Research article

DFT and TD-DFT investigation of calix[4]arene interactions with TFSI⁻ ion

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A R T I C L E    I N F O

Keywords:
Materials chemistry
TFSI⁻ ion
Electronic proprieties
Gap energy
TD-DFT
Non covalent interactions

A B S T R A C T

Understanding the interactions of the calix[n]arene molecules with a variety of invited chemicals entities is getting very important. In this context, we have studied a new host-guest such as the interaction of the calix[4]arenes with the bis (trifluoromethylsulfonyl) imide TFSI⁻ ion. The energy gap has decreased from 3.53 eV to 2.11 eV indicating the reliability of the electrochemical evaluation of HOMO and LUMO energy levels. In a predominant number of cases, we obtain the spatial accumulation of HOMO and LUMO at the interface of phenol groups. Then, according to the QMBO charge distribution of these host-guests interactions, we have demonstrated the direction of charge transfer between the CX[4] molecule and the TFSI⁻ ion. More importantly, the non covalent interactions (NCI) have been investigated that the endo-cavity position of the TFSI⁻ is the most stable position between all these host-guests. By using DFT quantum methods, we have identified as a suitable host for TFSI⁻ which can be used in the electronic technology.

1. Introductions

The calix[n]arenes (abbreviated as CX[n]) are macro-molecules based on the phenol groups with a special cavity to encapsulate chemicals entities. The selective admission of the invited molecules in the macro or microscopic systems facilitates the recognition of the electrostatic and magnetic properties of the several cationic or anionic guests [1, 2, 3, 4, 5]. In this context, the CX [4] have been chosen in our study because of their own chemical composition and the hydrophobic cavity form [6]. This chemical material synthesized by a specific diameter and height, which facilitates the interactions with the diversity of the chemical species (cationic, anionic or neutral guests) and small molecules [7, 8, 9]. Recently, these molecules and especially their own cavity entice the researchers of those adaptations with most of the materials to be functional in the medical [10, 11, 12] or microbiological field [13, 14]. Also, these molecules are used in the activation of the solubility of monomers in the specific media and in the pharmaceutical drug delivery. Surrounding this strategy, we have explored the photo-physical properties of the complexes CX[4]-TFSI⁻ (endo-cavity and exo-cavity position of the TFSI⁻ ion). By using the DFT calculations, we have described the dynamic stabilities of the CX[4]-TFSI⁻ (CX[4]-T) complexes. The electronic properties have explained the phenomenal rigidity and conductivity of each host-guest. The recognition of the weak or the strong and the nature of the interactions of such guest with the CX [4] molecule have been analyzed by the non-covalent RDG function.

2. Computational detail

The stable structures of the CX[4]-TFSI⁻ complexes have been optimized with the DFT method by using the global hybrid Generalized Gradient Approximation (GGA) functional B3LYP [15, 16, 17]. This method together with the empirical Becke and Johnson damping dispersion corrections D3BJ in combination with the 6-31-G(d) basis set [18, 19] is implemented in Gaussian 09 program [20] and Gauss View [21] as a visual program. The binding energies of CX [4]-TFSI⁻ complexes are calculated including the BSSE (Basis Set Superposition Error) counterpoise (CP) corrected energy of Boys and Bernardi [22], which considers the fragments relaxation terms and the changes of the conformations in such geometric form. The binding energies are given by the following formula:

\[ \Delta E_{\text{CX}[4] \cdot \text{TFSI}^-} = E_{\text{CX}[4] \cdot \text{TFSI}^-} - E_{\text{CX}[4]} - E_{\text{TFSI}^-} + \text{BSSE} \]

where \( E_{\text{CX}[4]} \), \( E_{\text{TFSI}^-} \), and \( E_{\text{CX}[4] \cdot \text{TFSI}^-} \) are the total energies of the host and the guest or the host-guest molecules.

The absorption spectra of the CX[4]-TFSI⁻ complexes have been calculated using TD-DFT method at CAM-B3LYP-D3/6-31+G(d) level. In
this part, there are explanations in depth for the variation of the wavelengths of an endo-complex towards another exo-complex. The electronic study of the calixarene-TFSI complexes are very effective to show the specificity of each invitation to these materials. The stabilization energies and intermolecular interactions of CX4-TFSI complexes have been stimulated at the same theoretical framework. Molecular orbital analysis has been studied to explain the variation of the gap energy and the electron delocalization passing from TFSI 1 to TFSI 4. The study of the type of the interaction between the calixarene molecule and TFSI ion is better explained using the non-covalent-interaction-reduced density gradient (NCI-RDG). The NCI-RDG [23] analysis presents the graphical visualization of the non-covalent interaction regions in real-space, distinguishing hydrogen bonds and determining of the Van der Waals or repulsive interactions using simple color codes. The charge transfer has been also studied theoretically between the CX(4) molecule and the TFSI ion by the NBO analysis.

3. Calix[n] arene–TFSI complexes investigate by DFT method

3.1. Stable geometries and adsorption energy analysis

The development of encapsulation for the cyclic macromolecules such as β-Cyclodextrine, Cucurbit[n]urils and calixarenes is very effective to find the best host-guest suitable for the applications in the very intersect fields (technological [24] or vital [25, 26]). In this category, we have encapsulated the calix[4]arene molecules by the TFSI ion. We have optimized all the structures at the B3LYP-D3/6-31+G(d) level of theory. Moreover, we discussed our results for the CX4-TFSI complexes based on the position of TFSI ion. The computational study indicates that the CX4-TFSI possesses 4 host-guests (See Fig. 1), their binding energies are illustrated in Table 1. We have calculated the binding energy of the supra-molecular complexes such as the bis(trifluoromethanesulfonyl) imide (TFSI) molecules located at the endo-cavity position inner of the phenol ring, the ionic molecule located in the exo-cavity position such as the fluorine branch directed to the center of the cavity, the TFSI ion located between the two phenolic outside the cavity. In this position, these molecules are well balanced by the hydrogen bonds. This molecule has been located outside the cavity such as the oxygen atom directed to the lower edge level. The optimization of the CX4-TFSI 1 gives a binding energy value equal to 52.23 kcal/mol. The no-destruction of the OH network ensures the conservation of the C4 symmetry group in this material. Moreover, the energy of the CX4-TFSI 2 (endo-cavity) is higher than the first supra-molecular complex, this value is equal to 52.090 kcal/mol. The stability of this complex is due to the interaction of the fluorine atoms with the cyclic network of the hydrogen bonds. Moreover, the interaction of this fluorine extremity with the hydrogen atoms connected to the phenol groups promotes the stability of this material.

The weak electrostatic interaction of the TFSI ion with the macrocycle molecule leaves this complex placed third considering the stabilization order in comparison with others complexes. The complex CX4-TFSI 3 has a lower value of the binding energy in comparison with CX4-TFSI 2 and CX4-TFSI 1. The value of adsorption energy is equal to 36.09 kcal/mol. The interaction of the hydrogen bonds of the two oxygen atoms connected to the sulfur atoms ensures the stability of this chemical compound. We note that, the interaction of the H-phenyl rings and the oxygen-fluorine groups was forming the weak CH–π hydrogen bonding. The complex CX4-TFSI 4 is the most stable system among all the other host-guests. The binding energy in this chemical material is around to 53.28 kcal/mol. The stability of this structure is due to the interaction of oxygen atom connected to the sulfur atom with the hydrogen OH bond located at the lower edge levels. The larger stability of this complex is due to the dipole-dipole interaction between the TFSI 4 ion and the calixarene molecule. This interaction promotes the reduction of the symmetry groups from C4 to the C1.

3.2. Electronic absorption spectra

The UV-visible spectrum of the four supra-molecules complexes will be studied to understand the electronic behavior of the different host-guests. Fig. 2 shows the superposition of the absorption spectra in the order of stability for these chemical compounds. From these spectra, we have found that the CX(4)–T complexes have an effective wavelength λ in the vicinity of 246 nm for the complexes CX(4)–T4, T2 and 244 nm, 243 nm for the CX(4)–T1 and CX(4)–T3 molecules successively. In this region we can deduce that the complex CX(4)–T2 has a more intense peak in comparison with the other supra-molecular complexes. The red-shifted of the wavelengths λ in the spectrum CX(4)–TFSI 2 is explained as, this molecule has a high sensibility to the calixarene molecules and a lower excited state than other host-guests. The TD-DFT computation for the interest complexes predicts a strong electronic transitions located at 256 nm corresponding to the CX(4)–T4. Fig. 2 shows the S0 to S1 transition of the four endo-VS-exo cavity complexes located near the 220, 221, 219 and 218 successively. The different wavelengths λ of the three most stable complexes is almost 1 or 2 nm. This region is characterized by the population of electronic charges surrounding the phenol groups located in the front of the invited ion. In addition, we notice, the appearance of the three corrective peaks having an important intensity in the neighborhood of 202 nm for the CX(4)–T1/T2 complexes and 198 nm for the CX(4)–T3 complex. In this zone there is a regrouping of electron charges in the region of the interaction between the invited and the calix[4]arene molecule.

The π to π* transitions were expected to co-exist at the low wavelength λ levels near to 189 nm for CX(4)–T1/T3/T4 and 187 nm for CX(4)–T2. These transitions may be due to the interaction of the aromaticity of the phenyl group with the organic compound. Finally, the supra-molecular complexes CX(4)–T2/T3 have two low red-shifted peaks located in the vicinity of 182 nm. Concerning the CX(4)–T2 complex, this peak may be due to the presence of the electronic charges at the center of the cavity linked to the existence of the H-bonding between the Host and the guest. In the same framework, the CX(4)–T3 complex is specified by the maximum electronic charges in the region of interactions between the fluorine group and the OH group of the CX(4) molecule.

3.3. FMO study

The ability to donate an electron; HOMO and the ability to accept an electron; LUMO is a very interesting study to take into account the chemical stability part of these macro-cycle complexes. The energy gap (Eg) is an important parameter which explains the chemical stability of these organic compounds very well. In Fig. 3, we have depicted the Frontier Molecular Occupied HOMO and the Frontier Molecular Unoccupied LUMO+1, LUMO+2, LUMO+3 and LUMO+5 of CX4–TFSI T1/T2/T3/T4. We have calculated all these FMO at the TD-CAM-B3LYP-D3/6-31+G(d) level of theory. From Fig. 3, we can see that, the occupied frontier specified by the red color, or the unoccupied frontier specified by the blue color. The HOMO-LUMO energy gaps of the four stable complexes are equal to 3.53eV (f = 0.4747), 2.95eV (f = 0.2376), 2.83eV (f = 0.2968) and 2.11eV (f = 0.2986) successively. According to Fig. 3, we find that, it’s very clear that the HOMO orbital of the CX(4)–T2/T3 is well delocalized and present at the levels of the phenol group. The majority of the charge population in the CX(4)–T3 complex co-exists in the phenol groups located in the front of TFSI ion. Concerning the second host-guest, we show that, the maximum of the charge distribution is located around the TFSI ion. The region in the front of the host has a low delocalization. On the contrary, the organic compound CX(4)–T4 is characterized by a delocalized frontier orbital located on the levels of the phenol group aromaticity near the TFSI ion. The FMO L1–1/2/3/5 of these chemical compounds are characterized by unequal charge distributions at the phenol group levels in the region of CX(4)–TFSI interaction. The complex CX4–TFSI 1 is characterized by an electronic delocalization situated in the symmetrical phenol groups. The π–binding
Fig. 1. Optimized geometries of and CX[4]-TFSI’ (CX[4]-TFSI 1 (a), CX[4]-TFSI 2 (b), CX[4]-TFSI 3 (c), CX[4]-TFSI 4 (d)) (Top views and Bottom views) structures using B3LYP-D3/6-31+G(d) method.
orbital LUMO+3 of the CX[4]-TFSI-2 is characterized by an important population of the charge distributions. The population of the charge delocalized of the CX[4]-TFSI-2 complex is located in all symmetrical phenol groups. The complex CX[4]-TFSI-3 has a π-electronic charge located at the phenol group in front of the invited molecule. The unoccupied molecular orbital LUMO+1 is specified by the existence of a maximum electronic population located around the two phenol groups in the face of the invited molecule. Moreover, from Table 2 we can deduce

| Complexes | \( E_b \) | BSSE | \( E_b \) (with BSSE) |
|-----------|----------|------|---------------------|
| CX4-TFSI 1 | 54.23    | 0.0028 | 52.23              |
| CX4-TFSI 2 | 52.09    | 0.0020 | 52.09              |
| CX4-TFSI 3 | 36.09    | 0.0046 | 36.09              |
| CX4-TFSI 4 | 53.28    | 0.0022 | 53.28              |

Fig. 2. Absorption spectrum of caption of TFSI\(^-\) ion by calix[4]arene molecule (Blue: CX[4]-TFSI-4, Green: CX[4]-TFSI-2, Red: CX[4]-TFSI-1, Magenta: CX[4]-TFSI-3) (according to the order of stability).

Fig. 3. Frontier Molecular Orbital analysis of CX[4]-TFSI(CX[4]-T) complexes.
that, the CX-[4]-TFSI-4 is very conductive than other host-guests and CX
[4]-TFSI-3 is less conductive than others. This information explains why
the extremity of the TFSI-4 (endo-cavity) forms a hydrogen bond with the
cyclic group of the calixarene molecule. This system is very interesting in
the optical and electrical
fields. The energy gaps of these host-guests
passing from CX-[4]-T1 to CX-[4]-T4 appear as a support to understand
the active property of these molecules. It is very important that, the en-
ergy gaps are very high in the system or co-exists the H-binding between
the cyclic net work of calixarene and the TFSI-ion. Hence it is thermo-
dynamically effective for the transfer of an electron to occur. The lowest
value of the HOMO-LUMO+1 gap presented the transition from the
ground to the first excited state of the CX[4]-TFSI-3 complex. This trans-
ition is due to the weak transfer of conjugated electrons between these
two molecules. We conclude that, this material has a low kinetic stability
and chemical reactivity in comparison with other complexes. From to
Fig. 3, we deduce that, it exists an anti-bonding $\pi$ character that has a
maximum of electronic charges between the CX[4]-T4 and the calixarene
molecule. The diagram of energy gap shows that these materials solve
many problems in the optoelectronic field.

### 3.4. Charge NBO analysis

The charge NBO has been calculated to identify the characteristic of
the charge transfer [34, 35] between the ionic molecule and the calix[4]
arene material. The NBO charge calculation of the CX[4]-TFSI 1/3
complex is equal to -0.978e. The $Q_{NBO}$ of the CX[4]-TFSI 2/4 chemical
compounds is equal to -0.956e. In addition, the $Q_{NBO}$ value of CX[4]-T1
and CX[4]-T3 inclusions explains the weak interaction between the guest
and the host in this case. On the contrary, from Table 3, we find that the
NBO loads of the supra-molecules complex CX[4]-T2/T4 is very impor-
tant, which explains the strong interaction between the TFSI$^-$ and the
calix[4]arene molecule. From $Q_{NBO}$ calculation, we can clearly explain
that, the TFSI$^-$ ion has a good stability in endo-cavity position of calix-
arene in comparison with exo-cavity position. We notice that always
$Q_{NBO}<0$. This result shows that the transfer of charge always takes place

**Table 2**

Main transition states, the corresponding assignments, $E_g$ and oscillator strength,
for all compounds, recorded with TD-DFT/CAM-B3LYP/6-31+G(d) method.

| Compounds | Electronic Transitions | Energy gap (eV) | Oscillator strength(f) | Molecular orbital |
|-----------|------------------------|-----------------|-----------------------|------------------|
| TFSI1     | S0→S1                  | 2.83            | 0.2968                | H→L$^+$2        |
| TFSI2     | S0→S1                  | 2.95            | 0.2376                | H→L$^+$3        |
| TFSI3     | S0→S1                  | 3.53            | 0.4747                | H→L$^+$1        |
| TFSI4     | S0→S1                  | 2.11            | 0.2986                | H→L$^+$5        |

**Table 3**

NBO charge analysis of CX[4]-TFSI complexes.

| CX4-TFSI | $Q_{NBO}$ (e$^-$) |
|----------|-----------------|
| CX4-TFSI 1 | -0.978          |
| CX4-TFSI 2 | -0.956          |
| CX4-TFSI 3 | -0.978          |
| CX4-TFSI 4 | -0.956          |

![Fig. 4](image-url) NCI isosurfaces for the inclusion complexes CX[4]-TFSI 1 (a), CX[4]-TFSI 2 (b), CX[4]-TFSI 3 (c) and CX[4]-TFSI 4 (d). The iso-surfaces were constructed with RGD = 0.5 au and the blue-red colors scaling from -0.01 to -0.01 au.
from CX[4] to TFSI− ion.

3.5. NCI-RDG theory

The non covalent interaction (NCI) via reduced-density-gradient (RDG) facilitates the understanding of the nature of interactions in different types of encapsulated complexes. In this context, we have discovered the type of the interactions between the CX[4] and the TFSI− ion. The NCI isosurface for CX[4] and the specific encapsulated system (TFSI−1, 2, 3 and 4) are shown in Fig. 4. Moreover, the van der Waals interactions are shown with a green spot, the strong hydrogen bonding interactions are shown with a blue spot and the repulsive steric forces are visualized by the red spot. From Fig. 4, we note that the CX[4]:TFSI−1 complex is characterized by the existence of the green spots located between the -S-H groups of the guest and the O–H groups of the host. The strong repulsive steric interactions are observed at the phenol five membered rings which are represented by the red spots. The strong H-bonding interactions have been conserved at the lower edge level. This existence of hydrogen bonds appears as blue spots (See Fig. 4-a). In the case of the inclusion complex CX[4]:TFSI−2, Fig. 4-b explains that, the weak van der Waals type of attractive interactions co-exist in the area of the –C-F3 group interaction and the four symmetric phenol groups. There is also the appearance of the weak van der Waals bonds surrounding the CH2 group of the calix[4]arene molecule. This type of attractive interactions is indicated by the green disk. In addition, it shows the appearance of the red spots in each phenol group center which explains the existence of the repulsion steric force in such zone. The existence of the weak intra-molecular hydrogen bonding interactions at the cyclic O–H net work are specified by the red spot. Concerning the CX[4]:TFSI−3 complex, we found that the interaction between the TFSI−3 ion and the calix[4]arene molecule is very weak, of van der Waals nature. The interrelation of this ion with the endo space of the cavity is explained by the red bond –S-O–H (CX[4]-TFSI−4) and –CF–H (CX[4]-TFSI−2).

4. Conclusion

A DFT calculation of the binding energy of the CX4-TFSI− complexes show that the CX[4]:TFSI− (endo-cavity) interaction is the most stable system. The absorption spectra explain that the ionic material CX4-TFSI−4 is a good candidate for the optoelectronic application. The NCI theory helps to discover the great, typical and the nature of the CX[4]:TFSI− interactions. This theory has demonstrated the great stability of the TFSI− (endo-cavity position) in comparison with TFSI− (exo-cavity position). The interrelation of this ion with the endo space of the cavity is explained by the strong bonds –SO–H (CX[4]-TFSI−4) and –CF–H (CX[4]-TFSI−2).

Declarations

Author contribution statement

B. Gassoumi, G. Ghalla and R. Ben. Chaabane: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Funding statement

This work was supported by the Tunisian's Ministry of high education and scientific research.

Competing interest statement

The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.

Acknowledgements

In this work, we were granted access to the HPC resources of the FLMSN, ‘Fédération Lyonnaise de Modélisation et Sciences Numériques’, partner of EQUIPEX EQUIP@Meso and to the ‘Centre de calcul CC-IN2P3’ at Villeurbanne, France.
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