First-principles and Molecular Dynamics simulation studies of functionalization of Au$_{32}$ golden fullerene with amino acids

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With the growing potential applications of nanoparticles in biomedicine especially the increasing concerns of nanotoxicity of gold nanoparticles, the interaction between protein and nanoparticles is proving to be of fundamental interest for bio-functionalization of materials. The interaction of glycine (Gly) amino acid with Au$_{32}$ fullerene was first investigated with B3LYP-D3/TZVP model. Several forms of glycine were selected to better understand the trends in binding nature of glycine interacting with the nanocage. We have evaluated various stable configurations of the Gly/Au$_{32}$ complexes and the calculated adsorption energies and AIM analysis indicate that non-Gly, z-Gly and also tripeptide glycine can form stable bindings with Au$_{32}$ at aqueous solution via their amino nitrogen (N) and/or carbonyl/carboxyl oxygen (O) active sites. Furthermore, cysteine, tyrosine, histidine and phenylalanine amino acids bound also strongly to the Au$_{32}$ nanocage. Electronic structures and quantum molecular descriptors calculations also demonstrate the significant changes in the electronic properties of the nanocage due to the attachment of selected amino acids. DFT based MD simulation for the most stable complex demonstrate that Gly/Au$_{32}$ complex is quite stable at ambient condition. Our first-principles findings offer fundamental insights into the functionalization of Au$_{32}$ nanocage and envisage its applicability as novel carrier of the drugs.

Drug delivery has gained a lot of attention as it can minimize the side effects of various drugs. Practically, an ideal drug delivery system is an increasingly effective disease treatment by delivery of a drug to target organs and lowering the side effects as well as sustained release formulations in which the drug is released over a period of time in a controlled manner. In particular, the development of drug delivery strategies has focused on nanoscale particles to improve bioavailability, selectivity and release rate. Nano drug delivery technology has been fully applied in the treatment course of tumor, diabetes, vascular disease and cancer patients, and can provide some guidance for clinical treatment work. Recently, Nanoscience has been used vastly in drug delivery and lots of research groups have been working on this topic for years to make biocompatible and safe drug carriers delivering a handful of dug to predetermined target cells in a controlled release manner.

Various drug delivery systems based on different nanomaterials, such as biodegradable polymers, calcium phosphate, nanoporous silica, Fe$_3$O$_4$ and carbon based nanomaterials have been investigated in the past decade. Among all of the nanoparticles, gold NPs have gained a lot of attentions since they exhibit anticancer activities along with being a drug carrier. Also, they have interactions with DNA and RNA and other biological molecules. Gold nanoparticles with different shapes and sizes show their unique physical and chemical properties for transporting and unloading the pharmaceuticals. Last but not the least, the gold nanoparticles can be used for hyperthermia since they can absorb near IR and generate heat.

However, the gold nanoparticles suffer from low drug loading capacities. Recent discoveries of three dimensional gold based particles, including gold clusters including Au$_{19}$, Au$_{32}$, Au$_{50}$ and Au$_{72}$ or golden fullerenes have provided new opportunities to use gold based carriers. Apart from the tetrahedral Au$_{19}$ and Au$_{20}$ clusters, 

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a new class of golden cages has been proposed using both experimental and theoretical methods. Among the three sizes, Au$_{32}$ has been the most intriguing cluster because of its ground state geometry which is an icosahedral golden fullerene. The report showed that the spherically hollow Au$_{32}$ conformation with nearly 0.9 nm diameter is structurally similar to C$_{60}$. On the other hand, the ground state Au$_{32}$ geometry has been proposed to be aromatic in nature.

The other part of drug delivery involves release time, which strongly depends on carrier and drug interaction. Thus, exploring the interactions between biologically relevant molecules and Au$_{32}$ surface seems to be of crucial importance for designing efficient drug carriers. The predictions from theoretical point of view help the scientists to understand the loading and the release time, loading capacities and even loading and release mechanism. Moreover, knowing the interactions can predict the delivery selectivity and efficiency. Meanwhile, since amino acids are the elementary component of biomolecules and can also mimic the chemical properties of complicated biomolecules, it is important to understand the interaction between drug carrier and typical amino acids. Among all the 20 elementary amino acids, glycine might be the best candidate since it is small which makes it possible to investigate their interactions with gold fullerene with a rational computational cost. However, as biomolecules such as proteins are much more complicated than the glycine amino acid we have also considered cysteine, tyrosine, histidine and, phenylalanine amino acids which consist amino nitrogen, carboxyl oxygen, carbonyl oxygen, hydroxyl oxygen, imidazole, sulfur and, aromatic rings.

Herein we study the interaction between fullerene-like gold nanocage and selected amino acids by using the first-principles calculations at the DFT-B3LYP level of theory. DFT calculations have been tremendously applied and shown by several researchers to successfully predict the structural and electronic properties of molecular systems interacting with nanostructured based materials. Adsorption energy is calculated and amino acids binding to the gold fullerene nanocage is investigated. We also analyzed the electronic structures and molecular properties for the energetically most favorable complexes. Our results indicate that selected amino acids can form stable bindings to the gold nanocage via their potential active sites. Our theoretical predictions have shed light toward the functionalization of gold nanocages with biomolecules and provide us a rational design and preparation of novel gold-based nanomaterials for drug delivery.

Computational Methodology

The first-principles DFT method was employed to calculate the structure and energy of the systems under study. All calculations were performed using the ORCA quantum chemistry code (version 3.0.0) at the B3LYP/def2-TZVP level of theory. In order to consider the long range non-local interactions, the atom-pairwise dispersion correction (D3) with Becke–Johnson damping (B) was utilized for the systems studied here. The optimized atomic structures of the considered systems were obtained by the density-fitting (resolution-of-the-identity approximation) and chain-of-spheres methods (RIJCOSX) to accelerate the calculations without loss of accuracy. Furthermore, we utilized the [SD (60, MDF)] effective core potential (ECP) for the Au atoms. The nucleus effective charge method was also used for the Au atom. The solvent effects have been described by conductor-like screening models (COSMO) to consider the electrostatic interaction of molecules with solvent. The adsorption energy was calculated as the difference between the energy of Gly/Au$_{32}$ complexes and the sum of the energies of the corresponding glycine and Au$_{32}$ fullerene ($E_{\text{ads}} = E(Gly) + E(Au_{32}) - E(Gly/Au_{32})$). To eliminate the basis set superposition errors (BSSEs) the Boys–Bernardián counterpoise (CP) scheme was utilized. It should be noted that positive/negative adsorption energy indicates endothermic/exothermic process, respectively.

We used the quantum mechanical descriptors (QMD) to describe the electronic properties of the glycine molecule and its complexes with the Au$_{32}$ nanocage consisting of ionization potential (IP), electron affinity (EA), global hardness ($\eta$) and energy gap ($E_g$). According to the Koopmans theorem, the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) energy level are related to the IP and EA, respectively as follows:

$$\text{IP} \approx -E_{\text{HOMO}} \quad \text{and} \quad \text{EA} \approx -E_{\text{LUMO}}$$

(1)

The global hardness is calculated using the relation:

$$\eta = \frac{\text{IP} - \text{EA}}{2}$$

(2)

The accuracy of our implemented method is validated elsewhere.

Chemical bonds nature was investigated and characterized using the quantum theory of atoms in molecules (QTAIM) scheme or AIM theory which its theoretical basis has been detailed elsewhere. AAIM analysis was carried out to deeply understand the interaction natures between Au$_{32}$ fullerene and different amino acids. We used ORCA software to obtain the wave-function used in the bonding analysis at the B3LYP level of theory. The topological analysis, the evaluation of local properties and surface electrostatic potential maxima and minima points were calculated and visualized using the Multiwfn program package.

Based on the AIM theory, when two atoms form a bond, a bonding critical point (BCP) appears between the formed bonds. At the BCP, the $\rho(r)$ and the sign of its Laplacian determines whether the charge is concentrated as in covalent bonds ($\nabla^2 \rho(r) < 0$) or is depleted as in closed shell (electrostatic) interactions ($\nabla^2 \rho(r) > 0$). For the shared interactions, the accumulation of electron density can be easily observed across the line that links the involving nuclei whereas for the closed shell interactions the accumulation of charges can be seen at the terminal of interacting nuclei and the BCP in the middle of the bond accounts for the depletion of the electron density.

According to the Bader's theory, at the BCPs, the total energy density ($H(\nabla^2 \rho)$) is related to the Laplacian of the electron density by the following equation:
where \( G(r_{BCP}) \) denotes the kinetic energy density which is always a positive value. A pure covalent bond is usually represented by a negative Laplacian and a negative energy density while pure closed shell bonds such as strong hydrogen bonds and ionic bonds are characterized by positive values for Laplacian and energy density. Their intermediated i.e. partially covalent and highly polar bonds, are classified by positive Laplacian and negative energy density. However, it is noteworthy to mention that there are also exceptions which do not completely follow the aforementioned rules regarding the type of bonds and these criteria should be considered together with the other analysis.

We have further performed first-principles MD simulation using the Spanish Initiative for Electronic Simulations with Thousands of Atoms (SIESTA) code which employs the Born-Oppenheimer dynamics approximation for simulation of molecular systems. To describe the interaction between electrons and ions we utilized the norm-conserving Troullier-Martins pseudopotential and the generalized gradient approximation (GGA) parameterized by Perdew, Burke and Ernzerhof (PBE) is employed for the exchange-correlation potential. The localized atomic orbitals for valence wave functions and double-\( \zeta \) basis sets are employed, with a cutoff energy of 125 Ry for system under study. The Velocity Verlet algorithm was employed for solving the Newton's equations of motion. The NVT ensemble was used with a Nosé–Hoover thermostat to maintain the temperature at 300 K.

### Results and Discussion

#### Interaction between non-ionic and zwitterion glycine and Au\(_{32}\)

We start by describing the equilibrium and electronic structures of golden fullerene and non-ionic glycine (non-Gly) molecules at the gas phase obtained using the B3LYP-D3/TZVP theoretical model. Figure 1 shows the optimized geometries of considered molecules. The average Au–Au bond length in golden fullerene is calculated to be 2.8 Å which agrees well with other theoretical studies. Further, the calculated bond lengths of glycine molecule are in good agreement with the literature studied. The charge analysis of optimized geometries depicts various active sites for molecules under consideration. It was found from the atomic charge analysis that the Au atoms placed at the center of the pentagon ring (Au-pent) are positively charged hence this half-occupied local state can consequently act as an acceptor entity. This finding reveals that point charges upon the Au\(_{32}\) nanocage can improve the binding capability and therefore increase the binding strength between the glycine and the nanocage. On the other hand, the O and N atoms of the glycine are found to be potential electron rich centers and therefore one can expect to glycine molecule attacks to positively charged Au atoms (Au-pent) of golden fullerene via these active sites.

To find the most stable geometries of glycine molecule interacting with golden fullerene, we approached the glycine molecule to the positively charged Au atom (Au-pent) via its N and O active sites. Then, all the considered structures were fully optimized at the B3LYP-D3/TZVP level of theory. The bonding energy and equilibrium distance (the closest atoms of the interacting molecules) for energetically favorable complex are determined to be about −1.364 eV (−31.454 kcal/mol) and 2.336 Å, respectively. Figure 1c represents the most stable geometry of non-Gly/Au\(_{32}\) complex obtained at the gas phase. As it can be found from the figure, glycine has a significant accumulation and the charges mainly accumulate at the terminal regions of the involved atoms in the bonding region between N and Au atom, there is no evidence of charge accumulation and the charges mainly accumulate at the terminal regions of the involved atoms in the bonding process. This binding behavior can also partially be explained by considering the geometrical parameters of the mentioned values for the isolated fragments at the optimized geometry of complex according to the below formula:

\[
\Delta \rho = \rho_{\text{total}} - (\rho_{\text{lab}} + \rho_{\text{ads}})
\]

The electron density difference plot for the formed complex is illustrated in Fig. 1d.

The accumulation of electron is represented by yellow color while the electron depletion is shown by blue. It can be seen from the figure that, in the bonding region between N and Au atom, there is no evidence of charge accumulation and the charges mainly accumulate at the terminal regions of the involved atoms in the bonding process. This binding behavior can also partially be explained by considering the geometrical parameters of the amino group of the glycine before and after the adsorption process. It can be seen from the optimized structure of the glycine after the adsorption that the N–C bond length has been increased from 1.452 Å to 1.469 Å. Since the highest occupied molecular orbital (HOMO) of the glycine is mainly located on the nitrogen of the amino group, hence this atom serves as the center of the reactivity and the increase in N-C bond length, facilitates the transfer of electrons from the glycine molecule to the Au\(_{32}\) surface. Upon the adsorption of glycine, the HOMO of the adsorbate donates electron to the lowest unoccupied molecular orbital (LUMO) of the substrate which is mainly localized on the pentagon of the Au\(_{32}\) molecule in conjunction with some electron back donation to the unoccupied states of the adsorbate molecule as can be seen from the accumulated electrons around the carbon atoms of the glycine. This charge transfer in turn polarizes the substrate and gives rise to the electrostatic interaction between the involved molecules which is expected to plays important role in the attachment of glycine onto the Au\(_{32}\) surface.
To further clarify the nature of the formed bond between considered atoms, analysis of the electronic charge density ($\rho(r)$) and its Laplacian ($\nabla^2 \rho(r)$) was carried out using the AIM theory.

Figure 1e illustrates the optimized structure of glycine/Au$_{32}$ complex and the BCPs are shown by orange color. Table 1 lists the calculated results including the Laplacian and energy densities of complex under consideration.
As can be seen from the Fig. 1e and Table 1, a BCP is evident in the bonding region of nitrogen and Au atom with the $V^p(r)$ value of 0.214 which shows a strong depletion of charge at this critical point and $H(r_{BCP})$ of about $-0.011$ a.u which shows that there exists an attractive interaction between the two nuclei. Based on these values, this bond has a highly polar nature similar to the situation for coordination and ionic bonds. This is completely in line with the results of charge transfer analysis based on the Hirshfeld method which suggests the transfer of 0.21 e from the glycine to the Au$_{32}$ and also the results of charge decomposition analysis (CDA) (not shown here) that the HOMO of the glycine and LUMO of the Au$_{32}$ exhibit the highest contribution to the charge transfer between the two molecules. This corroborates the fact the lone pair of the nitrogen donates electrons to the LUMO of the Au$_{32}$ which resembles a coordination bond rather than a normal covalent bond. These findings reveal a strong interaction between glycine molecule and gold fullerene thus one can conclude that non-Gly/Au$_{32}$ complex is energetically stable at the gas phase.

In order to investigate the influence of solvent on the stability of considered complex we have performed similar calculation procedure for the interacting molecules at aqueous solution. All molecular systems were optimized separately at aqueous solution and then focused on the interaction between two entities. Schematic representation of optimized structure of non-Gly/Au$_{32}$ fullerene complex in aqueous solution is given in Fig. 2. Our $ab$ initio results based on the DFT-B3LYP-D3 method with TZVP basis set indicate that considered complex is stable at aqueous solution with the adsorption energy of about $-1.444$ eV and bonding distance of 2.255 Å. It can be clearly found from the binding information that non-Gly/Au$_{32}$ fullerene complex is energetically more stable in aqueous solution than the gas phase. Furthermore, we have evaluated the binding strength of non-Gly molecule interacting with Au atom placed at the hexagon ring of the Au$_{32}$ nanocage. After full structural optimization on the molecular systems with B3LYP-D3/TZVP model, the binding energy was found to be about $-1.40$ eV which indicates that non-Gly prefer to be bound to the Au atom of pentagon ring of the nanocage.

We now consider the interaction between zwitterion glycine (z-Gly) amino acid (Fig. 2c) and gold fullerene at aqueous solution. We modeled the z-Gly/Au$_{32}$ nanocage complex so that O atom of carboxyl group was placed above the Au atom of pentagon ring while H atom of NH$_3$ group was positioned adjacent to the Au atom of hexagon ring. Full structural optimization was carried out for estimation of adsorption energy for z-Gly/Au$_{32}$ complex. The calculated adsorption energy at the B3LYP-D3/TZVP level showed that z-Gly prefers to be adsorbed at the center of pentagon of golden fullerene via its O atom (see Fig. 2d) with adsorption energy of $-1.112$ eV and the optimum interacting distance of about 2.267 Å. As a result, the adsorption energy values are highly negative for both forms of glycine amino acid interacting with Au$_{32}$ fullerene suggesting the thermodynamic favorability of respective complexes in aqueous solution though non-Gly counterpart undergoes stronger interaction with the Au$_{32}$ fullerene. Indeed, formation of such complexes seems to be experimentally possible from energetics point of view.

We have further calculated the isosurface corresponding to HOMO and the LUMO states for molecular systems under study (see Fig. 2). In the case of golden fullerene the HOMO state are delocalized on the Au atoms placed at the center of both hexagon and pentagon rings while the LUMO states are mainly localized on the Au atoms of the pentagon center as represented in Fig. 2e. Figure 2f represents the HOMO electronic state to be placed at the center of both hexagon and pentagon rings while the LUMO states are mainly localized on the Au atom placed at the hexagon ring of the Au$_{32}$ fullerene. Indeed, formation of such complexes seems to be experimentally possible from energetics point of view.

Furthermore, the energy gap, global reactivity descriptors and dipole moment values have also been calculated for non-ionic/zwitterion glycine-Au$_{32}$ complexes. A comparison of HOMO–LUMO gap and global reactivity descriptors for non-Gly and z-Gly indicate glycine to be quite stable with energy gap around 6.635–7.395 eV and $\eta$ value of 0.146–0.135 eV. The gold fullerene is stable due to its HOMO–LUMO gap of about 2.177 eV and $\eta$ value of 0.459 eV, while Au$_{32}$ fullerene is more reactive than glycine amino acid. Glycine adsorption onto the gold fullerene shows a small decrease in HOMO–LUMO energy gap and increase in $\eta$ value indicating that glycine adsorption renders slightly stability to Au$_{32}$ nanocage. Adsorption of non-Gly onto the Au$_{32}$ fullerene demonstrate more reduction in energy gap value than that of z-Gly counterpart which can be conducive with the rather higher adsorption energy. Furthermore, it was found that the presence of nanocage for loading of glycine facilitates in the increased reactivity of glycine amino acid compared to pristine glycine. The calculated dipole moment which tolerates correlation with adsorption energy values shows that non-Gly/Au$_{32}$ complex has slightly higher dipole moment value than the z-Gly/Au$_{32}$ complex. This finding confirms also the greater extent of adsorption

| Glycine | Anionic Glycine | Tripeptide glycine |
|--------|----------------|------------------|
| BCP1 | BCP1 | BCP2 | BCP3 | BCP1 | BCP2 |
| $\rho(r)$ | 0.066 | 0.088 | 0.057 | 0.065 | 0.070 | 0.053 |
| $V^p(r)$ | 0.214 | 0.275 | 0.228 | 0.267 | 0.278 | 0.213 |
| $H(r)$ | $-0.011$ | $-0.022$ | $-0.006$ | $-0.009$ | $-0.012$ | $-0.005$ |
| $G(r)$ | 0.064 | 0.091 | 0.063 | 0.075 | 0.081 | 0.058 |

Table 1. Calculated Laplacian and energy densities with B3LYP-TZVP model for Glycine/Anionic Glycine/Tripeptide glycine-Au$_{32}$ complexes.
of non-Gly amino acid interacting with Au$_{32}$ fullerene compared to the z-Gly one and also more solubility of the non-Gly/Au$_{32}$ complex rather than other counterpart.

**Interaction between tripeptide glycine and Au$_{32}$.** To extrapolate the upshots of the current work for larger biological systems that reasonably mimic -for example- the behavior of proteins upon the attachment to Au nanoparticles, we have selected a tripeptide model of the zwitterionic glycine (see Fig. 3) and evaluated the interactions between this biomolecule with Au$_{32}$ nanocage to obtain more realistic results. From the optimized
structure of the most stable configuration of tripeptide glycine upon the interaction with Au$_{32}$ fullerene, strong deformations can be observed in the surface of Au$_{32}$ structure together with the formation of bonds between the oxygen atom of the carboxyl and carbonyl groups and the neighboring Au atoms as represented in Fig. 3b. This adsorption configuration accompanies by the release of about $-2.058 \text{ eV}$ of energy which is high enough to be considered as a stable complex. However, to unveil the nature of the interactions between the two molecules and illustrate the type of the formed bonds, we have performed AIM analysis similar to the previous section and tabulated the results including the Laplacian and energy densities in Table 1. Visualization of the BCPs for the considered complex as shown in Fig. 3c illustrates the appearance of two BCPs within the region of formed bonds which we marked as BCP1 and BCP2. Based on the values in Table 1, both BCPs have positive Laplacian and negative energy densities which represent an intermediate interaction similar to the situation for neutral glycine. However, the magnitude of attractive interactions differs considering the values of energy density for the BCPs. This is in line with the calculated bond lengths where the bond corresponding to the BCP1 has shorter length than that of the BCP2.

**Figure 3.** Optimized structures of (a) Tripeptide glycine molecule and (b) Tripeptide Gly/Au$_{32}$ complex. (c) The bonding critical point (BCP) for the formed tripeptide glycine/Au$_{32}$ complex.
The geometrical parameters of the adsorbed amino acid also corroborate the above statements regarding the type of the interactions as the C=O bond lengths where the O-Au bond has been formed are between the range of 1.24 to 1.26 Å which are typical bond lengths for the carbon and oxygen double bond. In this vein, the hybridization of the C=O bond within the region of bond formation did not change which resembles a charge transfer mediated electrostatic interaction together with some structural deformations within the involved molecules. This is consistent with the results of charge transfer analysis which illustrates the transfer of 0.4 e from the amino acid to the Au$_{12}$ surface. As a result we could consider the Au$_{12}$ nanocage as a suitable carrier with retained biological potency which made it target for further chemical modification in the drug delivery technology.

**Interaction between ionic glycine and Au$_{32}$.** The interaction between ionic forms of glycine amino acid and Au$_{32}$ fullerene has also been taken into consideration. For this end, individually optimized structures of anionic and cationic glycine molecule were selected and the stability of the formed glycine/Au$_{32}$ complexes was evaluated through adsorption energy calculations and electronic structure analysis. The optimized structures for the most stable adsorption configurations of cationic and anionic glycine are illustrated in Fig. 4. It can be seen from the figure that the cationic form of glycine does not show any bond formation with the Au$_{32}$ cage skeleton and no structural deformations was seen after the adsorption of cationic form (See Fig. 4b). Indeed, both the Au$_{32}$ and cationic glycine retained their isolated-optimized structures and the cationic form positioned near the Au$_{32}$ surface. This orientation can be easily explained by calculating the molecular electrostatic potential (MEP) maps of the isolated fragments at their optimized geometry as can be observed in Fig. 4b,c. It can be seen from these plots that the local surface maxima are mainly distributed over the pentagon rings of the Au$_{32}$ and the NH$_3$ of the cation. Meanwhile, surface minima are mainly located near the hexagonal rings of the Au$_{32}$ while the glycine molecule does not show any surface minima mainly because of its cationic nature. In this regard, the cationic glycine tends to orient above the Au$_{32}$ in a way that its surface maxima approaches close to the surface minima of the Au$_{32}$ in a maximally ESP complementary manner to increase the electrostatic attraction, as can be clearly seen from the optimized structure of the complex system.

The situation on the other hand is slightly different for the anionic glycine where the adsorption of this molecule is accompanied by the release of about $-2.699 \text{ eV}$ of energy which is considerably higher than that for the cationic counterpart together with strong deformations within the structures of both adsorbate and substrate as can be seen from the Fig. 4d. From the optimized structure it is evident that the glycine anion tilted at the carbon atom which is attached to the amino group with the tilt angle of about 108° after the attachment to the Au$_{32}$ molecule and three bonds have been formed between the carboxyl and amino groups and their adjacent Au atoms. The length of all the formed bonds slightly exceeds the sum of the covalent radii of the respective atoms involved within the bond and these bonds are unlikely to be normal covalent. To clarify the nature of the formed bonds, similar to previous sections AIM analysis was carried out and the results have been given in Table 1. Based on the Laplacian and energy density values for all of the BCPs as shown in Fig. 4e, the formed bonds are within the domain of intermediate bonds as no electron accumulation can be signified from the positive Laplacian while the negative energy densities inform the existence of attractive interactions between the involved atoms of the bonds.

These bonds are mainly governed by one-sided electron donation from the glycine anion to the substrate along with the deformations within the structures of adsorbate and substrate. This electron donation is evident by calculating the transferred charge between the fragments which shows the transfer of 0.8 e from the glycine to the Au$_{32}$.

**Interaction of cysteine, histidine, phenylalanine and tyrosine with Au$_{32}$.** We now consider other amino acids (cysteine, histidine, phenylalanine and tyrosine) interacting with Au$_{32}$ fullerene and evaluate the binding properties of formed complexes. Similar calculations procedures has been carried out for systems under study. Various orientations considering the potential active sites of interacting molecules such as sulfur and aromatic rings of amino acids and pentagon/hexagon Au atom of golden fullerene have been considered for interacting systems and full structural optimization has been performed followed by the adsorption energy estimation. Our B3LYP-D3/TZVP calculations results demonstrate that all considered amino acids bound strongly to the Au$_{32}$ fullerene surface with adsorption energy of about $-1.0 \text{ eV}$ ($-23.06 \text{ kCal/mol}$).

Cysteine amino acid prefers to bind to the fullerene nanocage through its sulfur atom to the Au atom at the pentagon tip with adsorption energy of $-1.172 \text{ eV}$ comparable to the Gly/Au$_{32}$ complex. The calculated bonding distance was estimated to be about 2.448 Å which is close to the experimental value of Au-S bond length for the adsorbed thiocyanates on the Au surface$^{22}$. The optimized geometry of Cys/Au$_{32}$ complex is shown in Fig. 5a. Further investigation about the bond nature of the attached molecule has been carried out by AIM analysis. The result indicates a positive value of 0.187 and negative value of $-0.018$ for Laplacian and energy density, respectively, which indicates that there exists similar interaction nature (partially covalent and highly polar bonds) for complex under study.

For three other amino acids, our *first-principles* calculations reveal that these molecules were attached to gold fullerene through their C atom in aromatic ring as represented in Fig. 5b–d. The calculated equilibrium distances of Au–C were estimated to be about 2.5 Å which is comparable to similar system (Au dimer adsorbed on the benzene ring) with DFT method at B3LYP/LACVP** level of theory$^{23}$. The AIM analysis for tyrosine/Au$_{32}$ complex demonstrated a similar trend for the Laplacian (0.077) and energy density ($-0.005$) values indicating similar interacting nature between interacting systems.

The calculated adsorption energies accompanied with charge transfer and optimum distances between interacting systems for more stable configurations are listed in Table 2. As can be seen from the obtained results one can conclude that the interaction natures of selected amino acids are also typical for the chemisorption and the complexes are energetically stable.
First-principles molecular dynamics (MD) simulation. As a rule of thumb the first-principles molecular dynamics (MD) simulation has been carried out to assess the binding nature as well as the stability of Gly/Au$_{12}$ complex at ambient condition. MD simulation can reliably explore the global minima state of complex under investigation with realistic simulation of interacting systems by considering explicit water molecules at ambient condition. To this aim, we have further employed an expensive MD simulation based on DFT-D method for the energetically favorable complex (non-Gly/Au$_{12}$) obtained by B3LYP-D3/TZVP model. The simulation system is filled with the non-Gly/Au$_{12}$ complex and 25 water molecules beside the complex as depicted in Fig. 6. We then performed 10 ps of simulation times with 1.0 fs time-step in order to obtain the optimized structures of the system under study at ambient conditions.

Our DFT-D based MD simulation results show that the aqueous solution (explicit solvent) affects the stability of glycine isomers and slightly on the geometries of the complex at ambient condition. We found that Gly molecule

![Figure 4. Optimized structures of (a) non-glycine molecule, (b) non-Gly/Au$_{12}$ complex, (c) z-glycine molecule and, (d) z-Gly/Au$_{12}$ complex, at aqueous solution obtained with the B3LYP-D3-TZVP level of theory.](image)
bound to the Au-pentagon atom via its amino –NH₂ active site which confirms the current DFT-B3LYP optimization results for glycine molecule attached to the Au atoms of pentagon and hexagon ring. The calculated average value of Au–N bond length (2.504 Å) reveals that the equilibrium bond distance slightly enlarge (the optimized bond length of 2.255 Å). Meanwhile, we observe that the H atom from hydroxyl –OH group of glycine attack to the O atom of water molecule at about 250 fs of simulation time and new bond forms between glycine and water molecules. The average value of O-H bond in glycine was found to be enlarged to 1.386 Å (the initial bond length was 0.973 Å) which indicate that O-H bond was dissociated in the aqueous solution during the simulation time (See Fig. 6c,d). Indeed, one can conclude that the zwitterion glycine isomer is the stable form in the aqueous solution at room temperature. These findings according to the first-principles MD simulation indicate that Gly-Au₃₂ complex is quite stable and thus it is possible to be used as suitable nanocarrier at ambient condition.

**Conclusions**

We have investigated the interaction between Au₃₂ fullerene and glycine amino acid with the state-of-the-art DFT calculations at the B3LYP-D3/TZVP level of theory. Various possible orientations have been considered for a glycine molecule attaching to the surface of Au₃₂ nanocage. Our DFT calculations at the gas phase demonstrated that non-Gly molecule strongly bound to the outer surface of the nanocage on the top site directly above the Au-pent atom via its amino N active site. The calculated adsorption energy and bonding distance provide sufficient evidence to conclude that the adsorption of glycine onto Au₃₂ nanocage is exothermic and the complex is energetically stable. Furthermore, non-Gly amino acid has stronger interaction with Au₃₂ fullerene in aqueous solution than the gas

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**Figure 5.** Optimized structure of (a) cysteine/Au₃₂, (b) tyrosine/Au₃₂, (c) histidine/Au₃₂ and (d) phenylalanine/Au₃₂ complexes.

| Systems            | Eads (eV) | Hirshfeld Charge Transfer (e) | Bonding Distance (Å) |
|--------------------|-----------|-------------------------------|----------------------|
| Cysteine/Au₃₂      | -1.172    | 0.281                         | 2.448                |
| Histidine/Au₃₂     | -1.096    | 0.217                         | 2.427                |
| Phenylalanine/Au₃₂ | -1.085    | 0.165                         | 2.551                |
| Tyrosine/Au₃₂      | -1.109    | 0.190                         | 2.504                |

Table 2. Calculated adsorption energies, charge transfer and bonding distances for cysteine/tyrosine/histidine/phenylalanine-Au₃₂ complexes.
phase and the respective complex is energetically more stable. Meanwhile, it was found greater extent of adsorption of non-Gly molecule interacting with Au32 nanocage rather than z-Gly counterpart. The calculated electronic structures and charge analysis revealed that the presence of point charges in the Au32 nanocage make them suitable candidate for strong binding to glycine amino acid. Furthermore, the global hardness, energy gap and ionization potential of Gly/Au32 complex were decreased which increase the reactivity of the respective system.

In order to simulate a realistic model of biomolecule interacting with Au32 nanocage we have considered a tripeptide glycine system with zwitterionic structure. Our DFT-D3 based optimization procedure showed that tripeptide glycine bind strongly to the Au32 nanocage with binding energy of about $-2.0 \text{ eV}$ via its carbonyl and carboxyl O atoms active sites. This indicated that tripeptide glycine can form energetically more stable complex rather than both individual non-Gly and z-Gly counterparts. As the rule of thumb for more complicated systems such as proteins we have further evaluated the interaction between Au32 nanocage and other amino acids (cysteine, histidine,

Figure 6. Schematic illustration of (a) simulation box filled with 25 water molecules and the energetically favorable configuration of non-Gly/Au32 complex and (b) snapshot of non-Gly/Au32 complex at 10 ps of simulation time. Time evolution of equilibrium bonding distance between (c) Au and N atoms of Au32 nanocage and glycine molecule and (d) H and O atoms of the glycine.
phenylalanine and tyrosine) including sulfur, imidazole and aromatic groups. Full structural optimization demonstrated that all selected amino acids strongly bound to the Au_{12} cage skeleton with bonding distances of about 2.5 Å and adsorption energy values of about —1.0 eV indicating that the complexes are energetically stable.

A comprehensive DFT based MD simulation at ambient condition has been carried out starting from energetically favorable complex (non-Gly/Au_{12}) obtained with B3LYP-D3/TZVP model. The simulation results demonstrated that Gly/Au_{12} complex was quite stable in aqueous solution while non-Gly isomer contribute to a proton transfer phenomenon with adjacent water molecules and thus z-Gly form was found to be the stable isomer in the ambient condition. From the present findings one can predict that proteins which contain hydroxyl oxygen, amino nitrogen and carbonyl oxygen active sites can form stable bindings with Au_{12} fullerene via its potential active sites. Our first-principles DFT results provide a well-grounded understanding for the possible formation of complex between bio-molecules and Au_{12} fullerene and can be benchmark for directing experimental efforts for the development of drug delivery at the nanoscale.

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Author Contributions
M.D.G. designed the project. R.A. and H.T.L. modeled the systems. M.M. derived the theory. M.D.G., wrote the code and M.D.G., H.T.L. and R.A. performed calculations. All authors contributed to the data analysis and writing of the paper.

Additional Information
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