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ORIGINAL ARTICLE

One-pot synthesis, X-ray crystal structure, and identification of potential molecules against COVID-19 main protease through structure-guided modeling and simulation approach

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Received 2 March 2022; accepted 31 August 2022
Available online 15 September 2022

KEYWORDS
Dihydropyrimidinones; SARS-CoV-2 Mpro; Hirshfeld surface analysis; NBO analysis; Molecular docking

Abstract Although antimicrobial resistance before the Covid-19 pandemic is a top priority for global public health, research is already ongoing on novel organic compounds with antimicrobial and antiviral properties in changing medical environments in connection with Covid 19. Thanks to the Biginelli reaction, which allows the synthesis of pyrimidine compounds, blockers of calcium channels, antibodies, antiviral, antimicrobial, anti-inflammatory, or antioxidant therapeutic compounds were investigated. In this paper, we aim to present Biginelli’s synthesis, its therapeutic properties, and the structural–functional relationship in the test compounds that allows the synthesis of antimicrobial compounds. Both the DFT and TD-DFT computations of spectral data, molecular orbitals (HOMO, LUMO) analysis, and electrostatic potential (MEP) surfaces are carried out as an add-on to synthetic research. Hirshfeld surface analysis was also used to segregate the different intermolecular hydrogen bonds involved in the molecular packing strength. Natural Bond Orbital (NBO) investigation endorses the existence of intermolecular interactions mediated by lone pair, bonding, and anti-bonding orbitals. The dipole moment, linear polarizability, and first hyperpolarizabilities

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Peer review under responsibility of King Saud University.

https://doi.org/10.1016/j.arabjc.2022.104230
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have been explored as molecular parameters. All findings based on DFT exhibit the best consistency with experimental findings, implying that synthesized molecules are highly stable. To better understand the binding mechanism of the SARS-CoV-2 M\textsuperscript{pro}, we performed molecular docking, molecular dynamics (MD) simulations, and binding free energy calculations.

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1. Introduction

In 1843, Pietro Biginelli was discovered. The reaction is based on the condensation of aldehydes, urea, and 1,3-dicarbonyl compounds in an acidic medium. As a result of the synthesis, heterocyclic systems of dihydropyrimidinones are formed, which play an important role in the construction of biomolecules such as DNA and RNA. In addition, in the fields of application, the synthesis and development of materials with optical properties, the design of polymers, dyes, and adhesives are also included. The most widely used area of biological impact is medicine. The multicomponent nature of the Biginelli reaction provides large product diversity bearing different pharmacophoric groups in the structure of dihydropyrimidines, which ensure optimal supramolecular interactions with a specific biological target (molecular recognition) and triggers or blocks its biological response. Investigations through molecular manipulations reveal that this class of compounds demonstrate antiviral, anti-filarial, antifungal, analgesic, antileishmanial, antiproliferative, antitumor, anti-convulsant, antibacterial, anti-inflammatory, anti-hypertensive, melanin-concentrating hormone 1 receptor antagonist, anti-HIV, antiepileptic, anti-diabetic, anti-SARS, anti-malarial, pages-1 inhibitors, antihypoglycemic, antituberucleral, TRPA1 antagonist, miscellaneous, potassium and calcium channels, and a1a adrenergic antagonists. Similarly, the structural core of quinoline is frequently associated with medicinal applications such as anticancer, antimicrobial, HIV-1 integrase inhibition, HIV protease inhibitors, antileishmanial activity, NK-3 receptor antagonists, PLT antagonists, and antimalarial activity. As a result of these studies, the use of various types of drugs such as riboflavin, idoxuridine, aminophylline, emivirine, 5-fluorouracil, methylthiouracil, batzelladine A, B based on dihydropyrimidines has allowed widespread use in medicine. Due to the utmost importance of dihydropyrimidines in pharmaceuticals, robust, efficient, cost-effective, and “green” chemical synthesis and transformations of dihydropyrimidine scaffolds were developed (Kaur et al., 2015; Nagarajaiah et al., 2016; Terracciano et al., 2015; Yadlapalli et al., 2012; Zhu et al., 2010). (See Scheme 1 Scheme 2.)

In the biological aspects of the motif, fixed molecules are needed that can easily be obtained for evaluation/study, and in 1930, the woolly-containing activity of these molecules was patented. These molecules bore similarities to the clinical spectrum of nifedipine. Biginelli researched by analogy with their analogs, and later it turned out that they showed the similarity of natural sea alkaloids with batzelladine B. All three building blocks are aldehyde, active methylene, and urea and are caused by the molecular diversity of dihydropyrimidines, which have a multicomponent reaction. The potent pyrimidine molecules produced by the Biginelli reaction have been successfully used and experimented with in stress and pollution-induced diseases. The following are selective molecules having significant activity, and they are examined with clinically used drugs in vivo/in vitro, establishing QSAR (Ashok et al., 2007; Chiang et al., 2009; Chitra et al., 2010; Deshmukh et al., 2009; Kidwai et al., 2005; Rajanarendar et al., 2010). There are several reasons that Biginelli’s reaction has been studied in recent decades (Bais et al., 2020; de Fátima et al., 2015; Gireesh et al., 2013; Ismaili et al., 2008; Kaur et al., 2017; Li et al., 2020; Liu et al., 2019; Mokale et al., 2010; Rani et al., 2016; Sawant and Sarode, 2011; Silva et al., 2015).

As well as the inhibitory potential of antimicrobial and antibacterial pyrimidine derivatives has been evaluated against SARS-CoV-2 (Khan et al., 2021). Considering the importance of this class of compound, herein, two compounds are synthesized, characterized, and used in different computational approaches to unveil their anti-SARS-CoV-2 activity. Computational drug design has been a good platform for predicting the binding capacity of compounds to different therapeutic targets and thus saves time and cost that are usually required in experimental testing. In this study, the docking studies are revalidated by more sophisticated molecular dynamics simulation.

![Scheme 1](image-url)
and binding free energy analysis. In a nutshell, the findings obtained in this study could be useful for experimentalists to test the compounds against SARS-CoV-2 and in particular to be tested in enzymatic assay against the SARS-CoV-2 main protease enzyme.

2. Materials and methods

2.1. Instrumentation

The NMR spectra were recorded at 25 °C from solutions in DMSO-d$_6$ on a Bruker-300 spectrometer operating at 300 MHz. The IR spectra were obtained on a Specord 75IR instrument from samples dispersed in mineral oil. The progress of reactions and the purity of products were monitored by TLC.

2.2. X-ray diffraction study

Single crystals of compounds I and IV suitable for X-ray analysis were obtained by double recrystallization from ethanol (Kurbanova, 2010). The X-ray diffraction data were acquired on a Bruker APEX II CCD diffractometer (100 k), λMoK$_\alpha$ irradiation, graphite monochromator, φ- and ω-scanning, 20max = 56°. The structures were solved by the direct method and refined by the least-squares procedure in an anisotropic approximation for non-hydrogen atoms. Hydrogen atoms in the hydroxy and amino groups were localized by the Fourier difference syntheses, and their positions were refined in an isotropic approximation with fixed positional and thermal parameters. The positions of the other hydrogen atoms were calculated based on geometry considerations and were refined in isotropic approximation with fixed positional and thermal parameters. All calculations were performed using the SHELXTL PLUS and SADABS software packages (Sheldrick, 2003, n.d.). The complete sets of crystallographic data for compounds I and IV were deposited at the Cambridge Crystallographic Data Centre (entry nos. CCDC 694,407 and CCDC 707349).

2.3. Computational studies

Computational studies of the compounds were performed to evaluate their binding affinity for different anti-SARS-CoV-2 targets. These targets include the SARS-CoV-2 main protease enzyme, RNA-dependent RNA polymerase (RdRp), and envelop (E) protein. The crystal structures of these targets were retrieved from protein data banks and underwent preprocessing in UCSF Chimera 1.15 (Pettersen et al., 2004). The PDB IDs used for retrieval of the proteins are the main protease enzyme (6y2e), RdRp (7D4F), and E (7M4R). During preprocessing, all co-crystalized ligands were removed, and the structures were relaxed using the steepest descent and conjugate gradient algorithms using default parameters.

2.4. Molecular docking and dynamics simulation studies

Molecular docking of the compounds with the targeted enzymes was conducted through AutoDock Vina software (Trott and Olson, 2010). Before that, the compound structure was drawn in ChemDraw 12.0 (Cousins, 2011) and processed into.pdb format. The proteins were also imported into the docking software. The grid box was set around the active site of the enzyme along a 15 Å on XYZ axis. The number of compounds docked was set at 100. Each compound conformation is associated with binding energy, and the one with the lowest score was complexed with the best-docked enzyme. Visualization of complexes was done in UCSF Chimera 1.15 (Kaliappan and Bombay, 2016) and Discovery studio v2021 (Biovia, 2017). Further, the best-docked complex for both compounds was subjected to molecular dynamics simulation for 300 ns. Simulations were performed via AMBER20 software (Case et al., 2020). In the simulation, the protein parameters were prepared using ff14SB (Maier et al., 2015), while the compounds were parameterized using GAFF (He et al., 2020). Energy minimization of complexes was accomplished via steepest descent and conjugate gradient steps, each of which was run for 1000 cycles. Heating was achieved at 310 K for 100 ps. Equilibration was done for 200 ps. SHAKE and
Langevin were used to constrain hydrogen bonds and maintain temperature during a production run, respectively (Izaguirre et al., 2001; Krätutler et al., 2001). CPPTRAJ was applied to analyze simulation trajectories (Roe and Cheatham III, 2013). MMPBSA.py module of AMBER was considered for MMGBSA and MMPBSA binding free energies estimation on 100 frames of simulation trajectories (Genheden and Ryde, 2015; Miller et al., 2012).

3. Results and discussion

3.1. Chemical synthesis

The Biginelli reaction of salicylaldehyde derivatives can lead to the formation of 2 types of dihydropyrimidines (Kurbanova, 2010). The reaction of 5-bromosalicylaldehyde with ethyl acetoacetate and urea, depending on the conditions, ethyl 4-(5-bromo-2-hydroxyphenyl)-6-methyl-2-oxo-1,2,3,4-tetrahydropyrimidine-5-carboxylate (I) or ethyl 8-bromo-2-methyl-4-oxo-3,4,5,6-tetrahydro-2H-2,6-methanobenzo[g][1,3,5] oxadiazocine-11-carboxylate (II). Ester, was formed when its structure was proved by IR and NMR spectroscopy and X-ray crystallographic analysis performed by double X-ray diffraction data.

Ethyl 4-(5-bromo-2-hydroxyphenyl)-6-methyl-2-oxo-1,2,3,4-tetrahydropyrimidine-5-carboxylate (I). Yield 75 %, mp = 19–191 °C. IR spectrum, v, cm⁻¹: 3325, 1745, 1660. ¹H NMR spectrum, v, ppm: 1.30 t (3H, CH₂CH₃), 2.57 s (3H, CH₃), 4.25 q (2H, CH₂O), 5.70 s (1H, CH), 7.70–7.00 m (4H, Harom), 8.00 s (1H, NH), 9.41 s (1H, NH), 10.11 s (1H, OH). (See Fig. 1).

Ethyl 8-bromo-2-methyl-4-oxo-3,4,5,6-tetrahydro-2H-2,6-methanobenzo[g][1,3,5] oxadiazocine-11-carboxylate (II) Yield 55 %, mp = 197–199 °C. IR spectrum, v, cm⁻¹: 3350, 1740, 1680. ¹H NMR spectrum, v, ppm: 1.27 t (3H, CH₂CH₃), 2.35 s (3H, CH₃), 3.27 d (1H, CH), 4.16 q (2H, CH₂O), 4.46 d (1H, CH), 6.77–7.21 m (3H, Harom), 8.36 s (1H, NH), 8.41 s (1H, NH). X-ray crystallographic analysis was performed by double crystallization of compound I in ethyl alcohol (Kurbanova et al., 2009). The monoclinic, yellow crystals, with sizes 0.20 × 0.20 × 0.20 mm³, one striped: a = 9.3207(2), b = 16.841(2), c = 10.0908(13) Å, β = 116.678(1), V = 1415.3(3) Å³, space group P2₁/m, Z = 4, ds = 1.667 g/sm³, µ = 2.922 mm⁻¹ were obtained.

For comparison, three-component condensation of 5-bromosalicylaldehyde with acetylacetone and thiourea was examined. Under analogous conditions, the products were I- 1-[4-(5-bromo-2-hydroxyphenyl)-6-methyl-2-thioxo-1,2,3,4-tetra-
2018). The compounds I and IV were subjected to the Hirshfeld surface mapping across the normalized contact distance \( d_{\text{norm}} \) shown in Fig. 4. The color schemes shown in the map are red, blue, and white. The red \( (d_{\text{norm}} \) is negative) and blue \( (d_{\text{norm}} \) is positive) colored regions on the surface are from shorter and longer contacts than van der Waals radii respectively, whereas the white areas indicate the van der Waals radii of the contacts, or \( d_{\text{norm}} = 0 \) (McKinnon et al., 2007). In the brighter red zone, hydrogen bonding is responsible for the strong interactions, but electron density in the blue zone represents the weaker interactions. Qualitative measures of morphological structure, such as the “shape index,” can be sensitive to even the tiniest changes in surface curvature (or curvedness). The concave region with red triangles in the shape index of atoms of a molecule represents the \( \text{C} - \text{H} \cdot \pi \) and \( \pi \cdot \pi \) stacking interactions, while convex regions with blue triangles represent aromatic ring atoms inside the surface. For compounds I and IV, massive green flat areas with blue outlines on the relatively curved surfaces were observed due to the \( \pi \cdot \pi \) stacking interactions as shown in Fig. 3.

The intermolecular percent contribution of all kinds of contacts to the overall Hirshfeld surface area was calculated using 2D fingerprint plots (Seth, 2018). Fig. 4 shows the 2D fingerprint of all the contacts. The major contribution of both I and IV is from \( \text{H} \cdot \text{H} \) contacts, 38.0 %, and 37.1 % respectively, to the total Hirshfeld surface area. The remaining contributions of C1 are from O\( \cdots \text{H} \), C\( \cdots \text{H} \), Br\( \cdots \text{H} \), Br\( \cdots \text{C} \), and N\( \cdots \text{H} \) contacts, which contribute about 12.3 %, 9.6 %, 8.1 %, 2.1 %, 1.7 % respectively to the Hirshfeld surface area. The remaining contributions of C2 are from Br\( \cdots \text{H} \), O\( \cdots \text{H} \), H\( \cdots \text{S} \), C\( \cdots \text{H} \), C\( \cdots \text{C} \), Br\( \cdots \text{C} \), Br\( \cdots \text{Br} \), and Br\( \cdots \text{S} \) contacts, which contribute about 8.6 %, 7.2 %, 5.6 %, 5.6 %, 2.7 %, 1.7 %, 0.9 %, 0.6 % respectively to the total Hirshfeld surface area. Spikes arise in the fingerprint plots showing greater intermolecular interactions in the crystal packing of both I and IV.

3.4. 3D energy frameworks analysis

Energy framework analysis reveals the quantitative study of interaction energies and supramolecular design of molecules in a crystal. A cluster of molecules within 3.8 \( \text{Å} \) of the radius was built around a single I and IV compound. The scale factors used for developing the energy framework for B3LYP/6-31G(d,p) electron densities are \( k_{\text{ele}} = 1.057 \), \( k_{\text{pol}} = 0.740 \), \( k_{\text{disp}} = 0.871 \), \( k_{\text{rep}} = 0.618 \) (Edwards et al., 2017). Table 1 summarizes the computed interaction energies in kJ/mol and the symmetry operations for eight molecules of I. The blue-colored molecule positioned at 10.44 \( \text{Å} \) has the greatest total interaction energy (-74.7 kJ/mol), whereas the green-colored one positioned at 10.09 \( \text{Å} \) has the lowest total interaction energy (0.9 kJ/mol). Similarly, as shown in Table 1, the computed energy interactions and symmetry operations for seven IV molecules are also given in kJ/mol.

S-axis is shown in Fig. 5. The Coulomb interactions in I look to be comparable in strength as R values are closer for both. The calculated interaction energies for electrostatic, polarization, dispersion, and repulsion of I are -111.0, -31.0, -173.1, and 176.6 kJ/mol, respectively. The calculated interaction energies for electrostatic, polarization, dispersion, and repulsion of C2 are -194.4, -53.1, -235.8, and 299.4 kJ/mol, respectively. The total energies of I and IV are -182.1 and -265.2 kJ/mol respectively. The dispersion energy has the highest value among all interaction energies in I and IV, i.e., the dispersion interaction energy dominates over the electrostatic Coulomb interaction energy. This is because of the presence of the bromo group, which has a large electron cloud in each compound. The more electrons an atom or molecule has (the most polarizable), the stronger dispersion forces are. The scale factors for benchmarked energies used for the construction of energy models were taken from Mackenzie et al. (Mackenzie et al., 2017).

The visualization of different interaction energies like Coulomb interaction energy, dispersion energy, and total interaction energy is represented by red, green, and blue colors, respectively, for the compounds, I and IV along different axes as shown in Fig. 6. The cylinders in the energy framework represent the relative strengths of molecular packing in several directions. An overall scale factor is used to contract or expand the size of the cylinders in the framework (Turner et al., 2015). There is an absence of cylinders in a particular direction due to the exclusion of a few interactions under certain threshold energy. These weaker interactions have been ignored only to make the figures less crowded.

3.5. Structure optimization

The synthesized compounds (I and IV) are optimized at the B3LYP/6-31G(d,p) level of theory using Gaussian 16 (Frisch et al., 2016) together with GuassView 6.0 (Dennington et al., 2016) and their fully optimized geometries are presented in Fig. 7.

3.6. Frontier molecular orbital (FMO) analysis

The FMO’s of all compounds calculated at B3LYP/6-31G (d, p) DFT level are given in Fig. 8 along with values in Table 2. The values of energy gaps for compounds I and IV are very close with a minor difference (5.050, 4.668 eV) as shown in Table 2. The compound IV shows a smaller \( \Delta E \) value of 4.668 eV due to the greater \( \pi \)-conjugation than the I compound. Contrarily, the highest \( \Delta E \) value of 5.050 eV is seen in I. Thus, the HOMO-LUMO energy gap shows the following...
increasing order for all compounds: IV < I. The graphical diagrams are shown in Fig. 8 where red and yellow colors indicate the negative and positive phases of molecular orbitals, respectively.

3.7. Global reactivity descriptors

The ionization potential and electron affinity values in Table 3 illustrate that compound, I, show greater ionization potential than electron affinity revealing greater electron donating ability in comparison to the accepting nature. Overall, the electron affinity values are positive for both the compounds and a good gesture for their use in charge transfer reactions. The global electrophilicity values reveal that the electron donor capability (ω-) of these compounds are larger than the electron accepting capability (ω+). Similarly, the compound IV shows greater softness value followed by I with a minor difference. The electronegativity decreasing order observed among I and IV compounds is as IV > I. The synthesized compounds are harder

Fig. 3 Hirshfeld surface mapped of d_i, d_e, d_{norm}, curvedness, shape index, and fragment patch of compounds I and IV.
due to large values of chemical hardness than the lower values of global softness.

These findings reveal a high degree of chemical stability and little reactivity. With the increase in the chemical potential ($\mu$), chemical stability increases while reactivity decreases. The decrease in chemical potential for both the compounds is in the order: $[\text{IV} (\mu = -3.552) > \text{I} (\mu = -3.548)]$. This order indicates the lowest chemical potential ($\mu$) for the compound I making it kinetically less stable and more reactive. Similarly, the molecules with smaller softness are less stable and more reactive.

Fig. 4 2D fingerprint plots of I and IV displaying contacts for specific pairs of atoms contributing majorly.
The results of the global reactivity parameters show both compounds have a higher donating potential, more stability, and a poorer accepting ability and thus are appropriate to participate in charge transfer reactions. Moreover, the purpose of performing DFT computation of these compounds is to endorse experimental studies, cross-check experimental findings, as well as to evaluate stability, reactivity, intramolecular or intermolecular charge transfer, and other parameters are suitable for predicting potential photovoltaic characteristics (Pardasani et al., 2003; Pasha et al., 2007). Table 3 presents the global reactivity descriptors and accepting and donating aptitude of all compounds and endorses the relation with photovoltaic characteristics.

### 3.8. UV–visible study

To observe the electronic transitions, UV–vis spectroscopy is a powerful method, which also estimates the charge transfer probability in a compound and assignments of molecular orbitals corresponding to the transition (Siddique et al., 2021). Both, experimental and theoretical tools are used to study the UV–Visible transitions. The time-dependent DFT (TD-DFT) is performed at B3LYP functional in conjunction with the 6-31G(d,p) basis set. Table 4 represents the absorption maxima, oscillator strength ($f$), and molecular assignments. Both experimental and theoretical absorption values are close to each other with minute differences. The compound I shows maximum absorption at 267.3 nm with an oscillator strength of 0.217 which is nearly equal to the experimental $\lambda_{max}$ of 267.4 nm. Compounds IV shows absorption maxima at 247.8 nm bearing a very small difference with oscillator strengths of 0.210. Overall, the computed UV–vis studies are in fine contrast with experimental absorption values. The HOMO to LUMO transition is the characteristic of photovoltaic materials, ΔE gap and chemical hardness indicate molecules are a little bit harder requiring
a large amount of energy for reactivity which is an indication of kinetic stability. The biological charge transfer and photovoltaic capability of C1 and C2 are in the order IV > I making them potential candidates.

### 3.9. Molecular electrostatic potential (MEP) analysis

The 3D representation of electron density on all compounds is explored via MEP analysis following equation (1).

\[ V(r) = \sum_A \frac{Z_A}{R_A - r} \int \frac{\rho(r') dr'}{r' - r} \]  

(1)

Here, \( V(r) \) is molecular electrostatic potential, \( Z_A \) is charge density over the nucleus, \( A \) placed at \( R_A \), \( \rho(r') \) defines the electronic density function, and \( r' \) is the integration variable (Muthu and Prabhakaran, 2014; Okulik and Jubert, 2005).

The blue and red colors are the indications of nucleophilic and electrophilic centers, respectively. The magnitude of electrostatic potential is increased in order of
red < white < blue (Mahalakshmi and Balachandran, 2015). The electrophilic attack is easy on the red site while the blue site facilitates the nucleophilic attack. The 3D-MEP analysis is performed at B3LYP/6-31G(d,p) functional of all synthesized compounds and pictorial representation is given in Fig. 9. The oxygen, nitrogen, and sulfur atoms show negative potential (blue color), while mostly hydrogen and partially carbon and bromine atoms exhibit positive potential (red color). The white color indicates the mean potential being between two boundaries. The red and blue colors represent the different reaction sites.

### 3.10. NBO analysis

The charge transfer interactions within the bonds are studied using the natural bond orbital (NBO) analysis. The estimate of the off-diagonal NBO Fock matrix elements $F(i,j)$ is given by this interaction energy. This could be calculated using the second-order perturbation method (Foster and Weinhold, 1980; Glendening et al., 1998). The one-center lone pairs and two-center bonds from NBO analysis show a precise representation of chemical bonding for a stable molecular species, which corresponds to a single Lewis structure. The non-Lewis set involves unoccupied valence nonbonding (LP*) and

| Compounds | Ionization Potential (eV) | Electron Affinity (eV) | Electronegativity (eV) | Hardness (eV) | Chemical Potential (eV) | Electrophilicity $\epsilon^{-}$ donor capability (eV) | $\epsilon^{-}$ acceptor capability (eV) | Softness (eV) |
|-----------|---------------------------|------------------------|-----------------------|--------------|-------------------------|-----------------------------------------------|-----------------------------------------------|-------------|
| I         | 7.674                     | -0.578                 | 3.548                 | 4.126        | -3.548                  | 1.526                                         | 3.815                                         | 0.267       | 0.121        |
| IV        | 7.592                     | -0.488                 | 3.552                 | 4.040        | -3.552                  | 1.561                                         | 3.842                                         | 0.290       | 0.124        |
Table 4  Absorption values ($\lambda_{\text{max}}$), oscillator strength ($f$), and molecular assignments are obtained at TD-DFT/B3LYP/6-31G (d,p) level of DFT in chloroform solvent.

| DFT | Exp. $\lambda_{\text{max}}$ (nm) | f (oscillator strength) | MO contributions |
|-----|-------------------|------------------|-----------------|
|     |                   |                  |                 |
| 267.3 | 267.4              | 0.217            | H-1 $\rightarrow$ L (86 %), H-2 $\rightarrow$ L (4 %), |
| 247.8 | 248.9              | 0.210            | H-1 $\rightarrow$ L + 1 (3 %), H $\rightarrow$ L (2 %), |
|      |                   |                  | H-2 $\rightarrow$ L + 1 (53 %), H-1 $\rightarrow$ L + 2 (12 %), H $\rightarrow$ L + 2 (27 %), |

Extra-valence-shell Rydberg (RY*) orbitals also with the valence anti-bonds (BD*). The shortfall of the Lewis-type NBOs (bonds and lone pairs) denoting the density matrix can be computed with the occupancy of these NBOs. The delocalization energy, $E_{ij}$, is calculated as:

$$E_{ij}(2) = E_{ij} = qi \frac{F(i,j)^2}{(e_i^2 - e_j^2)}$$

Where $E_{ij}$ is the energy of hyper conjugative interactions, $qi$ is the occupancy of the contributing (Lewis type) orbital, $e_i$ and $e_j$ are the energies of the donating and accepting orbitals, and $F_{ij}$ is the off-diagonal element of the Fock matrix in the NBO basis (Frisch et al., 2000; Kleinman, 1962). An NBO analysis has been carried out to define the interior of the intramolecular hybridization and delocalization of electron density within the molecule.

Intermolecular hyper conjugative interactions are formed by overlapping the orbitals between the bonds of I (C–H), (C–O), (N–H), (C–C) and IV (C–H), (C–O), (C–N), (C–S), (C–C) and anti-bonding of I (C–C), (C–O), (C–Br), (N–C) and IV (C–C), (C–S), (C–N), (C–O), (N–C), (C–H) orbitals, which result in intramolecular charge transfer (ICT) stabilizing the molecular system and leading to strong delocalization. The strong intramolecular hyper conjugative interactions of the electrons of I (C–H), (C–O), (N–H), (C–C), and IV (C–H), (C–O), (C–N), (C–S), (C–C) and anti-bonding of I (C–C), (C–O), (C–N), (C–Br), (N–C) and IV (C–C), (C–S), (C–N), (C–O), (N–C), (C–H) bonds of the ring lead to stabilization of some part of the ring as given in Table 5. For example, in I, the intermolecular hyper conjugative interaction of σ (N2–H22) distributes to π* (C1–N6) and in IV σ (C15–S16) distributes to σ* (C15–S16), leading to stabilization of 21.129 KJ/mol and 10.91 KJ/mol, respectively.

This enhanced further conjugation with an anti-bonding orbital of I π (C4–C5) and IV π (C4–C5) which led to a strong delocalization of 98.742 KJ/mol and 23.24 KJ/mol with π* (C7–O8) in C1 and π* (C1–C6) in IV, respectively. The most important highest energy, related to molecule I, is electron-donating from π* (C13–C14) to the anti-bonding acceptor π* (C12–C17) and for C2, electron-donating from π* (C1–C6) to the anti-bonding acceptor π* (C2–C3) with stabilization energies of 845.796 and 204.94 KJ/mol, respectively. The magnitude of charge transferred in C1 from LP (2) O9 to the anti-bonding π* (C7–O8) and LP (1) N2 to the anti-bonding π* (C1–O10) shows that the stabilization energy is 6.38 KJ/mol and 10.89 KJ/mol, respectively. Similarly, in C2, the charge transferred from LP (1) N14 to the anti-bonding σ* (C15–S16) and LP (2) O8 to the anti-bonding π* (C4–C5) with stabilization energies of 52.70 and 28.53 KJ/mol, respectively.

3.11. Optical properties

Nonlinear optical materials may find use in optoelectronic devices, networking, signal manipulation, telecommunications, and other fields. The strength of the donor–acceptor group, as well as the sequence in which they are placed, affects the amount of nonlinear optical efficiency. To put it another way, organic NLO materials have a push–pull conjugated structure because of the π-electron conjugated system they contain, bearing electron donor and acceptor groups on different ends. The present work has been extended to explore the optical properties of the I and IV complexes for their remarkable utilization in optoelectronic. The optoelectronics properties include dipole moment ($\mu_{\text{tot}}$), polarizability ($\alpha_{\text{tot}}$), and first-order hyperpolarizability ($\beta_{\text{tot}}$) (Kurtz et al., 1990) and are determined by the following equations:

![Fig. 9](image-url)  The 3D MEP representation of I and IV at B3LYP/6-31G(d,p) level of DFT.
The greater hyperpolarizability of the molecule is optically active in the X-direction. The perpendicular components contribute minimally, indicating that the molecule is optically active in the X-direction. The results clearly show that the largest component of hyperpolarizability is mainly attributed to the greater intramolecular charge transfer while compound I formed hydrogen bonds with Asn142, Ser144, and Gly143. Similarly, compound IV formed hydrogen bonds with Phe 140, Gly143, and Cys145. The binding mode and chemical interactions of the compounds with the enzyme are given in Fig. 10.

### 3.12. Molecular docking study

The docking was conducted to examine the binding affinity of compounds with different targets of the SARS-CoV-2 virus. Of the different targets used, it was found the main protease enzyme was the best-docked enzyme for the compounds used, as shown in Table 7. Compound I had binding energy of $-12.67$ kcal/mol for the SARS-CoV-2 main protease enzyme, while compound IV had binding energy of $-11.24$ kcal/mol. Both the compounds dock well inside the binding cavity of the enzyme and are engaged by hydrophilic and hydrophobic interactions. The compound I formed hydrogen bonds with Asn142, Ser144, and Gly143. Similarly, compound IV formed hydrogen bonds with Phe 140, Gly143, and Cys145. The binding mode and chemical interactions of the compounds with the enzyme are given in Fig. 10.

### 3.13. Molecular dynamics simulation

As the compounds showed stronger binding affinity with the SARS-CoV-2 main protease enzyme compared to others, only main protease enzyme compounds best docked complexes were subjected to molecular dynamics simulation analysis. According to the molecular dynamic simulations, compound I is more dynamically stable compared to compound IV. The
The root mean square deviation (RMSD) reached a maximum level of 2 Å. The compound IV, on the other hand, showed significant deviation (up to 7 Å) in the first 100 ns. Followed that, the complex was found very stable towards the end of the simulation. The residue-wise fluctuations were measured via the root mean square fluctuation (RMSF), which indicated that in the presence of compound I, the enzyme remained very stable compared to compound IV. The RMSD and RMSF of compounds are given in Fig. 11.

Table 6  Dipole moment (Debye), polarizability (au), and nonlinear first order hyperpolarizability (au) of C1 and C2 complexes.

| Parameter   | Value | Parameter   | Value |
|-------------|-------|-------------|-------|
| $\mu_x$     | 1.543 | $\beta_{xxx}$ | 15.49 |
| $\mu_y$     | 0.000 | $\beta_{xyy}$ | 29.25 |
| $\mu_z$     | 0.000 | $\beta_{zzz}$ | 479.7 |
| $\mu_{tot}$ | 1.543 | $\beta_{xyz}$ | 50.63 |
| Polarizability (au) | | | |
| $\alpha_{xx}$ | 191.6 | $\beta_{xxx}$ | 15.49 |
| $\alpha_{xy}$ | 7.690 | $\beta_{xyy}$ | 29.25 |
| $\alpha_{xz}$ | -15.34 | $\beta_{xzz}$ | 479.7 |
| $\alpha_{yz}$ | 202.2 | $\beta_{xyz}$ | 50.63 |
| $\alpha_{zz}$ | -8.241 | $\beta_{zzz}$ | 15.49 |
| $\alpha_{tot}$ | 170.7 | $\beta_{xyz}$ | 29.25 |

Table 7  Docking score of compounds with different SARS-CoV-2 targets. The energy value is in kcal/mol.

| Compound | Main Protease | RdRp | E Protein |
|----------|---------------|------|-----------|
| I        | -12.67        | -6.35 | -7.65     |
| Compound IV | -11.24        | -5.65 | -6.65     |

Fig. 10  Binding of the compounds at substrate binding cavity of SARS-CoV-2 main protease enzyme. Compound I is shown by a red stick while compound IV is pink. The enzyme is on hydrophobicity surface. The binding interactions of compounds are also given.
3.14. MMPB/GBSA analysis

Different binding free energies of the compounds with the enzyme active site residues were also estimated. Again, compound I was ranked as the best-docked molecule with the main protease enzyme. The binding free energies of compounds are tabulated in Table 8. The net binding free energy of compound I is $-59$ kcal/mol in MMGBSA and $-59.553$ kcal/mol in MMPBSA. The energy value of compound I is better than that of compound IV, which has a net energy of $-46.69$ kcal/mol in MMGBSA and net energy of $-41.03$ kcal/mol in MMPBSA.

4. Conclusions

In this paper, we carried out several computation analyses to treat test dihydropyrimidines due to the possible benefits that can hit today’s prominent pandemic problem.

Furthermore, from the DFT theoretical calculations, the optimized geometrical parameters are analyzed and are in good agreement with the XRD results. We found that the HOMO-LUMO energy gap affects the stability of compounds. Similarly, NBO and natural atomic charge analyses were performed to explore the stabilization energy of different inter- and intramolecular interactions within the systems and the net charge distribution, respectively. Insights from the DFT study of these compounds reveal their good stability by using other analyses like global indices of reactivity and molecular electrostatic potential analysis. Molecular docking and MD simulations were used to assess the binding affinity and stability of these drugs to the binding site of the SARS-CoV-2 receptor. Through molecular docking studies with various types of interactions, substantial binding affinity for both examined compounds was established. Following the computational techniques, the substrates were subjected to MD simulations to measure the rigidity of the protein structure, fluctuations caused by interactions, and overall structural stability by calculating RMSD.

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.

Appendix A. Supplementary material

Supplementary data to this article can be found online at https://doi.org/10.1016/j.arabjc.2022.104230.

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