}$ while another important parameter the maximum charge transfer index was calculated by the formula $\Delta N_{\text{max}} = -\frac{\mu}{\eta}$.

As the higher value of HOMO energy and the lower value of LUMO energy imply better electron donating and accepting tendencies respectively, it is evident that DHCQ has the most electron- donating tendency while BDCQ has the most electron-withdrawing tendency. The HOMO-LUMO energy gap of all the three molecules is more or less comparable and DHCQ shows the least value while BDCQ showing a relatively larger band gap. Further BDCQ demonstrates the highest ionization potential and electron affinity indicating the least chemical reactivity. Conversely, DHCQ has the lowest ionization potential and electron affinity making it the most reactive system. This was further corroborated by the lowest value of the HOMO-LUMO energy gap for DHCQ. Electrophilicity index ($\omega$), which reflects the reduction of energy during the maximal flow of electrons between two systems\textsuperscript{20} would show us whether the species is capable of chargedonation.\textsuperscript{21} As a thumb rule, a species with lesser `\omega' would be more reactive and
nucleophilic and those with a higher value would go on to become a better electrophilic system. The studies revealed that DHCQ promises to be a better nucleophile while BDCQ showed a better propensity of an electrophile. This implies that BDCQ will be able to receive an electron doublet so as to react with other nucleophilic systems. Both DHCQ and DCQ demonstrated lower values of absolute electron negativity (χ) making them more basic as compared to BDCQ. This is an important aspect considering the fact that the basicity of the side chain governs the intracellular lysosomal activity of the drug molecules.

The three-dimensional HOMO-LUMO plots of the molecules are illustrated in Figure 2. It is evident that the HOMO density is populated over the aromatic core and slightly onto the chlorine and nitrogen of the side chain with no population seen over the rest of the basic side chain. On the other hand, the LUMO population is by large populated over the whole basic side chain with no apparent population over the aromatic core.

In the HOMO-LUMO diagram, the green and red colors symbolize the negative and positive charge fields respectively. This can be further substantiated by plotting the Density of States (DOS) plot which describes the number of states available at various energy levels (Figure 3). The DOS plots also visualize the occupied and virtual orbitals of the molecules and also compute the HOMO-LUMO energy gap. The relatively lower values of HOMO-LUMO energies of DHCQ, DCQ, and BDCQ at 4.019, 4.081, and 4.353 eV respectively indicate ease of transfer of electrons in the molecules.

**Mulliken Charge Analysis**

The Mulliken atomic charges demonstrate the electron charge distributions in molecules. The most negative site of the molecule will be the one most susceptible for an electrophilic attack while the most positive sites will be more liable for nucleophilic attack. The Mulliken analysis for the title molecules was performed computationally at DFT-B3LYP/6-311g and the results were tabulated in Table 3.

Our results showed N17 and O42 positions as the most negative sites for DHCQ and N17 and N30 positions for both DCQ and BDCQ indicating possibilities of electrophilic attacks at these sites. Conversely, the C8 position has been found to be the most positive and hence liable for nucleophilic attack in all three molecules. The structure-activity relationship of potential drug molecules including susceptibility towards nucleophilic and electrophilic attack and other intermolecular interactions can also be validated by performing molecular electrostatic potential (MEP) studies. MEP plots help to picturize regions of variable charges by using color grading. It also envisages the size, shape, and reactive behaviors of molecules. These three-dimensional diagrams visualize negative, positive, and neutral electrostatic potential regions by virtue of the color gradient; with red and blue colors representing the most electronegative and electropositive regions respectively. The regions with zero potential are shown in green color. The MEP plots of DHCQ, DCQ
and BDCQ were generated computationally and are presented below in Figure 4.

**Spectral Studies**

Even though spectral studies are not always considered for geometrical and structural optimization, FTIR data can corroborate the optimized geometry by providing positive force field constants. Here, the FTIR scans of the title molecules were performed computationally using the DFT-B3LYP/6-311g basis set. The computed FTIR spectra of the molecules showed the characteristic peaks as expected (Figure 5). There were no negative peaks reported suggesting that the most optimized structures predicted were indeed corresponding to the global minimum energy. DHCQ showed the characteristic OH stretching vibration at 3700 cm$^{-1}$ and a broad peak at around 1600 cm$^{-1}$ corresponding to C=C and C=N stretching modes. The notable C-Cl peak has been credited to the high intense peak around 650 cm$^{-1}$. DCQ showed the typical CH vibrations around the 3000 cm$^{-1}$ mark and C-Cl peak close to 600 cm$^{-1}$. The characteristic twin peaks for the terminal amino group have been visualized at 3500 cm$^{-1}$, followed by CH stretching modes at around 3000 cm$^{-1}$. Other notable
peaks were noted at 1650 and 700 cm\(^{-1}\) corresponding to C-N and C-Cl stretching modes.

**ADME Studies**

The evaluation of ADME parameters namely Absorption, Distribution, Metabolism and Excretion provide crucial pre-clinical information to assess the suitability drug-like molecules. It is imperative that the potential drug system reaches the target site in the stipulated amount and be available in the correct quantity till its pharmacological functions are performed.\(^{22}\) The ADME parameters of DHCQ, DCQ, and BDCQ were assessed using the SwissADME web tool to evaluate the flexibility, lipophilicity, size, polarity, solubility, and saturation. The critical limits of the above-listed parameters were visualized using the bioavailability radar (Table 4). It also provides useful acumen regarding pharmacological attributes namely gastrointestinal absorption (HIA), blood-brain-barrier (BBB) permeability, topological polar surface area (TPSA), etc. The bioavailability radars of DHCQ, DCQ, BDCQ and the parent molecules such as CQ and HCQ were depicted in Figure 6 for comparison.

From Table 4, it is evident that none of the lead molecules or parent drugs showed any Lipinsky violations indicating overall druglikeness. However, DHCQ and DCQ showed one potential violation in the leadlikeness index while BDCQ showed none. The overall physiochemical parameters of the metabolites and drug systems are comparable. Of the three metabolites, DHCQ possesses 3 hydrogen bond acceptors and 3 hydrogen bond receptors making it more reactive with the receptor sites. Conversely DCQ and BDCQ possess 2 each of these. The results reveal that both DHCQ and BDCQ have lower logP values compared to DCQ and the parent drugs. This is advantageous as the toxicity of the drug systems decreases as logP value decreases and systems with logP value around 2.2 are more suitable for oral administration.\(^{23-25}\) It is worth noting that CQ has a logP value around 4.2 well above the toxicity cut-

### Table 4: The physiochemical properties of lead molecules compared against the parent molecules.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>CQ</th>
<th>HCQ</th>
<th>DHCQ</th>
<th>DCQ</th>
<th>BDCQ</th>
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<tbody>
<tr>
<td>MW</td>
<td>319.87</td>
<td>335.87</td>
<td>307.82</td>
<td>291.82</td>
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<td>23</td>
<td>21</td>
<td>20</td>
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<tr>
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<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>#Rotatable bonds</td>
<td>8</td>
<td>9</td>
<td>8</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>#H-bond acceptors</td>
<td>2</td>
<td>3</td>
<td>3</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>#H-bond donors</td>
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<td>3</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Molar refractivity, MR</td>
<td>97.41</td>
<td>98.57</td>
<td>88.86</td>
<td>87.7</td>
<td>77.99</td>
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<tr>
<td>TPSA</td>
<td>28.16</td>
<td>48.39</td>
<td>57.18</td>
<td>36.95</td>
<td>50.94</td>
</tr>
<tr>
<td>Consensus Log P</td>
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<td>3.29</td>
<td>2.77</td>
<td>3.58</td>
<td>2.83</td>
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<td>ESOL Log S</td>
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<td>Moderate</td>
<td>Soluble</td>
<td>Soluble</td>
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<tr>
<td>GI absorption</td>
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<td>High</td>
<td>High</td>
<td>High</td>
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<td>BBB permeant</td>
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<td>Yes</td>
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<td>CYP1A2 inhibitor</td>
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<td>CYP2D6 inhibitor</td>
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<td>CYP3A4 inhibitor</td>
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<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td>log Kp (cm/s)</td>
<td>-4.96</td>
<td>-5.81</td>
<td>-6.23</td>
<td>-5.38</td>
<td>-5.84</td>
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<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Synthetic Accessibility</td>
<td>2.76</td>
<td>2.82</td>
<td>2.6</td>
<td>2.54</td>
<td>2.35</td>
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</table>
off limit of 3.0. Among the metabolites and parent molecules, DHCQ possesses the highest TPSA value due to the presence of polar hydroxyl group in the terminal position of the side chain. Studies revealed that an increased TPSA value and a lower logP value ensure better bioavailability of the drug. The value of TPSA is an important parameter considering the fact that as TPSA increases the blood-brain-barrier (BBA) index decreases. However, all the five molecules considered here have TPSA values well below the cut-off value of 90 Å² making them BBB permeant. The metabolites, by and large, showed comparable inhibitory actions towards the major cytochromes as compared to the parent drugs.

The BOILED-Egg plot is the graphical output of the SwissADME software which showcases the gastro-intestinal absorption and CNS absorption rates of the potential drug molecules Figure 7. It is inferred that those systems that appear in the egg's white would exhibit better gastro-intestinal absorption and those in the yellow yolk are better drugs for the central nervous system applications. In the BOILED-Egg plot, the molecules located in the ‘egg white’ will have more GI absorption, whereas those in the ‘yolk’ have more BBB permeation and CNS penetration. Molecules in the grey area have often shown poor activity as GI or CNS drugs. Further, the compounds that are colored blue get pumped out of the brain while those in red remain in the brain. The BOILED-Egg plot suggests that both the metabolites and the parent drugs have BBB permeation and have an impact on the CNS system.

**CONCLUSION**

Though both CQ and HCQ are known to possess good antiviral and anti-arthritis properties, they suffer from various side effects and prolonged half-life. Three major metabolites of HCQ namely DHCQ, DCQ, and BDCQ were selected for structural, spectral, and physiochemical assessment computationally. The optimized structure and geometrical parameters were computed, and the data showed concordance with the experimental results available in the literature. The HOM-LUMO energies of DHCQ, DCQ and BDCQ were computed along with the global reactivity descriptors such as electronic chemical potential, absolute electro negativity, global softness, global hardness, electrophilicity. Of the three metabolites, DHCQ showed the lowest ionization potential and electron affinity making it the most reactive system. This was further substantiated by having a lower HOMO-LUMO energy gap. DHCQ also exhibited the lowest value of absolute electro negativity (χ) making it more basic. This is an important aspect considering that the basicity of the side chain governs the intracellular lysosomal activity of the drug molecules. The Mulliken charge analysis showed N17 and O42 positions as the most negative sites for DHCQ and N17 and
N30 positions for both DCQ and BDCQ indicating possibilities of electrophilic attacks at these sites. The assessment of ADME properties revealed that none of the metabolites showed any Lipinsky violations indicating good overall drug likeness. Of the three metabolites, DHCQ possesses 3 hydrogen bond acceptors and 3 hydrogen bond receptors making it more reactive with the receptor sites. The results reveal that both DHCQ and BDCQ have lower log P values compared to the parent drugs which make them less toxic and suitable for oral intake. Both metabolites and parent drugs demonstrated very low values of TPSA and hence increased BBB permeation. In general, it can be concluded that DHCQ may be considered as a better lead molecule for the potential drug development and docking studies compared to DCQ and BDCQ.

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

ABBREVIATIONS
HCQ: Hydroxychloroquine; CQ: Chloroquine; SARS: Severe acute respiratory syndrome; DHCQ: Desethyl hydroxychloroquine; DCQ: Desethyl chloroquine; BDCQ: Bis-desethyl chloroquine; ADME: absorption, distribution, metabolism, and excretion; TPSA: Topological polar surface area; BBB: Blood-brain-barrier; HOMO: Highest occupied molecular orbital; LUMO: Lowest unoccupied molecular orbital; GI: Gastrointestinal; DFT: Density functional theory; MEP: Molecular electrostatic potential; HIA: Gastrointestinal absorption; FTIR: Fourier-transform infrared spectroscopy.

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