Thermal and hydrodynamic properties of coronavirus at various temperature and pressure via molecular dynamics approach

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Abstract
COVID-19 is an epidemic virus arising from a freshly discovered coronavirus. Most people involved with the coronavirus will experience slight to moderate respiratory disease and recover without needing particular therapy. In this work, the atomic stability of the coronavirus at different thermodynamic properties such as temperature and pressure, was studied. For this purpose, the manner of this virus by atomic precession was described with a molecular dynamics approach. For the atomic stability of coronavirus description, physical properties such as temperature, total energy, volume variation, and atomic force of this structure were reported. In molecular dynamics approach, coronavirus is precisely simulated via S, O, N, and C atoms and performed Dreiding force field to describe these atoms interaction in the virus. Simulation results show that coronavirus stability has reciprocal relation with atomic temperature and pressure. Numerically, after 2.5 ns simulation, the potential energy varies from −31,163 to −26,041 eV by temperature changes from 300 to 400 K. Furthermore, this physical parameter decreases to −28,045 eV rate at 300 K and 2 bar pressure. The volume of coronavirus is another crucial parameter to the stability description of this structure. The simulation shows that coronavirus volume 92% and 14% increases by 100 K and 2 bar variation of simulation temperature and pressure, respectively.

Keywords Coronavirus · Molecular dynamic simulation · Temperature effect · Thermodynamic properties

Introduction
COVID-19 is an epidemic virus arising from a freshly discovered coronavirus. Most people involved with the coronavirus will experience slight to moderate respiratory disease and recover without needing particular therapy. Historically, coronaviruses were reported in the 1930s for the first time when some infection bronchitis virus has appeared from domesticated chickens shown [1] (see Fig. 1). Working at nanoscales levels such as nanoparticles dispersed in the base fluid might help to investigate the virus configuration [2, 3] through the porous medium [4]. The structure of coronaviruses is massive, occasionally pleomorphic (varying in shape), mainly spherical, particles with bulbous surface schemes [5]. The electron micrographs show that the envelope of coronavirus becomes visible as a distinct pair of electron-dense shells (is used to scan the virus particle, shells which are comparatively opaque to the electron beam) [6, 7]. The viral envelope contains a lipid bilayer that it would surround, and spike structural proteins are attached [8]. The proportion of envelope/spike/membrane in the lipid bilayer is about 1:20:300 [9]. Indifferently a coronavirus particle contains 74 surface spikes [10]. A subset of coronaviruses has a shorter spike similar to surface protein (hemagglutinin esterase) [11]. The spikes on the surface of coronavirus are homotrimers of the S protein, which is formed of an S1 and S2 subunit. One of the class I fusion protein is the homotrimeric S protein that intercedes the receptor binding
and membrane fusion between the cell and virus that be a host. The head of the spike is the S1 subunit forms, and it consists of the receptor-binding domain. One of the subunit forms (S2) is the stem, which attaches the spike in the viral envelope and on protease activation abilities to fusion. The M and E protein are significant in shaping the viral envelope and preserving its structural shape. In this envelope, there is the nucleocapsid, which is contained from multiple copies of the nucleocapsid (N) protein, bound to the positive-sense single-stranded RNA genome in a going on beads on a string type organization [12]. The preservation of the virus outside the host cell is done by the nucleocapsid, membrane proteins, and lipid bilayer envelope [13]. At this time, there are no particular vaccines or cure for coronavirus.

Nevertheless, various factors, such as temperature and pressure, appear to affect the manner of this virus and change its stability. Many works have reported the characteristics of this virus [14]. Lu et al. phylogenetic research report that bats might be one of the original hosts of coronavirus. An animal sold in Wuhan might represent an intermediate host, making possible the emergence of the virus in humans [15].

Furthermore, the analysis related to the structure of this research group reported that coronavirus could have the ability to connect to the angiotensin-converting enzyme two receptors in humans [16, 17]. It is not known how the coronavirus behaves in various environmental conditions. In this work, the atomic stability of coronavirus with the molecular dynamics (MD) technique was simulated. MD method is the computer simulations capable of predicting the dynamical evolution of molecules and atoms. In this process, simulated particles are permitted to interact in the simulation box and give a sight of the dynamic changes of the system. All molecular dynamics simulations in this research were done via employing the Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) provided by Sandia National Laboratories [23–25].

Methods

In these molecular dynamics simulations, it is used equilibrium MD simulations are used to describe the atomic stability of coronavirus in various temperatures and pressures. The molecular dynamics simulation method is a numerical approach to calculate the dynamical evolution of molecules and atoms. In this process, simulated particles are permitted to interact in the simulation box and give a sight of the dynamic changes of the system. All molecular dynamics simulations in this research were done via employing the Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) provided by Sandia National Laboratories [23–25]. To use of this MD simulation package to estimate the atomic stability of coronavirus with various temperatures ($T = 300–400$ K) and pressures ($P = 0–2$ bar), the coronavirus molecules were simulated by the atomic accuracy (Fig. 2). This nuclear structure was visualized by Open Visualization Tool (OVITO) [26].

In these MD simulations, aqueous environment is prepared by the Packmol package [27, 28]. Figure 3 shows the simulation box at the front, top, and perspective views. For coronavirus MD simulation in an aqueous environment, the directions $x$, $y$, and $z$ were used to show the periodic boundary conditions [29]. To duplicate the canonical phase-space dispersion, the use of this thermostat strives. This was accomplished by altering the equations of motion to contain a non-Newtonian term to preserve the
total kinetic energy of the atomic system constant. For the modified equation of atom movement, it can be written:

\[
\frac{dv(t)}{dt} = \frac{F(t)}{m} - \zeta v(t)
\]  

(1)

where \(\zeta\) is the thermodynamic friction coefficient, defined by Eq. (2):

\[
\frac{d\zeta}{dt} = \frac{1}{Q} \left[ \sum m v(t)^2 - (X + 1)k_B T \right]
\]  

(2)

In this equation, \(Q\) is a parameter that clears the time scale of the temperature fluctuation, and \(X\) is the number of degrees of freedom. After 2.5 ns MD simulation, the simulated atomic structures were equilibrated at initial condition (initial temperature and pressure). Furthermore, to MD simulate the atomic structure of coronavirus, the Dreiding force field was used [30]. The best choice of this interatomic force field is to affect biological structures. The potential energy for coronavirus atoms in the Dreiding force field was shown as a superposition of bonded and non-bonded ones. Non-bond interaction between atoms in the Dreiding force field is described by the Lennard–Jones (LJ) potential [31]:

\[
U(r) = 4\varepsilon \left[ \left( \frac{\sigma}{r_{ij}} \right)^{12} - \left( \frac{\sigma}{r_{ij}} \right)^{6} \right] \quad r \ll r_c
\]  

(3)

Both \(\varepsilon\) and \(\sigma\) parameters rely on the type of atoms in simulated structures in MD simulations. The energy and cutoff radius and length scale parameter, for different atoms in coronavirus simulation, are written in Table 1.

The bonded interaction in the Dreiding force field consists of bond strength, bond angle bend, and dihedral angle torsion terms. The bond, angle, and dihedral strength stretch in this force field described by harmonic oscillator equations as (4), (5), and (6):

\[
E_r = \frac{1}{2} k_r (r - r_0)
\]  

(4)
\[ E_0 = \frac{1}{2} k_0 (\theta - \theta_0) \]  
\[ E = K(1 + d \cos(n \phi)) \]  
\[ v(t + \Delta t) = v(t) + \frac{a(t) + a(t + \Delta t)}{2} \Delta t + O(\Delta t^3) \]  

These sympathetic constants in coronavirus MD simulation were selected at 300 (kcal/mol)/Å² and 100 (kcal/mol)/degree², while \( d = +1 \) or \(-1 \). Furthermore, the equilibrium value of angles and atomic bond length in these MD simulations is reported in Tables 2 and 3.

To aqueous environment definition in this computational work, \( \text{H}_2\text{O} \) molecules are simulated by the SPC model. This simple model can describe the atomic evolution of the \( \text{H}_2\text{O} \) structure very well in the presence of \( \text{model} \). This simple model can describe the atomic evolution of the \( \text{H}_2\text{O} \) structure very well in the presence of \( \text{model} \). After atomic modeling and force field implementation, the atomic manner of coronavirus in Table 1

| Parameter | \( r_d / \text{Å} \) |
|-----------|-------------------|
| C–C bond  | 1.530             |
| N–C bond  | 1.462             |
| O–C bond  | 1.420             |
| S–C bond  | 1.800             |
| O–N bond  | 1.352             |

Table 2 The bond strength of the equilibration distance for coronavirus structure in these MD simulations [30]

| Parameter | \( \theta_0 / \text{degree} \) |
|-----------|-------------------------------|
| C–C–C angle | 109.471                     |
| C–N–C angle | 106.700                     |
| C–S–C angle | 92.100                      |
| N–C–C angle | 109.471                     |
| N–C–N angle | 109.471                     |
| O–C–C angle | 109.471                     |
| O–C–O angle | 109.471                     |
| S–C–C angle | 109.471                     |
| N–O–C angle | 104.510                     |
| O–N–C angle | 106.700                     |

Table 3 The bond angle bend of the equilibration angle for coronavirus structure in these MD simulations [22, 30]
Step A Coronavirus in the aqueous environment was simulated at 300 K and $P = 0$ bar with $\Delta t = 1$ fs (time step); 150 Å length in $x$, $y$, and $z$ directions. In the equilibration process, the NPT ensemble with 0.01 and 1 for temperature and pressure damping rate (respectively) for 2.5 ns was used [33, 34].

Step B The temperature and pressure of simulated structures increase to 400 K and 2 bar after 2.5 ns. For the MD study of coronavirus stability in defined conditions, parameters such as temperature, total energy, and volume of coronavirus were reported.

Results

Equilibration of simulated structures

The atomic structure of coronavirus with $\text{H}_2\text{O}$ molecules in $T = 300$ K and $P = 0$ bar is studied, and final position of these structures is used in the second step of this MD study for the first time [35–37]. The results showed that the atomic model of coronavirus, which used in this MD study, is adopted with the Dreiding force field. Physically, atomic structures' stability is described by temperature and potential energy of them at $T = 300$ K and $P = 0$ bar. Temperature variation of simulated structures depicted in Fig. 4. From this figure, it can be said that virus and $\text{H}_2\text{O}$ molecules temperature equilibrated after 2.5 ns. Figure 5 displays the potential energy variation of simulated structures in initial condition (temperature and pressure). It can be seen that the atomic structures' potential energy converged after 2.5 ns. Numerically, the potential power of coronavirus in aqueous environment converged to $-31,163$ eV. From the result of this section of this report, it can be said that the Dreiding force field has excellent ability in coronavirus MD simulation.

Atomic stability of coronavirus in various temperature and pressure

For the atomic stability study of coronavirus in various temperature and pressure, first, temperature and potential energy variation in simulated structures were reported. Figure 6 shows that atomic structures' temperature reaches 350 and 400 K after 2.5 ns. As can be seen, a similar atomic manner for pressure variation of these structures and pressure of them reach after 2.5 ns to 1 bar and 2 bar. By equilibrium of atomic arrangements in the initial condition, the potential energy variation of these atoms was reported. These MD results show that by increasing the temperature of structures, their atomic stability decreases. Numerically, by increasing the temperature of coronavirus in the aqueous environment from 300 to 400 K, the potential energy of them decreases from $-31,163$ to $-26,041$ eV (Fig. 7 and Table 4). Physically, by increasing the temperature in the MD simulation box, various atoms in coronavirus structure move away and interatomic distance increases in this virus. The distance of atoms in simulated structures has a reciprocal relation with potential energy. So, by increasing the temperature, the potential energy of the system and stability of them decreases.
Furthermore, by increasing the pressure in simulated structures, the distance between atoms decreases. After this atomic phenomenon, repulsive interaction between atoms occurs, which causes an increase in atomic distance between them. By this atomic evolution, the potential energy and atomic stability of coronavirus in the aqueous environment decreases. Numerically by increasing the pressure from 0 to 2 bar, the potential energy of coronavirus in the aqueous environment decreases from $-31,163$ to $-28,045$ eV (see Fig. 8 and Table 5).

![Fig. 8](image)

**Fig. 8** Potential energy variation of simulated structures at various pressures as a function of simulation time

| Pressure/bar | Potential energy/eV |
|--------------|---------------------|
| 0            | $-31,163$           |
| 1            | $-29,607$           |
| 2            | $-28,045$           |

The atomic volume of coronavirus is proportional to its atomic stability in the aqueous environment. These MD results show that by MD simulation time passing, the volume of coronavirus increases (see Fig. 9). Physically, by increasing the simulation temperature and pressure, the distance of atoms in coronavirus structure increases and the stability of this virus decreases. From Fig. 10, by increasing the temperature from 300 to 400 K, the volume of coronaviruses increases from 179,091 to 343,437 Å$^3$. Pressure increasing has a similar effect on virus atomic volume and so on atomic stability. These simulation results show that by increasing the pressure from 0 to 2 bar, the volume of simulated structure changes from 179,091 to 205,283 Å$^3$ (Fig. 11 and Table 6). This atomic behavior indicates that the temperature and pressure increase can be used for this virus disinfection in medical applications. These physical processes can play an effective role in the elimination of

![Fig. 9](image)

**Fig. 9** Temperature increasing of coronavirus in an aqueous environment to 350 K and 400 K as a function of MD simulation time

![Fig. 10](image)

**Fig. 10** Temperature variation of coronavirus in an aqueous environment at various temperatures as a function of simulation time

| Temperature/K | Potential energy/eV |
|---------------|---------------------|
| 300           | $-31,163$           |
| 350           | $-27,663$           |
| 400           | $-26,041$           |

![Table 4](image)

**Table 4** Potential energy variations with temperature

| Temperature/K | Potential energy/eV |
|---------------|---------------------|
| 300           | $-31,163$           |
| 350           | $-27,663$           |
| 400           | $-26,041$           |
Thermal and hydrodynamic properties of coronavirus at various temperature and pressure via MD simulation results. Furthermore, from MD simulation results, it can be said that the temperature effect is more effective on atomic disinfection of coronavirus. The mutual force between the coronavirus and H$_2$O molecules describes virus evolution in MD simulations. In the final section of this study, the inserted force to coronavirus from the aqueous environment was calculated. Figure 12 shows this physical parameter for virus atoms in $T = 300$ K after 2.5 ns. The maximum rate of this parameter is 41 eV Å$^{-1}$. After temperature increasing from 300 to 400 K, the maximum rate of this force reaches 34 eV Å$^{-1}$ (Table 7). Physically, this parameter decreasing shows the interatomic potential decreasing, too. So it can be said that the atomic stability of the virus decreases by increase in temperature, and the coronavirus can be destroyed in high...
temperatures. Pressure increasing has a similar effect on the coronavirus atomic manner. By increasing the pressure from 0 to 2 bar, the maximum rate of inserted atomic force to coronavirus decreases from 41 to 37 eV Å⁻¹. Figure 13 shows the inserted force to coronavirus atoms in $T=300$ K and $P=2$ bar after 2.5 ns.

### Table 7 Inserted force to coronavirus as a function of temperature and pressure of simulated structures

| Temperature/K | Pressure/bar | Force/ eV Å⁻¹ |
|---------------|--------------|---------------|
| 300           | 0            | 41            |
| 350           | 0            | 36            |
| 400           | 0            | 34            |
| 300           | 1            | 39            |
| 300           | 2            | 35            |

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**Fig. 12** Inserted force to coronavirus atoms from the aqueous environment in $T=300$ K and $P=0$ bar after 2.5 ns

**Fig. 13** Inserted force to coronavirus atoms from the aqueous environment in $T=300$ K and $P=2$ bar after 2.5 ns
Conclusions

The present study investigates the atomic stability of coronavirus in an aqueous environment with MD simulations. In these calculations, coronavirus is shown via S, O, N, and C atoms and H2O molecules modeled by SPC. Also, to simulate interatomic force between virus atoms, the DreiDing force field has been selected. The result of these MD simulations shows that the increase in temperature and pressure causes coronavirus disinfection, which can be used for medical purposes. Results are as following:

- The total energy of this virus in aqueous environment converged to $-31,163 \text{ eV}$ after $2.5 \text{ ns}$ at $T=300 \text{ K}$ and $P=0 \text{ bar}$ which implies the atomic stability of simulated structures.
- The potential energy of coronavirus in the aqueous environment decreases by increase in temperature. This atomic manner shows that coronavirus stability and MD simulation temperature have a reciprocal relation.
- Numerically, by increasing the temperature from 300 to 400 K, the potential energy of coronavirus in the aqueous environment decreases from $-31,163$ to $-26,041 \text{ eV}$.
- The potential energy of coronavirus in the aqueous environment decreases by increasing the pressure. So atomic stability of these structures has reciprocal relation with MD simulation pressure.
- Numerically, by increasing the pressure from 0 to 2 bar, the potential energy of coronavirus in the aqueous environment decreases from $-31,163$ to $-28,045 \text{ eV}$.
- The volume of the coronavirus increases by increase in temperature in the MD simulation box. This atomic manner shows that unstability was growing in the virus by increase in temperature. Numerically, coronavirus volume increases from 179,091 to 343,437 $\text{Å}^3$ after temperature increasing from 300 to 400 K.
- The volume of the coronavirus in the aqueous environment increases by increase in pressure in the MD simulation box. Numerically, coronavirus volume increases from 179,091 to 205,283 $\text{Å}^3$ after pressure increases from 0 to 1 bar.
- Coronavirus and aqueous environment mutual atomic force changes from 41 to 34 $\text{eV} \text{ Å}^{-1}$ by the change in thermodynamic properties such as temperature and pressure variations.

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Data availability The data that support the findings of this study are available from the corresponding author upon reasonable request.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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