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New α-Hydrazinophosphonic acid: Synthesis, characterization, DFT study and in silico prediction of its potential inhibition of SARS-CoV-2 main protease

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A new α-Hydrazinophosphonic acid (HDZPA) has been synthesized and its molecular structure was determined using spectroscopic methods. The Density Functional Theory (DFT) at the B3LYP/6–31 G (d,p) level was utilized to determine the electronic properties, vibrational modes and active sites of the examined molecule. In this context, some quantum chemical parameters have been calculated in order to discuss the reactivity of the studied molecule. Also, the inhibition activity of the investigated α-Hydrazinophosphonic acid for SARS-CoV-2 main protease (M pro ) and RNA dependent RNA polymerase (RdRp) has been predicted using in silico docking.

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1. Introduction

The COVID-19 is an abbreviation of the Coronavirus disease 2019. At first, the COVID-19 is started as an epidemic in the Chinese city of Wuhan in December 2019, afterward it quickly propagated in Chinese territory and outside [1,2]. Two months later, the WHO declares the COVID-19 as a pandemic [3]. In general, the principal diseases caused by COVID-2019 are respiratory problems and gastrointestinal [4].

Presently, no known special, confirmed and efficient antiviral drug is discovered or developed. However, several drugs are also tested in clinical trials as effective treatments of COVID-19 patients such as the old antimalarial drug Hydroxychloroquine and the new antiviral drug developed for Ebola virus Remdesivir. In this context, a recent published paper reports that the treatment including Remdesivir and Chloroquine drugs inhibits the growth process of SARS-CoV-2 in vitro [5]. Furthermore, Remdesivir has proved an important efficacy against SARS-CoV-2 whether in vitro or in vivo and has also started its clinical experiments [6]. Also, several clinical tests realized in China show that Chloroquine presents a remarkable effect on the clinical results and viral clearance [7,8]. Another recent research suggests the use of Hydroxychloroquine as a potential treatment of COVID-19 patients [9]. In addition, the paper published by Didier Raoult et al. on March 2020 showed that the treatment containing Hydroxychloroquine and Azithromycin presents a significant reduction/disappearance of the viral load in COVID-19 patients [10].

The crystalline structure of SARS-CoV-2 main protease (M pro ) has been established at the first time by Liu et al., and the structure of M pro is available for public access in the Protein Data Bank (PDB) [11]. Generally, this enzyme is known by its vital role in the transformation of the translated polyproteins [12]. On the other hand, RNA-dependent RNA polymerase (RdRp) plays a pivotal role in virus replication and transcription of the viral genome [13]. So, M pro and RdRp can be considered as targets to discover therapeutic agents to COVID-19.

In general, the physicochemical and electronic characteristics of drug molecules can affect their chemical and biological activities. Additionally, the quantum chemical calculations using Density Functional Theory (DFT) method are largely employed to determine the active sites of drugs and to correlate their activity with various quantum chemical parameters [14–16]. In this context, several
biological activities of the bioactive molecules can be extensively studied using the DFT calculations [17-19].

The one-pot multicomponent reactions such as Mannich-type reaction and Kabbachnik-Fields reaction are largely and efficiently used in organic synthesis to prepare the biologically active compounds [20-24]. Several bioactive and pharmaceutical compounds have been synthesized using one-pot multicomponent reactions including antiviral [25], antitumor [26], anti-HIV [27], antimalarial and antiinsecticidal [28], antipsyhotic [29,30], antiparasitic [31], antidepressant [32], antibacterial and antimicrobial [33]. The Kabbachnik-Fields reaction is a three-component reaction including a carbonyl, an amine and a dialkylephosphite or a trialkylephosphite. This reaction is very important in drug discovery studies and mainly used to synthesize α-amino phosphonates.

This paper represents the results of an experimental and theoretical study of a new α-Hydrazinophosphonic acid (HDZPA). So, the experimental part consists to synthesize the desired compound via one-pot three-component reactions and to determine its molecular structure using spectroscopic methods such as UV-VIS, FT-IR, 1H NMR, 13C NMR, and 31P NMR. On the other hand, the HDZPA has been theoretically investigated to locate their active sites, electronic and vibrational properties by means of DFT method at the B3LYP/6–31G (d,p) level. The optimized molecular structures, the vibrational spectra, the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) properties, dipole moments (μ), Molecular Electrostatic Potentials maps (MEP), atomic charges, energy gap (ΔE Gap), hardness (η), local softness (σ), electronegativity (χ) and electrophilicity (ω) are calculated for of the studied molecule. Finally, the synthesized α-Hydrazinophosphonic acid was tested as a potential inhibitor for SARS-CoV-2 main protease (Mpro) and RNA dependent RNA polymerase (RdRp) using in silico docking to support drug discovery.

2. Experimental

2.1. Materials and spectroscopic details

All the chemical reagents employed for the HDZPA ligand synthesis were bought from Sigma-Aldrich and utilized without further purification. The UV–VIS spectrum of the synthesized HDZPA ligand was obtained in the range of 190–900 nm in aqueous solution by means of the Jasco V-650 UV–VIS spectrometer. In addition, the JASCO 4000 FTIR spectrometer was used to realize the FT-IR spectrum of the investigated ligand in solid state at room temperature, and the obtained vibration frequencies were listed in the range of 600–4000 cm⁻¹. NMR spectra were recorded on a Bruker Avance 300 spectrophotometer operating at 300 MHz (1H) and 75 MHz (13C) at 298 K using tetramethylsilane (0 ppm) as the internal reference. NMR spectroscopic data were recorded in CDCl₃ (δ = 7.26 ppm) using as internal standards the residual non-deuterated signal for 1H NMR and the deuterated solvent signal (δ = 7.71 ppm) for 13C NMR spectroscopy. Chemical shifts (δ) are given in ppm and coupling constants (J) are given in Hz. The following abbreviations are used for multiplicities: s = singlet, d = doublet, t = triplet, q = quartet, dd = doublet of doublets, and m = multiplet.

2.2. Synthesis of the α-Hydrazinophosphonic acid ligand

In this study, a new α-Hydrazinophosphonic acid ligand (HDZPA) was synthesized applying the Irani-Moeditzer method [34]. Accordingly, a mixture of 1.0 mmol of (4-methylphenyl)hydrazine and 2.0 mmol of H₃PO₃ has been dissolved in 50 ml of water and 25 ml of HCl, then the mixture was refluxed for 4 h at 100 °C (Fig. 1), whereas 2.0 mmol of formaldehyde 36% was added dropwise to this mixture. After the addition, the mixture was kept at the room temperature for another 3 h under reflux. Also, TLC analysis was used to check the completion of the reaction. After the reaction was completed, a rotary evaporator has been used to eliminate the solvent under reduced pressure. Finally, the obtained crude product was purified by means of chromatographic column of silica gel using ethyl acetate/MeOH (9:5:0.5, v/v). Moreover, the obtained brown solid (HDZPA) was characterized by UV–Vis, FT-IR, 1H NMR, 13C NMR, and 31P NMR.

Yield 81%, M.p. 79.22 °C; IR (ATR, ν(cm⁻¹)) : 3621 (O=H), 3315 (N–H), 3250 (C–H₆), 2980 (C–H₆), 2911 (P=O), 1629 (N=C=O), 1202 (P = O), 1168 (N–N), 1020 (N=C=O), 868 (C-P); 1H NMR (300 MHz, CDCl₃, δ(ppm)): 2.48 (s, 3H, -CH₃), 3.12 (d, 2J H,P = 11.8 Hz, 4H, -CH₂–), 5.08 (s, 4H, -OH), 7.18 (d, J = 7.9 Hz, 2H, H-Ar), 7.71 (d, J = 7.9 Hz, 2H, H-Ar), 8.92 (s, 1H, NH); 13C{1H} NMR: (75 MHz, CDCl₃, δ(ppm)): 23.9 (1C, -CH₃), 67.7 (2C, d, 1J C,P = 36.8 Hz, N–CH₂–), 114.1 (2C, CH₆), 129.9 (2C, CH₆), 133.3 (1C, CH₂–C₆H₅), 141.8 (1C, N–C₆H₅); 31P NMR: (121 MHz, CDCl₃, δ(ppm)): 8.77 (dt, 1J P,C = 87.1 Hz, 2J P,H = 116.4 Hz); UV–Vis (H₂O), λmax (nm): 244.76 and 327.25. MS (70 eV) m/z (%): 311 (M + 1, 7.51), 310 (M⁺, 100), 309 (M–1, 4.2) (see Supporting Information).

2.3. Computational details

Throughout this study, we applied the Gaussian 09 W program package to implement all quantum chemical calculations [35]. The geometry of HDZPA ligand was entirely optimized utilizing the DFT method through B3LYP hybrid functional at 6–31G (d,p) basis set [36,37]. Also, this theory has been utilized to determine the vibrational frequencies at the obtained optimized structure of HDZPA. We chose Density Functional Theory (DFT) with B3LYP/6–31G (d, p) level because the B3LYP functional has shown good results for organic molecules. Also, the obtained results with B3LYP/6–31G (d,p) level are in good agreement with the experimental data, especially with ATR-FTIR and UV–VIS data. The 1H, 13C and 31P NMR spectra of HDZPA in the presence of CDCl₃ as solvent are predicted using the GIAO method with the hybrid B3LYP at 6–31G (d,p) basis set, while the electronic spectrum in the water was predicted by using the Time-dependent DFT calculations (TD-DFT) with the B3LYP/6–31G (d,p) method. The VEDA 4 program has been used to carry out various vibrational parameters and PED calculations [38]. In addition, the calculated energies of the highest occupied molecular orbital (E₉(HOMO)) and lowest unoccupied molecular orbital (E₉(LUMO)) have been utilized to determine various quantum chemical parameters such as the energy gap (ΔE GAP), dipole moments (μ), hardness (η), local softness (σ), electronegativity (χ) and electrophilicity (ω). Moreover, the following equations were applied to compute the precedent parameters [39,40]:

\[ \Delta E_{\text{GAP}} = E_{\text{LUMO}} - E_{\text{HOMO}} \]  
\[ \eta = \frac{E_{\text{LUMO}} - E_{\text{HOMO}}}{2} \]  
\[ \sigma = \frac{1}{\eta} \]
3. Results and discussion

3.1. Spectral study

3.1.1. UV–vis analysis

The experimental and theoretical UV–vis spectra of HDZPA (Fig. 3) were obtained in water at room temperature and they reveal two absorption bands in the ultraviolet domain. The registered experimental spectrum of the investigated ligand illustrates a band at 244.76 nm related to the $\pi \rightarrow \pi^*$ transitions of the C = C group of the aromatic ring, which indicates that the electron orbital jumps from $\pi$ bonding orbitals to $\pi$ anti-bonding orbitals. Also, the band appeared at 327.25 nm is assigned to the $n \rightarrow \pi^*$ transitions related to the presence of heteroatoms (O, N and P) on the molecular structure of HDZPA, which indicates that the electron orbital jumps is from non-bonding orbitals to $\pi$ anti-bonding orbitals. In an atom or molecule, this type of transitions shows that electrons move from low energy levels to higher energy levels. Fig. 6 indicates that the experimental and calculated absorption spectra are in good agreement. Also, the calculated spectra shows a band at 256.07 nm associated to the $\pi \rightarrow \pi^*$ transitions. In addition, the calculated band observed at 320.07 nm is related to the $n \rightarrow \pi^*$ transitions of heteroatoms.

3.1.2. Vibrational analysis

The vibrational frequencies, the vibrational mode assignments and the potential energy distributions (PED) of the characteristic groups of HDZPA are presented in Table 1. These results were selected from the obtained experimental and calculated infrared spectra. Also, the calculated frequencies are scaled using a scaling factor of 0.9614 [41]. As a result, a good harmony has been observed between the scaling calculated frequencies and the experimental frequencies (Fig. 5). Generally, the objective of the vibrational analysis is to specify vibrational modes of a molecule. At B3LYP/6–31 G (d,p), the studied compound with 35 atoms donates (3N-6) i.e. 99 vibrational modes in the range 20.02–4126.00 cm$^{-1}$. So, the attribution of these 99 vibrational modes has been performed to establish some correlation between structure and spectrum. These 99 vibrational modes distinguish 34 stretching modes, 33 bending modes and 32 torsional modes. All frequencies below 600 cm$^{-1}$ are not presented in Table 1 owing to their very complex mode compositions. Therefore, the analysis of the IR spectra presented in Fig. 4 and Table 1 confirms the existence of the following vibrational modes:

3.2.1. Phosphonic acid group (-PO(OH)$_2$) vibrations

The phosphonic acid group (-PO(OH)$_2$) presents several vibrational modes such as $\nu(P = O)$, $\nu(-OH)$, $\nu(P-O)$, and $\nu(C-P)$. Generally, the vibrational modes related with this group are observed as medium to strong absorption peaks in IR spectra. The peak which is highest in wavenumber appeared at 3620 cm$^{-1}$ is attributed to the O–H stretching vibrations. Also, the $P = O$ stretching vibrations is observed at 1202 cm$^{-1}$, while the intense peak appeared at 1611 cm$^{-1}$ is related to the P–O stretching vibrations. The stretching vibrations of C–P are located at 684 cm$^{-1}$. Also, the presence of hydrogen bonds in the molecular structure of HDZPA can expanded the characteristic peaks of $\nu(P = O)$ and $\nu(P–O)$ and there corresponding frequencies can be down-shifted below expected regions.
3.2.2. N–H stretching vibrations

In general, the N–H stretching vibrations are facely determined by the apparition of a sharp peak at highest wavenumbers. For the investigated molecule, the asymmetric and symmetric stretching vibrations of N–H are located at 3315 cm⁻¹. Also, the N–H out of plane bending vibrations are situated at 621 cm⁻¹. The N–H bending vibrations of the secondary amine (N–H wagging) are observed between 652 and 910 cm⁻¹.

3.2.3. C–H vibrations

The molecular structure of HDZPA indicates the presence of the aromatic and aliphatic C–H stretching vibrations. So, the obtained peak between 3005 and 3250 cm⁻¹ can be attributed to the aromatic C–H asymmetric and symmetric stretching vibrations. Also, the asymmetric and symmetric stretching vibrations of the aliphatic C–H (CH₂ and CH₃ groups) are obtained at 2980 and 2848 cm⁻¹, respectively. Also, the out of plane and in plane bending vibrations of the aromatic C–H have been specified and presented in Table 1. In addition, all peaks appeared between 1400 and 1250 cm⁻¹ are assigned to the deformation modes of C–H. In experimental spectra, the CH₂ scissoring is attributed to 1495–1445 cm⁻¹, while the CH₂ wagging is assigned to 1358–1322 cm⁻¹.

![Experimental and calculated IR spectra of HDZPA.](image)

![Correlation diagram between the calculated and experimental wavenumbers of HDZPA.](image)

### Table 1

| Assignment (PED) | Experimental | Theoretical |
|------------------|--------------|-------------|
| ν O–H (98)       | 3621 (ν O–H) | 3788 3641   |
| ν N–H (100)      | 3315 (ν N–H) | 3425 3293   |
| ν C–Hα (96)      | 3250 (C–Hα) | 3353 3223   |
| ν C–Hγ (97)      | 3314 (C–Hγ) | 3142 3020   |
| ν C–Hβ (94)      | 3298 (C–Hβ) | 3117 2996   |
| ν C–Hγ (84)      | 2884 (ν C–Hγ) | 3108 2988   |
| ν C–Hβ (85)      | 1611 (ν P–OH) | 1674 1609   |
| ν C–H (25)       | 1411 (ν C–H) | 1459 1402   |
| α C–H (34)       | 1378 (α C–H) | 1393 1342   |
| ω C–H (39)       | 1351 (ω C–H) | 1382 1328   |
| ν N–Cα (52)      | 1269 (ν N–Cα) | 1283 1234   |
| ν P = O (33)     | 1202 (ν P = O) | 1242 1194   |
| ν N–N (22)       | 1168 (ν N–N) | 1186 1140   |
| ν C–H (24)       | 1084 (ν C–H) | 1125 1081   |
| ν N–Cβ (34)      | 1041 (ρ P–O) | 1066 1024   |
| ν N–CH₂ (3)      | 1020 (ν N–Cβ) | 1052 1011   |
| δ C–H (41)       | 748 (δ P–C) | 772 742     |
| δ C–H (17)       | 684 (δ P–C) | 714 686     |

ν: stretching, γ: rocking, α: scissoring, δ: twisting, ω: wagging, β: in plane bending, ρ: torsion, ρout: out plane bending, s: symmetric, as: asymmetric.
3.2.5. C–N vibrations

The attributed peaks of the aromatic and aliphatic C–N stretching vibrations of the studied ligand are situated at 1269 cm\(^{-1}\) and 1020 cm\(^{-1}\), respectively. The thin peak located at 822 cm\(^{-1}\) may be due to C–N band vibration. Also, the calculated in plane bending vibrations of C–N are observed at 468 and 497 cm\(^{-1}\).

3.2.7. N–N vibrations

The characteristic peak of the stretching vibrations of N–N group is appeared with a medium intensity at 1168 cm\(^{-1}\). Also, the peak situated at 791 cm\(^{-1}\) is associated to the symmetric wag. Moreover, the peak situated at 885 cm\(^{-1}\) may be assigned to the in-plane bending vibrations of C–N–N.

3.2.8. C–C and c = c vibrations

Generally, the C–C stretching modes of aromatic rings are appeared between 1400 and 1650 cm\(^{-1}\). Consequently, for the studied compound, the C–C stretching vibrations are observed at 1520 and 1742 cm\(^{-1}\) in the experimental IR spectrum. Also, the aromatic C = C semicircular stretching vibrations are appeared between 1625 and 1430 cm\(^{-1}\).

3.1.3. NMR analysis

The analysis of the experimental \(^1\)H NMR spectrum of HDZPA permits to determine the following characteristic signals: a singular signal located at 2.48 ppm is related to the three protons of \(-\text{CH}_3\) group. The 4H of the two aliphatic \(-\text{CH}_2\) – groups are appeared as a doublets signal at 3.12 ppm and this can be proved by the obtained values of integration (3.9 = 4H) and coupling constant \((J = 11.8\) Hz\) of the correspondent signal. Also, the 4H of the phosphonic acid groups \((–\text{OH})\) are observed as a singular signal at 5.08 ppm. Moreover, the observed signals in the region of 7.05–7.15 ppm are attributed to the aromatic protons \((–\text{CH}_2\text{Ar}, 4\text{H})\).

The proton of the \(-\text{NH}\) group is appeared as a singular signal at 8.92 ppm.

Furthermore, the elucidation of the obtained experimental \(^{13}\)C NMR spectrum of HDZPA shows the presence of the following characteristic signals: the carbon of the \(-\text{CH}_3\) group was observed as a singular signal at 23.89 ppm. The two signals situated at 67.70 ppm and 68.19 ppm can be associated to the carbons of the two \(\text{N}–\text{CH}_2\) – groups. The carbons of the aromatic ring are observed between 114.14 ppm and 133.37 ppm.

The experimental \(^31\)P NMR spectrum of HDZPA shows two signals at 8.42 ppm and 9.13 ppm, which confirms the presence of the phosphorus atoms in the molecular structure of ligand. Also, the obtained signals of the phosphorus atoms are revealed as a triplet due to their coupling with the two protons of the methylene group \((\text{N}–\text{CH}_2–\text{P})\). Indeed, in the case where the molecule contains a phosphorus atom, then this one couples with the nuclei of spin \(\frac{1}{2}\). The NMR signals are therefore duplicated. Thus we will observe coupling constants of the 3 J and 4 J type.

Thus, for HDZPA all the carbons and protons shown will couple with phosphorus. The most striking example is that of carbons of methylene groups \((\text{N}–\text{CH}_2–\text{P})\) for which there will be a coupling constant \(J = 13.9\) Hz.

On the other hand, the results presented in Table 2 show that the calculated NMR chemical shifts for HDZPA are in agreement with the experimental chemical shifts. On the other hand, we observe better correlations are obtained for the H and P atoms in solvent than the C atoms. Also, we observe in the \(^1\)H NMR results that the theoretical values were larger than those obtained in the experimental measurements.

3.2. DFT study

3.2.1. Optimized molecular structures

In Fig. 2(a) we present the obtained optimized molecular geometry of HDZPA ligand. Additionally, the calculated value of total energy for this molecule at the optimal structure was \(-43,437,641\) eV, which signifies the energy of the more stable conformation of the studied ligand.

3.2.2. Frontier molecular orbital analysis

Generally, the reactivity and the stability of drug molecules can be described significantly using the HOMO and LUMO orbitals and their energies. Also, the aptitude of a molecule to contribute electrons to an electrophilic species can be examined by HOMO, while the ability of a molecule to accept electrons from nucleophilic species may be determined using LUMO [42]. So, the capacity of a molecule to give electrons to an acceptor species is favored by the high values of \(E_{\text{HOMO}}\) [43], whereas the ability of a molecule to take electrons is preferred by the low values of \(E_{\text{LUMO}}\) [44]. On the other hand, the energy gap \((\Delta E_{\text{gap}})\) of drug molecules is an important parameter to determine their reactivity and stability [45]. Additionally, the calculated value of \(\Delta E_{\text{gap}}\) can explain the charge transport interactions in the molecule. In general, \(\Delta E_{\text{gap}}\) represents the necessary energy to excite the electrons of a molecule. Also, molecules can be highly chemically reactive, unstable and excited easily when \(\Delta E_{\text{gap}}\) is smaller, while it can be very stable and less chemically reactive if \(\Delta E_{\text{gap}}\) is very large [46].

Figs. 6(c) and 6(d) illustrate the calculated HOMO and LUMO orbitals for the HDZPA ligand. Generally, the green color characterizes the negative phases, whereas the red color indicates the positive phases [39]. So, we observe from Figs. 6(c) and 6(d) that the HOMO and LUMO are frequently located on the aromatic ring, the amino and methylene groups. On the outer hand, we observe from Table 3 that the HDZPA molecule has an elevated value of \(E_{\text{HOMO}}\) and \(E_{\text{LUMO}}\), which signifies that the HDZPA can liberate electrons to an acceptor molecule. Moreover, the calculated value of \(\Delta E_{\text{gap}}\) indicates the stability and the reactive of HDZPA.

3.2.3. Molecular surface electrostatic potential (MEP)

One of the helpful ways of quantum chemical calculations used to determine the active sites of drug molecules is the measure of
their molecular electrostatic potential (MEP) maps. In general, the MEP of molecules is associated to their electronic densities and can elucidate their chemical activities, electrostatic effects and partial charges. Also, MEP map is a visual method which can be used largely to identify the relative polarity of molecules and to establish their negative and positive electrostatic potentials [47]. Additionally, the total electrons density plotted with electrostatic potential surface can be used to characterize the charge density, the dimension and the form of the active sites and to locate the place of the chemical reactivity of molecules. So, the 3D maps presentation of the electrostatic potential variation is represented by a gradient of colors. In principle, the negative electrostatic potential regions related to the electrophilic reactions are graphically presented in yellow and red colors, while the positive electrostatic potentials corresponding to the nucleophilic reactions sites are illustrated in blue color. Moreover, the zones of nil potential are presented in green color and the evolution of the potential obeys the sequence red-orange-yellow-green-blue [48].

The calculated 3D MEP maps of HDZPA by means of DFT method are displayed in Fig. 6(b). The examination of the achieved MEP map elucidates that the yellow and red zones are situated on the O33, O32, O27, O29, O30, and O25, which proves that these atoms are the probable sites of the electrophilic reactions. Also, the aromatic ring is accounted as negative regions. Conversely, the green and blue colors are observed in the region of the carbon and hydrogen atoms, which represent the positive sites designated for the nucleophilic reactions.

### 3.2.4. Mulliken atomic charges

The estimation of the partial atomic charges of molecules can be provided by Mulliken charges analysis. Furthermore, the adsorptive sites of drug molecules can be proved by determination of their atomic Mulliken charges. In this context, the obtained values of the atomic Mulliken charges of HDZPA are regrouped in Table 4. The examination of the obtained results indicates that the oxygen and nitrogen atoms have the most negative charges, which is due to the molecular relaxation. As well, the hydrogen atoms cover the positive charges. In particular, the largest part of negative charges are localized on the O27, O25, O29, O30, O32, O33, N15 and N17 atoms of HDZPA, which are possibly the active centers of adsorption [49]. Generally, the heteroatoms (N and O) can share their pairs of electrons with acceptor molecules. In addition, the P24 and P35 atoms of HDZPA have the majority positive charges.

### 3.2.5. Dipole moment

The dipole moment (μ) can be used to evaluate the chemical reactivity of molecules. In general, the value μ designates the polarity of drug molecules, which is correlated to the fractional electric charge distribution in these molecules [50]. As well, the importance of μ illustrates in the mechanism of reactions and designates the ability of molecules to interact with other molecular species. In Table 3 we present the calculated value of μ for the examined ligand. The examination of this value indicates that the HDZPA ligand offer high abilities to interact with surrounding medium.
3.2.6. Global reactivity descriptors
The calculated values of \( E_{\text{HOMO}} \) and \( E_{\text{LUMO}} \) have been used to calculate the global reactivity parameters such as local softness (\( \sigma \)), hardness (\( \eta \)), electrophilicity (\( \omega \)) and electronegativity (\( \chi \)). Also, the relation between the stability of molecules and their global chemical reactivity can be determined using these parameters [51]. Generally, the chemical reactivity and stability of molecules can be evaluated measuring their hardness and local softness. In addition, the resistance of drug molecules against the deformation of their electron clouds or polarization can be measured by the hardness. Moreover, hard molecules are described by the elevated values of \( \Delta E_{\text{CAP}} \), whereas the soft molecules are expressed by the low values of \( \Delta E_{\text{CAP}} \). On the outer hand, the high gap energy value indicates that the molecule is more stable and low reactive, while the small gap energy value is related to high reactivity and low stability of the molecule. So, the \( \Delta E_{\text{CAP}} \) is an important parameter to determine the stability and the reactivity of molecules. From the presented values of \( \eta \) and \( \sigma \) in Table 3, we observe that HDZPA has an elevated value of \( \sigma \) and least value of \( \eta \), which indicates its high reactivity. Furthermore, the calculated value of \( \chi \) indicates that HDZPA is an electronegative species.

The electrophilicity of molecules estimate their abilities to take electrons. As well, the elevated values of \( \omega \) prove the better electrophility, whereas the low values of \( \omega \) signify a poor electrophile [52]. According to the electrophility value, we can classify the organic molecules into three categories: marginal electrophiles with \( \omega < 0.8 \) eV, moderate electrophiles with \( 0.8 < \omega < 1.5 \) eV and strong electrophiles with \( \omega > 1.5 \) eV [53]. From Table 3 we note that the examined ligand is a marginal electrophile with \( \omega < 0.8 \) eV.

We can observe clearly a good correlation between MEP analysis results and the calculated values of global reactivity descriptors.

3.3. Molecular docking analysis
The molecular docking between HDZPA ligand and the M\textsuperscript{pro} and RdRp receptors was executed to define the appropriate conformation of the HDZPA in the receptor and the secondary forces resulting between HDZPA and the active amino acids of the receptor. This leads to the development of new drug designs. Based on the minimum binding energy, the non-covalent bonds, \( \pi-\pi^* \) and \( \pi-\sigma \) interactions between the active amino acids of the M\textsuperscript{pro} and RdRp target receptors and the HDZPA ligand were tested.

Fig. 7 represents crystal structures of the best docked modes of M\textsuperscript{pro}-HDZPA and RdRp-HDZPA complexes. We can observe from Figs. 7(a) and 7(b) that the HDZPA ligand prefers to bind in the outer structure of M\textsuperscript{pro}, while it favors to bind in the inner pocket of RdRp.

Fig. 8 illustrates the detailed presentation of interactions of the investigated ligand with M\textsuperscript{pro} and RdRp. Generally, the HDZPA binds with M\textsuperscript{pro} and RdRp by means of various hydrogen and Van der Waals bonding interactions. For M\textsuperscript{pro}, we can observe from Fig. 8(a) that the HDZPA interacts through H-bonding with HIS163, GLY143, SER144 and LEU141 amino acids whereas bends with van der Waals forces with some amino acids such as GLN189, HIS41, HIS164, MET165, GLU166 and PHE140. Generally, the hydrophilic interactions included H-bonding whereas hydrophobic interactions implicated van der Waals forces. In case of RdRp, we can see from Fig. 8(b) that the HDZPA interacts with just one hydrogen bond with the A13 Nucleotide. Concerning van der Waals forces, HDZPA connects with GLY590, THR591, LEU758, TRP598, PHE812, MET601, CYS813, PHE594 and LYS593. The calculated values of the binding energy of HDZPA and some drugs have been presented in Table 5. Comparing the obtained values of binding energy of HDZPA with those of some drugs reported in the literature, we observe that our molecule presents the lowest values of binding energy in molecular interactions with M\textsuperscript{pro} and RdRp, which indicates that HDZPA has the better binding affinity and the M\textsuperscript{pro}-HDZPA and RdRp-HDZPA complexes are more stable than these formed for Chloroquine, Hydroxychloroquine and Remdesivir.

In conclusion, the in silico docking results revealed that HDZPA is expected as a potential compound to treat COVID-19. On the other hand, we recommend researchers to complete the in vitro and in vivo studies of HDZPA with novel corona virus in order to confirm the obtained docking results and to know the exact impact of the studied compound.

4. Conclusion
In this research, the \( \alpha \)-Hydrazinophosphonic acid has been synthesized in good yield using one-pot three-component reactions and characterized using spectroscopic methods. Also, the quantum chemical study of HDZPA ligand has been performed applying DFT method at B3LYP/6–31 G (d,p) basis set. In this context, the optimal structures, the vibrational spectra, the 3D MEP map, HOMO and LUMO orbitals, \( \Delta E_{\text{CAP}} \), dipole moments, Milliken atomic charges, hardness, local softness, electrophilicity and electronegativity have been calculated for the examined molecule. So, the essential findings are described below. Firstly, the experimental and theoretical IR spectra of the investigated molecule were completely analyzed and the vibrational modes have been attributed. The calculated values of \( E_{\text{HOMO}} \) and \( E_{\text{LUMO}} \) show that the HDZPA share their electrons to an acceptor species. Also, the obtained values of \( \Delta E_{\text{CAP}} \) indicate that the HDZPA is a reactive and instable species. The 3D MEP maps illustrate that the possibly sites of the electrophilic reactions are situated on the O33, O32, O27, O29, O30, and O25 atoms. According the Milliken charges results, the O27, O25, O29, O30, O32, O33, N15 and N17 atoms of HDZPA are possibly the active sites of interaction. The obtained values of the global reactivity descriptors demonstrate that the HDZPA ligand is a pow-
Fig. 8. 3D and 2D Binding-interaction diagrams of HDZPA ligand with SARS-CoV-2 main protease (a) and RNA dependent RNA polymerase (b).

| Compounds                  | MPro Binding energy in Kcal/mol | RdRp Binding energy in Kcal/mol |
|----------------------------|---------------------------------|--------------------------------|
| This work (HDZPA)          | −6.00                           | −7.70                          |
| Chloroquine [54]           | −4.9                            | −5.4                           |
| Hydroxychloroquine [54]   | −5.5                            | −5.6                           |
| Remdesivir [55,56]         | −4.96                           | −7.60                          |

**Author Statement**

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Nadjib Chafai: Conceptualization, Methodology, Validation, Supervision, Project administration, Resources and Investigation, Theoretical studies and Review.

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**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.molstruc.2021.130480.

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