Physico-chemical study of complexation of silver ion (Ag\(^+\)) by macrocyclic molecules (hexa-Helicenes) based on statistical physics theory: new description of a cancer drug

Manel Ben Yahia\(^1\) & Mohamed Ben Yahia\(^2\)

In recent papers, it is found that the silver-[6]Helicene complex can be used as a cancer drug but the interaction silver-hexaHelicene has not yet proven. The idea of this paper is to investigate the complexation process of the [6]Helicene by the silver metal (Ag\(^+\)) using three types of adsorbates. Indeed, the adsorption of silver chloride, silver nitrate and silver sulfide into the sensor films deposited on the QCM electrode are measured at three temperatures (293–333 K). Films of the [6]Helicene were deposited on the QCM resonators using spin coating method in order to obtain uniform and homogenous sensor surface. Experimental results indicated that the [6]Helicene can form a stable complex with the silver ion and that the AgCl is the appropriate adsorbate for the complexation achievement. Actually, an advanced modeling analysis by means of statistical physics adsorption models is applied to explore the new vision of the complexation system. The values of the models parameters are deduced from fitting the experimental data with the developed models. They result in confirming the experimental findings by comparing the complexation energies of the three examined systems. In particular, for the silver nitrate, the Van-der-Waals parameters explained the isotherms drop at high concentration through the lateral interactions between the adsorbates. The adsorption energies analysis showed the highest interaction AgCl-[6]Helicene. Density functional theory (DFT) simulations showed that chemical bonds take place during the adsorption of silver chloride on hexaHelicene which confirms that the [6]Helicene can function as a chiral molecular tweezer of the univalent cationic silver.

In recent years, several experimental and theoretical examinations have been carried out on Helicenes and related molecules including the determination of their racemization barriers\(^1\), aromatic character and non-linear optical properties\(^2\). In addition, recent works emerged the transition metal-based helicenes as novel attractive chiral molecules for various reasons. Firstly, metals are powerful templates for assembling \(\pi\)-conjugated ligands into well defined molecular structures; simply by using the fundamental ideas of coordination and organometallic chemistry\(^4\). Secondly, novel characteristic can be shown when metallic particles are grafted onto \(\pi\)-helical structures\(^5\). In particular, these chiral molecules are involved in several research fields such as molecular recognition\(^5\), supra-molecular chemistry\(^6\) and catalysis of transition metal\(^2\) because of their extraordinary geometry and their electronic properties\(^3\). Finally, the complexation of Helicenes by metals suggests the development of future devices in optoelectronics as chiral waveguides and we can think of planning new materials for the modulation of information in telecommunications\(^7\).

Moreover, It has been shown that thin layers of adsorbed ions (Li\(^+\), Mg\(^{2+}\), Fe\(^{3+}\), K\(^+\), Na\(^+\)... ) based functionalized heptaHelicene or macrocycle molecules (e.g porphyrins) can be obtained on solid support of quartz

\(^{1}\)Physics Department Rabigh College of Science and Arts, King Abdulaziz University, Jeddah, P.O box 344, Rabigh, 21911, Saudi Arabia. \(^{2}\)Laboratory of Quantum and Statistical Physics, LR18ES18, Faculty of Sciences of Monastir, Monastir, 5000, Tunisia. e-mail: mbinyhea@kau.edu.sa; ben_yahia_med@hotmail.fr
point, it should be noted that up to now, a cation-π complex with [6]Helicene has not been proven24.

we choose to use the hexaHelicene as adsorbent material seeking for formation of more stable complex. At this molecular tweezer but the bond is not strong and the complex formed is not stable. For this reason, in this paper, under various temperatures10–12. The initial success of this experimental approach has encouraged us to study the tal adsorption isotherms by identifying the complexed masses of metallic ions for a wide range of concentration on the electrode surface13. Recently, the quartz crystal microbalance has been demonstrated more powerful than the other methodologies34. It involves the spin coating of Helicene volume on the quartz electrode to cover the overlapping electrodes portion. The objective is to acquire appropriate films which should be homogeneous over the whole surface of the quartz crystal15,16. Films formed in this way have several virtues: first, they are mechanically stable and tolerant of different solvents and reaction conditions used for common organic transformations15; second, they are adequately well ordered that globally most of the complexing Helicenes are attractive candidates for this application because they have been demonstrated to be dynamic adsorbent in the complexation processes46. Interestingly, it should be noted that the QCM technique was devoted for the achievement of experimental adsorption isotherms by identifying the complexed masses of metallic ions for a wide range of concentration under various temperatures10–12. The initial success of this experimental approach has encouraged us to study the complexation of silver ions on hexaHelicene and to investigate the microscopic properties of the formed complex. Thus, it appears that the complex Ag⁺–Helicene can serve as a drug in the fight against cancer17 but the adsorption of silver ion on Helicenes does not seem to be significantly studied despite the importance of this complex in the medical fields. This motivated us to study the experimental (also theoretical) possibility of hexaHelicene acting as a complexing sensor of the silver ion (Ag⁺). Hence, a study of adsorption at the solid-liquid interface is required18.

In reality, hexaHelicene is an interesting member of the polycyclic aromatic hydrocarbon family called carbo-Helicenes19,20. It is characterized by its helical geometry offering rise to two enantiomers (Fig. 1). Six benzeneoid rings offer ascent to scaffolding that finishes a total turn of the helix. In addition, the terminal rings of these Helicenes appear to be cofacials. The profile of these molecules shows the two rings of benzene overlapping like jaws of a crocodile (Fig. 2). So, it is intriguing to check if this small molecule has the adequate flexibility to complex the cationic metals5,20.

Actually, several examinations indicated that the heptaHelicene can complex the alkali metals (K⁺, Na⁺, Cs⁺, Li⁺…)10,21,22. However, it is exhibited that the binding of the silver (I) particle in the focal cavity of [7]Helicene shows that the counter ion is feebly coordinated23. In such coupling mode, the [7]Helicene is working as a chiral molecular tweezer but the bond is not strong and the complex formed is not stable. For this reason, in this paper, we choose to use the hexaHelicene as adsorbent material seeking for formation of more stable complex. At this point, it should be noted that up to now, a cation-π complex with [6]Helicene has not been proven24.

The study of the insertion mode of alkali silver into [6]Helicene cavity is the subject of this work. Indeed, the initial segment aims to achieve the adsorption isotherms of silver chloride, silver nitrate and silver sulphide on [6] Helicene layers using the quartz crystal microbalance technique18–20. The second part is committed to the modeling of these isotherms using the adsorption models developed through statistical physics treatment11,12,21,22. This modeling permits the determination of the best adsorbate regarding its stability and reactivity in a practical industrial application through the comparison of the adsorption capacities. The choice of the adsorbate-adsorbent couple is also checked by applying the density functional theory for interaction energy calculation26. The application of the DFT method permits foreseeing the most plausible structure of the resulting complex26.

**Figure 1.** Representation of the two spatial conformations (enantiomers) M and P of hexa–Helicene molecule.
surface of the quartz crystal will be preciously identified by the microbalance apparatus\(^2^7\). In fact, this technique requires cleaning of the quartz crystal and then covering the gold electrode with thin layer of Helicene.

**Preparation of adsorption cell.** The commercial quartz crystal was protected by resin layer which should be removed using a well-known cleaning protocol\(^2^8\): the material was rinsed with acetone and deionizer water followed by drying. Then, it was cleaned with a Piranha solution at room temperature for 1 to 5 minutes. After this cleaning treatment, the crystals were completely rinsed with deionizer water and ethanol, and then dried by applying high purity nitrogen to remove any remaining water. Note that the crystals used in the experiments are polished with a fundamental resonance frequency of 5 MHz and have a diameter of 2.54 cm.

Then, a modification of the active surface of the crystal was carried out by coating the hexaHelicene adsorbent: 30 µl of adsorbent was deposited at 3000 rpm for 1 min onto the quartz crystal surface. The covered crystals were then dried at 100 °C for 2 hours. After drying, the resonance frequency of the crystal coated with the adsorbent was measured. The difference between the frequencies of the virgin and the coated crystal was calculated. By the intermediate of this frequency difference, we calculated the mass of the deposited adsorbate. The design of such chemically modified electrodes is to control if the Helicene molecules were capable of forming stable complexes with silver ions in the close proximity of the electrode surface. Once the adsorption cell is prepared, we proceed with the isotherms measurement.

**Measurement of experimental adsorption isotherms.** In the reactor filled with the buffer solution (\(V_i = 130 \text{ ml}\)), the Adsorption cell ([6]Helicene coated on quartz crystal) was immersed until the adjustment

---

**Figure 2.** Profile of hexaHelicene molecule trying to ‘devour’ a silver metal.

**Figure 3.** Experimental setup of Quartz Crystal Microbalance strategy devoted for the achievement of experimental adsorption isotherms of silver chloride, silver nitrate and silver sulphide on hexa-Helicene molecules.
of the frequency. Once the QCM probe was immersed in water, the hydrostatic pressure caused a change in the resonant frequency. This frequency \( f_0 \) was measured by a Maxtek frequency-meter PM700 and was taken as reference for our measurement.

Then, stock solutions of adsorbates AgCl, AgNO\(_3\) and \((\text{Ag})_2\text{S}\) were prepared. Adding a volume \( V_{ad} \) to the initial volume \( V_i \) permitted to obtain concentration of solutions ranging from \( 10^{-3} \text{M} \) to \( 1.6 \times 10^{-2} \text{M} \). If we note by \( c_0 = 10^{-4} \text{mol/l} \) the concentration of the stock solution of the adsorbate already prepared and we need to obtain a final concentration in the reactor \( c_f = 10^{-2} \text{M} \), the additional volume was determined as follows:

\[
V_{ad} = \frac{c_f}{c_0} \cdot V_i
\]

The added volumes for each concentration were calculated according to the last equation. So, volumes of adsorbate solution were injected using a micropipette in order to increase the adsorbate concentration in the reactor. For example, to obtain the concentration \( 10^{-2} \text{M} \), 100 \( \mu \text{l} \) of adsorbate solution \( 10^{-4} \text{M} \) was injected into the reactor followed by magnetic stirring for 5 min to homogenize the solution. After each addition, the microbalance apparatus displayed the resonance frequency associated with the new concentration. This step was repeated according to the number of performed injections.

Furthermore, it is necessary to quantify the quartz resonant frequency by assigning the different parameters to the corresponding frequency in order to correctly exploit the experimental results. In fact, once the sensor is immersed in a liquid, the model which describes the response of a quartz is called the Kanazawa-Gordon model whose change of oscillation frequency is affected by the properties of the liquid which are the density and the viscosity of the solution, by the temperature, by the hydrostatic pressure of the solution and by the roughness of the adsorbent surface\(^{29-32}\). Then, the total frequency is written as follows:

\[
\Delta f = f_r - f_0 = \Delta f_m + \Delta f_F + \Delta f_p + \Delta f_r + \Delta f_{\text{mT}}
\]

where, \( \Delta f_m \) describes the influence of the mass change, \( \Delta f_F \) is the temperature effect, \( \Delta f_p \) is the pressure effect, \( \Delta f_r \) is the roughness effect and \( \Delta f_{\text{mT}} \) represents the liquid properties effects described by Kanazawa which are due mainly to the variation of the viscosity and the density of the solution.

In our examination, the temperature, the pressure and the roughness were irrelevant because the experimental measurements were achieved in a thermostatic cell at fixed temperature using polished quartz. Moreover, low impacts of solution properties were noted so they were not taken into account\(^{31}\). It can be concluded that the frequency change was mainly due to the deposited mass of adsorbate \( \Delta f = \Delta f_m \). In this case, the adsorbed mass was determined from the Sauerbrey’s relation which assumed that for small masses changes, the added mass can be treated as an additional mass of the quartz\(^{33}\). Thus, the frequency-mass relation is described as follows\(^{33}\):

\[
\Delta f = -C_f \cdot \Delta m
\]

where, \( C_f \) is the Sauerbrey’s constant \((\text{Hz}.\text{cm}^2/\mu\text{g})\), \( \Delta f \) is the frequency shift (Hz) and \( \Delta m \) is the additional mass deposited on the quartz (\( \mu\text{g/cm}^2 \)).

We can finally deduce the complexed quantity per unit of surface for each concentration and afterward, we obtain the equilibrium adsorption isotherms of silver chloride, silver nitrate and silver sulfide onto hexaHelicene which are illustrated in Fig. 4 at three different temperatures.

**Results of QCM measurements.** In view of the experimental isotherms profiles, we note the following results:

Firstly, the experimental adsorption isotherms indicated that the silver particles are captured by the [6] Helicene molecules at various temperatures. Therefore, it is demonstrated that the [6]Helicene can function as a sensor of silver particles.

Secondly, one can see that the behaviors of isothem profiles are totally different for the three adsorbates, despite all experimental data were accomplished under the same conditions and using the same adsorbent (hexaHelicene). One can conclude that the isotherms profiles depend on the adsorbate type which constitutes the dominant factor of the complexation phenomenon. Moreover, by looking at the adsorbed amounts of the three adsorbates, we hold that silver chloride presents the highest adsorption capacities and in this way it was the best adsorbate for Helicenes complexation.

Thirdly, we note that an expansion in the temperature causes an increase in the adsorbed amount for all adsorption systems. So, the temperature effect indicates that the Helicene complexation is an endothermic process whatever the adsorbate is.

Finally, it is noticed that the silver chloride isotherms present one saturation level indicating that only one layer of cationic ions Ag\(^+\) is adsorbed onto the solid support of [6]Helicenes while the chlorine ions (Cl\(^-\)) stay as spectator ions in solution and don’t impose any impact on the complexation mechanism. At high adsorbate concentration, we can see a stable saturation state for the AgCl which means that all hexaHelicene sites are occupied and that the AgCl-[6]Helicene interaction is strong so cutting this link is almost impossible. For silver nitrate adsorption, one saturation level followed with a drop is observed at the three temperatures which suggests the presence of an irreversible phenomenon after the saturation state. This can be explained by the fact that the AgNO\(_3\)-[6]Helicene bond is weak so the silver ions can escape from the crocodile jaws ([6]Helicene cavity) at any time which leads to the decrease of the adsorbed quantity at high concentration. The isotherms profile of silver sulfide is totally different from the others. Two saturation states are shown for the adsorption isotherms of Ag,S on hexaHelicene which leads to estimate the formation of two adsorbed layers. So, the use of silver sulfide leads to conclude that layer by layer adsorption\(^{15,34}\) takes place and that the anionic sulfur ions have an impact.
on the complexation mechanism: the [6]Helicene acts as a complexing adsorbent of the silver Ag⁺ and after the formation of the first layer, the adsorption of the anionic ion S²⁻ takes place until the total saturation at high concentration.

According to these explanations, one can conclude that silver chloride is the suitable material for the fabrication of a stable complex silver-hexahelicene.

In the next section, statistical physics treatment is proposed for modeling analysis of adsorption isotherms. The goal is to elaborate a theoretical description of these experimental results at the ionic scale.

**Statistical Physics Analysis**

In this section, the microscopic investigation of the three complexation systems is carried out by adopting advanced statistical physics models. This approach has been recently used with great success in adsorption isotherms modeling.

**Development of advanced models.** The experimental isotherms indicated that the adsorption of silver chloride and silver nitrate take place with single-layer formation whereas, L.B.L double-layers formation is concluded for the silver sulfide adsorption. Moreover, the silver chloride and the silver sulfide isotherms show stable saturation levels with no disturbances. These isotherms can be interpreted by the classical single-layer and double-layers models which are established on the basis of the ideal gas approach. However, one saturation level followed with a drop is observed for the silver nitrate isotherms. According to previous studies, this decrease in the adsorbed quantity after saturation is mainly due to the lateral interactions between the adsorbate particles at free state. This description requires the use of the chemical potential of real gas in which the lateral interactions are introduced.

Actually, the development of adsorption models using the statistical physics treatment requires the following assumptions:

1. The use of the grand canonical ensemble is necessary to assess the variation of adsorbed ions number in relation with the concentration.
2. Only the translation degrees of freedom are taken into account. All the other internal degrees of the particle (rotational, vibrational, electronic and nuclear) are neglected because the investigated metallic ion in this paper is monoatomic (Ag⁺).

**Figure 4.** Complexed quantity per unit of surface area $Q$ (µg.cm⁻²) describing the adsorption of silver chloride, silver nitrate and silver sulfide on hexa-Helicene molecules given at 293, 313 and 333 K.
It is known that the [6]Helicene has two spatial conformations \( M \) and \( P \) (Fig. 1)\(^\text{13}\). So, it is assumed that the receptor sites of the adsorbent are divided into two types and that a self-adsorption competition of the silver metal takes place between the two kinds of receptor sites, each with its relating energy level. Therefore, the statistical physics models developed in this paper are the single-layer model with two energy levels and the double-layers model on two types of sites.

**Single-layer adsorption.** For the single-layer adsorption model, it is assumed that the silver particles followed a monolayer adsorption process. They are adsorbed on the first adsorbent type \( H_{M1} \) with the first energy \((-\varepsilon_1)\) while the adsorption on the density \( H_{M2} \) of the second type of sites is carried out with the second energy level \((-\varepsilon_2)\). Then, the partition-function of the grand-canonical ensemble of Gibbs is written as\(^\text{21,35}\):

\[
z_{1gc} = \sum_{N_{1}=0,1} e^{-\beta(-\varepsilon_1 - \mu_1)} N_{1} = 1 + e^{-\beta(-\varepsilon_1 - \mu_1)}
\]

\[
z_{2gc} = \sum_{N_{2}=0,1} e^{-\beta(-\varepsilon_2 - \mu_2)} N_{2} = 1 + e^{-\beta(-\varepsilon_2 - \mu_2)}
\]

With \(z_{1gc}\) and \(z_{2gc}\) being the partition-functions of the two kinds of sites, \(\mu_1\) and \(\mu_2\) being the chemical potentials of adsorbed ions and \(\beta\) being the Boltzmann factor.

The next step of the statistical physics development comprises the calculation of the average numbers of occupation \(N_{1j}\) and \(N_{2j}\) of independent [6]Helicene sites. They are expressed as follows\(^\text{36,37}\):

\[
N_{1j} = \frac{H_{M1}}{1 + e^{-\beta(\varepsilon_1 + \mu_1)}}
\]

\[
N_{2j} = \frac{H_{M2}}{1 + e^{-\beta(\varepsilon_2 + \mu_2)}}
\]

For a classical single-layer adsorption with no disturbances at saturation, the mutual interactions between the adsorbate particles are neglected and then, we use the chemical potential of the ideal gas. Whereas, in the case of saturation level followed with a drop, we use the chemical potential of the real gas in order to introduce the lateral interactions between the adsorbate particles.

Finally, we calculate the analytical expression of the adsorbed quantity (Q) corresponding to the single-layer model with two types of sites resulting from the product of the two numbers of ions per sites \(n_{1j}\) and \(n_{2j}\) and the two average occupation numbers \(N_{01}\) and \(N_{02}\)\(^\text{21,37}\):

\[
Q = n_{1j} N_{01} + n_{2j} N_{02}
\]

The single-layer model expression established on the basis of the perfect gas (SMPG) is written as:

\[
Q_{SMPG} = \frac{n_{1j}H_{M1}}{1 + \left(\frac{c_1}{7}\right)^{n_{1j}}} + \frac{n_{2j}H_{M2}}{1 + \left(\frac{c_2}{7}\right)^{n_{2j}}}
\]

Where, \(n_{1j}\) and \(n_{2j}\) are the numbers of bonded ions per Helicene site. \(H_{M1}\) and \(H_{M2}\) are the densities of the two spatial conformations \(M\) and \(P\) of hexaHelicene. \(c_1 = Sc^{-361}\) and \(c_2 = Sc^{-362}\) are the energetic parameters.

The expression of the single-layer adsorbed quantity developed on the basis of the real gas law (SMRG) is written as follows:

\[
Q_{SMRG} = \frac{n_{1j}H_{M1}}{1 + \left(\frac{w_1 - bc}{c} e^{2aic} e^{-\frac{\varepsilon_1}{kT}}\right)^{n_{1j}}} + \frac{n_{2j}H_{M2}}{1 + \left(\frac{w_2 - bc}{c} e^{2aic} e^{-\frac{\varepsilon_2}{kT}}\right)^{n_{2j}}}
\]

This advanced model presents eight physicochemical parameters: the steric counting parameters \(n_{1j}\) and \(n_{2j}\), \(H_{M1}\) and \(H_{M2}\), the two energetic parameters \(w_{1}\) and \(w_{2}\) and the two lateral interactions parameters which are the cohesion pressure \(a\) and the covolume \(b\).

**L.B.L Double-layers adsorption.** The L.B.L double-layers adsorption is shown in the case of silver sulfide isotherms. In order to develop the systematic expression of this model, four energy levels were assumed:

- \((-\varepsilon_3)\) characterizes the adsorption of silver ions of the first layer on the first adsorbent type.
- \((-\varepsilon_4)\) reflects the adsorption of silver ions on the second spatial conformation of hexaHelicene.
- \((-\varepsilon_5)\) is the energy level corresponding to the adsorption of sulfur ions of the second layer on the first spatial conformation.
- \((-\varepsilon_6)\) is the energy level of sulfur adsorption on the second adsorbent type.
According to this supposition, the grand-canonical partition function of the L.B.L double-layers model on two
types of sites is composed as follows:

\[ z_{gc} = \sum_{N=0,1,2} e^{-β(−e_1−μ_1)N} = 1 + e^{β(e_1+μ_1)} + e^{β(e_1+e_2+2μ_1)} \]  

(11)

\[ z_{dc} = \sum_{N=0,1,2} e^{-β(−e_1−μ_2)N} = 1 + e^{β(e_2+μ_2)} + e^{β(e_2+e_4+2μ_2)} \]  

(12)

Following the statistical physics treatment described previously, we deduce the two analytical expressions Q of
the L.B.L double-layers model with four energy levels:

Based on ideal gas approach:

\[ Q_{DMPG} = n_1H_{M1} \left( \frac{c_{n1}}{1 + \left( \frac{c_{n1}}{c_{21}} \right)^2} \right)^{2n1} + n_2H_{M2} \left( \frac{c_{n2}}{1 + \left( \frac{c_{n2}}{c_{21}} \right)^2} \right)^{2n2} \]  

(13)

Based on real gas law:

\[ Q_{DMRG} = n_1H_{M1} \left( \frac{c_{w1}}{w_1(1−bc)α_{12}e^{2\alpha_a} τ_{FE}} \right)^{n1} + n_2H_{M2} \left( \frac{c_{w2}}{w_1(1−bc)α_{12}e^{2\alpha_a} τ_{FE}} \right)^{n2} \]  

(14)

\[ c_{1,2,3,4} = Se^{−βc_{1,2,3,4}} \]  

(15)

The coefficients \( c_{1,2,3,4} \) and \( w_{1,2,3,4} \) are energetic parameters written as function of adsorption energies:

\[ w_{1,2,3,4} = Se^{−βc_{1,2,3,4}} \]  

(16)

These two models present common steric parameters: \( n_1, n_2, H_{M1} \) and \( H_{M2} \). Each of them presents four ener-
getic factors which are \( c_1, c_2, c_3 \) and \( c_4 \) for the DMPG model and \( w_1, w_2, w_3 \) and \( w_4 \) for the DMRG model. The
DMRG model presents also two Van-der-Waals parameters (the cohesion pressure \( a \) and the covolume of particles \( b \)).

**Numerical results.** Numerical fitting program is applied to adjust the four models SMPG, SMRG, DMPG and
DMRG with all complexation isotherms.

Note that the results demonstrating that an adsorption isotherm is well fitted with statistical physics model
are: the correlation coefficient \( R^2 \) tends to the unit, the values of the residual root mean square \( RMSE \) are less than
3 and the fitting values of the Akaike information criterion \( AIC \) are the lowest quantities\(^{12,38}\)

The values of errors coefficients of adjustment (\( R^2 \), \( RMSE \) and \( AIC \)) are reported in Table 1.

According to Table 1, it is confirmed that the experimental isotherms of silver chloride are well adjusted by
the single-layer adsorption model (SMPG). The SMRG is found in acceptable correlation with the silver nitrate
adsorption. The L.B.L double-layers model with four energies (DMPG) is selected for the interpretation of the
silver sulfide complexation.

These numerical results are in accordance with the experimental findings given in section II.3: silver chloride
is confirmed to be the best adsorbate for hexaHelicene complexation because the anionic ions (Cl\(^−\)) do not influence
the adsorption mechanism. Moreover, during the silver chloride complexation there are no disturbances at
saturation which reflects the highest stability of the formed complexes.

The values of the physicochemical parameters are also deduced through the numerical adjustment. By the
intermediate of the models parameters values, the complexation mechanisms can be deciphered in order to select
the best adsorbate for [6]Hericene complexation.

**Results of statistical physics modeling.** Table 2 indicates the fitting values of all parameters affecting the
single-layer complexation of silver chloride and silver nitrate and the double-layers adsorption of silver sulfide.

It is necessary to comprehend the physical effect of all models parameters on the adsorption capacities in order
to compare the materials used for [6]Hericene complexation with microscopic manner:
The adsorption energy calculation is the most significant factor frequently applied to characterize the interaction type of adsorption system. The single layer model SMPG evolves two energetic parameters $E_1$ and $E_2$ which participate at the double-layers adsorption and prevents the Helicenes complexation. The presence of these parameters in the SMRG model (which depicts the silver nitrate adsorption on [6]Helicene) indicates high adsorbate-adsorbate interaction. This means that the attractive interaction between the silver ions and the solution is greater than the interaction with the adsorbing surface which leads to a reversible phenomenon at high concentration. Actually, the reversible phenomenon shown in silver nitrate adsorption isotherms (Fig. 4) is probably due to the weak binding $\text{AgNO}_3$-[6]Helicene. This deduct confirm the deduction from the numerical adjustment of experimental hexa-Helicene adsorption isotherms with the four statistical physics models (SMPG, SMRG, DMPG and DMRG).

- Generally, the higher the value of $n_1$ or $n_2$ is, the greater the adsorbed amount is since this parameter represents the number of particles per adsorbent site and it can estimate the accumulation level of silver particles on one site. So, a value of $n$ greater than one implies that it is easier to gather many silver particles on one [6]Helicene site. Undoubtedly, the adsorption system with the highest values of $n_1$ and $n_2$ is the most appropriate for [6]Helicene complexation in term of reproducibility. According to Table 2, we can clearly see that: $n_1(\text{AgCl}) > n_1(\text{AgNO}_3) > n_2(\text{AgCl}) > n_2(\text{AgNO}_3)$ and $n_1(\text{AgCl}) > n_1(\text{Ag}_2\text{S}) > n_1(\text{Ag})$. This means that the silver chloride is the most attractive with the hexa-Helicene sites in term of quantity. Besides, all adjusted values of $n_1$ and $n_2$ are inferior to one. So it can be concluded that the multi-ionic adsorption into the hexa-Helicene sites is impossible because of the electrostatic repulsion interaction between the silver ions. The fitted values of $n_1$ or $n_2$ are the lowest for silver sulfide because sulfur ions have an impact on the complexation process. The interaction between the first adsorbed layer of $\text{Ag}^+$ and the second layer of $\text{S}^2-$ disfavors the interaction $\text{Ag}^+\text{-}[6]\text{Helicene}$.

- The densities of receptor sites $H_{M1}$ and $H_{M2}$ are steric parameters associated with the effectively occupied [6]Helicene sites ($P$ and $M$). So, it is evident to note that the adsorbed quantity effectively increases as $H_{M1}$ and $H_{M2}$ values increase. As indicated in Table 2, the fitted values of $H_{M1}$ and $H_{M2}$ are the highest in the case of the silver chloride adsorption: $H_{M1}(\text{AgCl}) > H_{M1}(\text{AgNO}_3) > H_{M1}(\text{Ag}_2\text{S})$ and $H_{M2}(\text{AgCl}) > H_{M2}(\text{AgNO}_3) > H_{M2}(\text{Ag}_2\text{S})$. It is noticed that the two types of [6]Helicene sites are more attractive with silver chloride in accordance with the previous investigation of $n_1$ and $n_2$. In fact, the chlorine ions have no influence on the complexation procedure so there is a quick penetration of $\text{Ag}^+$ in the adsorbent pores contrary to the sulfur ions $\text{S}^2-$ which participate at the double-layers adsorption and prevents the Helicenes complexation.

- The cohesion pressure $a$ and the covolume $b$ are responsible of the instability of the complexes formed during the adsorption of silver nitrate on [6]Helicene. They describe the lateral interactions between the adsorbate particles at free state$^{16,11,21}$. The presence of these parameters in the SMRG model (which depicts the silver nitrate adsorption) indicates high adsorbate-adsorbate interaction. This means that the attractive interaction between the silver ions and the solution is greater than the interaction with the adsorbing surface which leads to a reversible phenomenon at high concentration. Actually, the reversible phenomenon shown in silver nitrate adsorbers isotherms (Fig. 4) is probably due to the weak binding $\text{AgNO}_3$-[6]Helicene. This deduct confirm the choice of silver chloride as best material for Helicenes adsorption but it requires confirmation through energetic investigation. Furthermore, we can see from Table 2 that the Van-der-Waals parameters $a$ and $b$ decline with ascending of temperature. This result indicates that the lateral interactions between the adsorbates and the disturbances during the complexation process are the lowest at high temperature. This leads to estimate that the silver nitrate complexation can be carried out if we increase the temperature. This fact cannot be achieved by means of the QCM setup because the gold electrode of quartz crystal is very sensitive to high temperatures.

- The adsorption energy calculation is the most significant factor frequently applied to characterize the interaction type of adsorption system. The single layer model SMPG evolves two energetic parameters $\epsilon_1$ and $\epsilon_2$. Thus the complexation energies ($-\Delta E_1$) and ($-\Delta E_2$) which correspond to the complexation of silver chloride on the two spatial conformations of hexaHelicene are calculated by the following relation:

$$-\Delta E_{1,2} = RT \ln (c_{1,2}/S(\text{AgCl}))$$

where, $R$ is the ideal gas constant ($8.3144621 \text{ mol}^{-1} \text{ K}^{-1}$), $T$ is the absolute temperature, $S(\text{AgCl})$ is the solubility of investigated adsorbate (AgCl) and $c_1$ and $c_2$ are the fitted value energetic parameters deduced from the adjustment with SMPG Model.

The SMRG model gives rise to two energetic parameters $w_1$ and $w_2$. Then, the complexation energies ($-\Delta E_1$) and ($-\Delta E_2$) of silver nitrate on the two kinds of hexaHelicene are calculated by the subsequent relation:

| Adsorption isotherms | AgCl-[6]Helicene | AgNO₃-[6]Helicene | Ag₂S-[6]Helicene |
|---------------------|------------------|------------------|------------------|
| Temperature(K)      | 293              | 313              | 293              |
| $R^2$               | 0.99             | 0.99             | 0.88             |
| RMSE                | 0.04             | 0.05             | 0.52             |
| AIC                 | 6.5              | 6.9              | 14.2             |
| $\text{SMPG}$       |                  |                  |                  |
| $\text{SMRG}$       |                  |                  |                  |
| $\text{DMPG}$       |                  |                  |                  |
| $\text{DMRG}$       |                  |                  |                  |

Table 1. The values of the correlation coefficient $R^2$, the residual root mean square coefficient RMSE and the Akaike information criterion AIC deduced from the numerical adjustment of experimental hexa-Helicene adsorption isotherms with the four statistical physics models (SMPG, SMRG, DMPG and DMRG).

$\text{SMPG}$

$\text{SMRG}$

$\text{DMPG}$

$\text{DMRG}$
\[ -\Delta E_{1,2} = RT \times \ln \left( \frac{S(\text{AgNO}_3)}{w_1,2} \right) \] (18)

With, \( S(\text{AgNO}_3) \) is the solubility \( \text{AgNO}_3 \). The fitted value of the energetic parameters \( w_1 \) and \( w_2 \) are deduced from the adjustment of experimental isotherms with the SMRG Model.

The L.B.L double-layers adsorption of silver sulfide offers four adsorption energies by the intermediate of the DMPG model. They are calculated by the following formula:

\[ -\Delta E_{1,2,3,4} = RT \times \ln \left( \frac{S(\text{AgS}_2)}{w_1,3,4} \right) \] (19)

With \( S(\text{AgS}_2) \) is the silver sulfide solubility.

The adsorption energies of the three complexation systems determined at three temperatures are given in Table 3.

For silver sulfide adsorption, it should be noted that \((-\Delta E_1)\) and \((-\Delta E_2)\) reflects the silver particles adsorption at the level of the first adsorbed layer i.e. they characterize the \( \text{Ag}^+ \cdot [6] \text{Helicene} \) interaction. However, \((-\Delta E_3)\) and \((-\Delta E_4)\) are related to the adsorption of the second layer of sulfur ions\(^{1,26}\). Therefore, it is clear that \((-\Delta E_3)\) and \((-\Delta E_4)\) are not useful for the comparison of the formed complexes and then, the choice of the best adsorbate for hexaHelicene complexation depends on the comparison of \((-\Delta E_1)\) and \((-\Delta E_2)\) of the three systems.

In view of Table 3, the adsorptions energies are ordered as follows:

| Physic-chemical variables | 293 (K) | 313 (K) | 333 (K) |
|-------------------------|--------|--------|--------|
| \( n_1 \)              | 0.91   | 0.95   | 1.01   |
| \( n_2 \)              | 0.96   | 0.94   | 0.99   |
| \( H_{sc} (\text{Ag cm}^{-2}) \) | 293    | 313    | 333    |
| \( H_{sc} (\text{Ag cm}^{-2}) \) | 293    | 313    | 333    |
| \( a (10^{-20} \text{J.mL.mol}^{-1}) \) | 293    | 313    | 333    |
| \( b (10^{-10} \text{mL.mol}^{-1}) \) | 293    | 313    | 333    |
| \( c_1 (\text{mol}^{-1}) \) | 293    | 313    | 333    |
| \( c_2 (\text{mol}^{-1}) \) | 293    | 313    | 333    |
| \( c_3 (\text{mol}^{-1}) \) | 293    | 313    | 333    |
| \( c_4 (\text{mol}^{-1}) \) | 293    | 313    | 333    |
| \( w_1 (\text{mol}^{-1}) \) | 293    | 313    | 333    |
| \( w_2 (\text{mol}^{-1}) \) | 293    | 313    | 333    |

Table 2. Fitting values of physic-chemical variables affecting the single-layer complexation of silver chloride and silver nitrate and the double-layers adsorption of silver sulfide given at 293, 313 and 333 K.

---

The L.B.L double-layers adsorption of silver sulfide offers four adsorption energies by the intermediate of the DMPG model. They are calculated by the following formula:

\[ -\Delta E_{1,2,3,4} = RT \times \ln \left( \frac{S(\text{AgS}_2)}{w_1,3,4} \right) \] (19)

With \( S(\text{AgS}_2) \) is the silver sulfide solubility.

The adsorption energies of the three complexation systems determined at three temperatures are given in Table 3.

For silver sulfide adsorption, it should be noted that \((-\Delta E_1)\) and \((-\Delta E_2)\) reflects the silver particles adsorption at the level of the first adsorbed layer i.e. they characterize the \( \text{Ag}^+ \cdot [6] \text{Helicene} \) interaction. However, \((-\Delta E_3)\) and \((-\Delta E_4)\) are related to the adsorption of the second layer of sulfur ions\(^{1,26}\). Therefore, it is clear that \((-\Delta E_3)\) and \((-\Delta E_4)\) are not useful for the comparison of the formed complexes and then, the choice of the best adsorbate for hexaHelicene complexation depends on the comparison of \((-\Delta E_1)\) and \((-\Delta E_2)\) of the three systems.

In view of Table 3, the adsorptions energies are ordered as follows:

| Physic-chemical variables | 293 (K) | 313 (K) | 333 (K) |
|-------------------------|--------|--------|--------|
| \( n_1 \)              | 0.91   | 0.95   | 1.01   |
| \( n_2 \)              | 0.96   | 0.94   | 0.99   |
| \( H_{sc} (\text{Ag cm}^{-2}) \) | 293    | 313    | 333    |
| \( H_{sc} (\text{Ag cm}^{-2}) \) | 293    | 313    | 333    |
| \( a (10^{-20} \text{J.mL.mol}^{-1}) \) | 293    | 313    | 333    |
| \( b (10^{-10} \text{mL.mol}^{-1}) \) | 293    | 313    | 333    |
| \( c_1 (\text{mol}^{-1}) \) | 293    | 313    | 333    |
| \( c_2 (\text{mol}^{-1}) \) | 293    | 313    | 333    |
| \( c_3 (\text{mol}^{-1}) \) | 293    | 313    | 333    |
| \( c_4 (\text{mol}^{-1}) \) | 293    | 313    | 333    |
| \( w_1 (\text{mol}^{-1}) \) | 293    | 313    | 333    |
| \( w_2 (\text{mol}^{-1}) \) | 293    | 313    | 333    |

Table 2. Fitting values of physic-chemical variables affecting the single-layer complexation of silver chloride and silver nitrate and the double-layers adsorption of silver sulfide given at 293, 313 and 333 K.

---
hexaHelicene was determined using the next expression:

\[ \Delta E = E(\text{complex}) - (E(\text{Ag}) + E(\text{hexaHelicene})) \]  

where, \( E(\text{complex}) \) is the pure electronic energy of the formed complex Ag- [6]Helicene, \( E(\text{Ag}) \) is the estimated energy for the silver metal and \( E(\text{hexaHelicene}) \) is the energy of the molecular structure of the [6]Helicene surface.

The interaction energy, \( \Delta E \), for the connections between the silver particles and the adsorbent surface of hexaHelicene was determined using the next expression:

| Molar adsorption energy (kJ mol\(^{-1}\)) | Temperature (K) | AgCl-[6]Helicene | AgNO\(_3\)-[6]Helicene | AgS-[6]Helicene |
|----------------------------------------|---------------|-----------------|-----------------|----------------|
| \(-\Delta E_1\)                        | 293           | 88.4            | 43.6            | 33.6           |
|                                        | 313           | 95.4            | 48.6            | 39.4           |
|                                        | 333           | 111.2           | 50.7            | 44.1           |
| \(-\Delta E_2\)                        | 293           | 77.9            | 38.4            | 29.4           |
|                                        | 313           | 91.2            | 40.2            | 33.2           |
|                                        | 333           | 98.4            | 49.6            | 38.4           |
| \(-\Delta E_3\)                        | 293           | —               | —               | 21.3           |
|                                        | 313           | —               | —               | 18.4           |
|                                        | 333           | —               | —               | 16.5           |
| \(-\Delta E_4\)                        | 293           | —               | —               | 20.4           |
|                                        | 313           | —               | —               | 18.5           |
|                                        | 333           | —               | —               | 16.2           |

Table 3. Estimated values of adsorption energies of the three complexation systems determined at three temperatures (293–333 K).

Density functional theory (DFT) calculation and results. In order to elaborate the complexation characteristics of silver on hexaHelicene, theoretical calculations were carried out at the density functional level of theory (DFT, B3LYP functional)\(^{36,39}\), employing the Gaussian 09 suite of programs\(^{39}\). The LanL2DZ basis set was utilized, and the optimizations were unconstrained. DFT simulations were performed to estimate the bond distances associated with the silver metal adsorption and the subsequent binding energies of the formed complex.

Actually, the proposed structure of the silver-[6]Helicene complex was anticipated on the basis of the profound conformational analysis i.e., six distinctive mutual positions of the [6]Helicene ligand and the cationic silver were considered during the geometry optimization\(^{36,37}\). In the model calculations, we optimized the molecular geometries of the free [6]Helicene ligand and its complex with silver particle. The structure of the mentioned [6]Helicene used in DFT calculation is depicted in Fig. 5a.

The interaction energy, \( \Delta E \), for the connections between the silver particles and the adsorbent surface of hexaHelicene was determined using the next expression:

\[ \Delta E = E(\text{complex}) - (E(\text{Ag}) + E(\text{hexaHelicene})) \]  

where, \( E(\text{complex}) \) is the pure electronic energy of the formed complex Ag-[6]Helicene, \( E(\text{Ag}) \) is the estimated energy for the silver metal and \( E(\text{hexaHelicene}) \) is the energy of the molecular structure of the [6]Helicene surface.

Results of DFT simulations are illustrated in Fig. 5b. From Fig. 5b, in the resulting Ag-[6]Helicene complex, the silver metal is bound by four relatively strong bonds to four carbon atoms from the two terminal benzene rings of the parent ligand via cation-π interaction. The bonds lengths ranged from 1.91 Å to 2.03 Å. In addition, we note an interaction energy value of \(-98.3\,\text{kJ/mol}\) which also confirms the formation of this complex. The estimated binding energy corresponded to covalent bonds which could be associated to chemisorption mechanism. This finding is consistent with the results obtained with the SMPG adsorption model (silver chloride adsorption) based on statistical physics approach.

Consequently, the DFT investigation has proven that high interaction takes place during the adsorption of silver chloride on hexaHelicene.

Conclusion and Recommendation
The aim of this work was to describe a cancer drug at the ionic scale. The ultimate result is that hexaHelicene can function as sensor of the cationic silver metal. Overall, despite all experimental isotherms are carried out by the intermediate of the QCM method and describe the same chemical process ([6]Helicene complexation), three different phenomena are observed. The adsorption of silver chloride is explained via the single-layer model on two types of sites SMPG, the silver nitrate adsorption is discussed by the intermediate of the SMRG model which takes into account the lateral interaction between the particles and the silver sulfide isotherms are in good correlation with the L.B.I. adsorption model DMPG.
The patterns of the three complexation frameworks are discussed by comparing the fitted parameters of the three advanced statistical physics models. The parameters numbers of silver ions per Helicene site ($n_1$ and $n_2$) demonstrated that the silver chloride is the best interactive material with hexaHelicene sites in term of quantity and that no accumulation occurs on one receptor site during the complexation mechanisms ($n_{1,2} < 1$). The densities of occupied sites $H_{\text{HM1}}$ and $H_{\text{HM2}}$ indicated that the two types of [6]Helicene sites P and M are more attractive with the silver chloride. The calculation of the complexation energies and the DFT optimizations confirm the highest stability of the AgCl-[6]Helicene complex. Actually, the interaction AgCl-[6]Helicene is found typical to a chemisorption process.

Correlating all QCM measurements and statistical physics discussions, we recommend the use of silver chloride as adsorbate material for a modern application of Helicenes complexation.

**Comparison with similar works.** Theoretically speaking, similar previous works on [7]Helicene (e.g. lithium-heptaHelicene and Potassium-heptaHelicene...)\textsuperscript{10,21} showed that the presence of chlorine ions has no impact on the complexation process which is in accordance with the case of silver chloride. However, the mentioned previous complexes are explained by a monolayer model that presents disturbances at saturation indicating that the complexes formed with heptaHelicene are unstable like the case of silver nitrate in the present investigation. This confirms that the use of hexaHelicene as complexing adsorbent is more successful and conceptive than the use of heptaHelicene. Moreover, despite the silver sulfide-hexaHelicene was found a stable complex, the L.B.L adsorption on Helicenes molecules is shown at the first time in this investigation. Thus, past works have demonstrated that the adsorption of cationic metals (Mg$^{2+}$, Zn$^{2+}$, and Fe$^{2+}$...) on porphyrins was carried out on the basis of the layer by layer approach in the case of nitrate ions presence\textsuperscript{11,12}. So, we can notice that an L.B.L adsorption on Helicenes can be only shown in the case of using an adsorbate involving sulfur ions because silver nitrate complexation resulted in a single-layer adsorption which indicated that the nitrate ions were not involved in the adsorption mechanism.

Received: 7 April 2020; Accepted: 28 May 2020;
Published online: 25 June 2020

**References**

1. Grimme, S. & Peyerimhoff, S. D. Theoretical study of the structures and racemization barriers of [n]helicenes (n = 3-6, 8). *J. Chem. Phys.* **204**, 411–417 (1996).
2. Schulman, J. M. & Disch, R. L. Theoretical Studies of the [N]Phenylenes. *J. Am. Chem. Soc.* **118**, 8470–8474 (1996).
3. Žádný, J. et al. Exploration of 9-bromo[7]helicene reactivity. *Tetrahedron* **69**, 6213–6218 (2013).
4. Elschenbroich, C. *Organometallics 3rd, Completely Revised and Extended Edition*, Wiley VCH, Munich (2006).
5. Janke, R. H., Haufe, G., Würthwein, E. U. & Borkent, J. H. Racemization Barriers of Helicenes: A Computational Study. *J. Am. Chem. Soc.* **118**, 6031–6035 (1996).
6. Nakai, Y., Mori, T. & Inoue, Y. Theoretical and Experimental Studies on Circular Dichroism of Carbo[n]helicenes. *J. Phys. Chem. A* **116**, 7372–7385 (2012).
7. Furche, F. et al. Circular Dichroism of Helicenes Investigated by Time-Dependent Density Functional Theory. *J. Am. Chem. Soc.* **122**, 1717–1724 (2000).
8. Xu, Y. et al. (P)-helicene displays chiral selection in binding to Z-DNA. *J. Am. Chem. Soc.* **126**, 6566–6567 (2004).
9. Wang, D. Z. G. & Katz, T. J. A [5]HELOL Analogue That Senses Remote Chirality in Alcohols, Phenols, Amines, and Carboxylic Acids. *J. Org. Chem.* **70**, 8497–8502 (2005).
10. Ben Yahia, M. et al. Adsorption of sodium and lithium ions onto helicenes molecules: Experiments and phenomenological modeling. J. Mol. Liq. 288, 110988 (2019).

11. Knani, S., Khalifa, N., Ben Yahia, M., Aouaini, F. & Tounsi, M. Statistical physics study of the interaction of the 5, 10, 15, 20-tetrakis(4-tolylphenyl) porphyrin (H2TTPP) with magnesium ion: New microscopic interpretations, Arab. J. Chem. 13, 4374–4385 (2020).

12. Ben Yahia, M. et al. Statistical studies of adsorption isotherms of iron nitrate and iron chloride on a thin layer of porphyrin. J. Mol. Liq. 248, 235–245 (2017).

13. Ballarin, B., Masiero, S., Seeger, R. & Tonelli, D. Modification of electrodes with porphyrin-functionalised conductive polymer. J. Electroanal. Chem. 449, 173–180 (1998).

14. Sener, G. et al. Quartz crystal microbalance based nanosensor for lysozyme detection with lysozyme imprinted nanoparticles. J. Biosens. Bioelectron. 26, 815–821 (2010).

15. Korposh, S., Selyanchyn, R. & Lee, S. W. Nano-assembled thin film gas sensors. IV. Mass-sensitive monitoring of humidity using quartz crystal microbalance (QCM) electrodes. Sens. Actuat. B. 147, 599–608 (2010).

16. Nuckolls, C. et al. Electro-Optic Switching by Helicene Liquid Crystals. J. Chem. Mater. 14, 773–776 (2002).

17. Huang, H., Zhou, J., Chen, S., Zeng, L. & Huang, Y. A highly sensitive QCM sensor coated with Ag+–ZSM-5 film for medical diagnosis. Sens. Actuat. B. 101, 316–321 (2004).

18. Johansson, M. P. & Patzschke, M. Fixing the Chirality and Trapping the Transition State of Helicene with Atomic Metal Glue. Chem. Eur. J. 15, 13210–1321 (2009).

19. Saleh, N., Shen, C. & Cassouss J. Helicene-based transition metal complexes: synthesis, properties and applications. Chem. Sci. 5, 3680–3694 (2014).

20. Dougherty, D. A. Cation–π Interactions in Chemistry and Biology: A New View of Benzene, Phe, Tyr, and Trp. Science. 271, 163–168 (1996).

21. Ben Yahia, M. et al. Study of the Interaction of [7]-Helicene with Alkaline Cations (K+ and Cs+): New Insights on Microscopic Adsorption Behavior. RSC Adv. 7, 44712–44723 (2017).

22. Knani, S., Mabrouk, N., Tounsi, M. & Yahia, M. B. Statistical modeling of adsorption isotherm of potassium onaza[7]helicene-coated gold electrode attached to quartz crystal microbalance. Sep. Sci. Techn. 54, 2386–2396 (2018).

23. Vijay, D. & Sastry, G. N. Exploring the size dependence of cyclic and acyclic π-systems on cation–π binding†. Phys. Chem. Chem. Phys. 10, 582–590 (2008).

24. Fuchter, M. J. et al. [7]-Helicene: a chiral molecular tweezer for silver(I) salts. J. Dalton Trans. 41, 8238–8241 (2012).

25. Di Natale, C. et al. The exploitation of metalloporphyrins as chemically interactive material in chemical sensors. Mater. Sci. Eng. C. 5, 209–215 (1998).

26. Sellouli, D. et al. Understanding the adsorption of Pb2+, Hg2+ and Zn2+ from aqueous solution on a lignocellulosic biomass char using advanced statistical physics models and density functional theory simulations. Chem. Eng. J. 365, 305–316 (2019).

27. O’Sullivan, C. K. & Guibault, G. G. Commercial quartz crystal microbalances theory and applications, Biosens. Bioelectron. 14, 663–670 (1999).

28. Köflinger, C. et al. A quartz crystal biosensor for measurement in liquids. Biosensors and Bioelectronics. 7, 397–404 (1992).

29. Wang, D., Moussavi, P., Hauser, P. J., Oxenham, W. & Grant, C. S. Quartz crystal microbalance in elevated temperature viscous liquids: Temperature effect compensation and lubricant degradation monitoring. Colloids and Surfaces A: Physicochem. Eng. Aspects. 268, 30–39 (2005).

30. Park, K., Koh, M., Yoon, C., Kim, H. & Kim, H. The behavior of quartz crystal microbalance in high pressure CO2. J. Super. Fluids. 29, 203–212 (2004).

31. Rocklein, M. N. & George, S. M. Temperature-Induced Apparent Mass Changes Observed during Quartz Crystal Microbalance Measurements of Atomic Layer Deposition. Anal. Chem. 75, 4975–4982 (2003).

32. Urbakh, M. & Daikhist, L. Influence of the Surface Morphology on the Quartz Crystal Microbalance Response in a Fluid. Langmuir. 10, 2836–2841 (1994).

33. Sauerbrey, G. Use of quartz vibration for weighing thin films of a microbalance. Z. Phys. 155, 206–212 (1959).

34. Lvov, Y., Ariga, K., Ichinose, I. & Kunitake, T. Assembly of multicomponent protein films by means of electrostatic layer-by-layer adsorption. J.Am.Chem.Soc. 117, 617–6223. (1995).

35. Aouaini, F., Knani, S., Ben Yahia, M. B. Formation of the advanced statistical physics models in order to interpret the complexation phenomenon at the microscopic scale.

36. Marquardt, W. D. An algorithm for least-squares estimation of nonlinear parameters. Chem. Phys. Lett. 316, 106–114 (2002).

37. Ben Lamine, A. & Bouazra, Y. Application of statistical thermodynamics to the olfaction mechanism. Chem. Senses. 22, 67–75 (1997).

38. Markwardt, W. D. An algorithm for least-squares-estimation of nonlinear parameters. J. Soc. Ind. Appl. Math. 11, 431–441 (1963).

39. Karabacak, M. et al. DFT calculations and experimental FT-IR, FT-Raman, NMR, UV-Vis spectral studies of 3-fluorophenylboronic acid, Spectrochim. Acta. Part A: Mol. Biomol. Spectr. 136, 306–320 (2015).

40. Chen, D. M., Liu, X., He, T. J. & Liu, F. C. Density functional theory study on the geometric, electronic and vibrational structures of alkaline metal porphyrin complexes. Chem.Phys.Lett. 101, 316–321 (2004).

41. Roy, D. R., Shah, E. V. & Roy, S. M. Optical activity of Co-porphyrin in the light of IR and Raman spectroscopy: a critical DFT investigation. Spectrochim. Acta. A. Mol. Biomol. Spectros. 190, 121–128 (2018).

Acknowledgements
This project was funded by the Deanship of Scientific Research (DSR), King Abdulaziz University, Jeddah, under grant No. (D-271 -662 -1441). The authors, therefore, gratefully acknowledge DSR technical and financial support.

Author contributions
Dr. Mohamed Ben Yahia is responsible for the achievement of the experimental part and the materials used (Quartz Crystal Microbalance apparatus) for the measurement of experimental adsorption isotherms of silver ions onto Helicenes molecules. 2- Pr. Manel Ben Yahia is responsible for the theoretical characterization of the experimental isotherms. She developed the advanced statistical physics models in order to interpret the complexation phenomenon at the microscopic scale.

Competing interests
The authors declare no competing interests.
Additional information

Correspondence and requests for materials should be addressed to M.B. or M.B.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2020