Impact of Doxorubicin on Self-Organization of Congo Red: Quantum Chemical Calculations and Molecular Dynamics Simulations

Klaudia Kwiecińska, Anna Stachowicz-Kuśnierz, Anna Jagusiak, Irena Roterman, and Jacek Korchowiec*

ABSTRACT: Quantum-chemical calculations and molecular dynamics simulation were applied to a model self-organization process of Congo red (CR) molecules in aqueous solution and the impact of doxorubicin (DOX) molecules on such a process. It was demonstrated that both pure CR/CR and mixed CR/DOX dimers were stable. Van der Waals interactions between aromatic units were responsible for a stacked dimer formation. An important source of stabilization in the CR/CR dimer was the polarization energy. In the CR/DOX mixed dimer long range, electrostatic interactions were the main driving force leading to complexation. An implicit solvent model showed that the formation of the CR/CR dimer was favored over the CR/DOX one. Molecular dynamics simulations demonstrated rapid complexation. In the pure CR system, short sequences of ribbon-like structures were formed. Such structures might be glued by hydrogen bonds to form bigger complexes. It was shown that the aromatic part of the DOX molecule enters CR ribbons with the sugar part covering the CR ribbons. These findings demonstrated that CR may find applications as a carrier in delivering DOX molecules; however, further more extensive investigations are required.

INTRODUCTION

Development of supramolecular chemistry opens up new horizons for many fields of science. The chemistry of such molecular systems is ruled by weak intermolecular interactions, e.g., van der Waals forces, hydrogen bonds, and π−π and electrostatic interactions. The presence of noncovalent interactions between organic molecules allows the formation of large self-assembled supramolecular structures.1−3 This kind of multimolecular structure is characterized by high elasticity and a large surface area. These properties enabled some supramolecular associates to form complexes with proteins, which in general are compounds that are unable to interact at sites other than the active site.4 Congo red (CR) is an example of a self-assembling supramolecular compound creating ribbon-like structures related to the association of flat, polyaromatic parts of these molecules. It belongs to the azo dye group, and it is used especially as an indicator and biological stain. CR has been used for over a century to demonstrate the presence of amyloidal deposits in tissue, e.g., in brain tissue, or in the heart5 or kidneys6 in the diagnosis of Alzheimer’s disease.

Amyloid fibers bind to Congo red, giving the effect of apple-green birefringence under a polarized light microscope, which remains the gold standard in recognizing amyloid deposits.7,8 Congo red is also often used as a model dye in evaluating the effectiveness of various materials that catalyze the degradation of organic pollutants in water.9,10 The structure of Congo red is symmetrical. Its central part consists of a biphenyl ring linked through azo bonds with naphthalene rings substituted with sulfone and amino groups. Individual molecules interact with each other to form larger supramolecular, ribbon-like structures. The polyaromatic structure of Congo red enables intercalation of various compounds with planar groups, e.g., the doxorubicin (DOX) drug.

Received: March 11, 2020
Accepted: July 14, 2020
Published: July 27, 2020
Doxorubicin is an anthracycline antibiotic commonly used as an anticancer drug. The mechanism of doxorubicin action is associated with the formation of a stable complex with DNA, thus inhibiting further cell division ultimately causing its death. Unfortunately, chemotherapy is not an ideal treatment method. Cytostatics used in this treatment destroy all rapidly dividing cells, including normal cells, which causes numerous side effects for the human body. Currently, new solutions are sought in order to reduce the therapeutic dose of the drug while maintaining its effectiveness and reducing its toxicity to normal cells. One of these possible solutions is the use of drug carriers.

The self-assembled ribbon-like structures that form elongated Congo red systems owing to their unique properties are used in targeted drug delivery to cancer tissue. Such systems are an example of a new type of ligand for proteins to which they bind using a nonclassical mode of interaction. An example is binding to antigen–antibody complexes while not binding to free antibodies. Research to date has demonstrated the possibility of using Congo red systems in vivo as potential drug carriers. At the same time, CR systems effectively bind to various molecules, including drugs, to form co-micelles with them. Research has shown that Congo red supramolecular assemblies are investigated and its possible use as a carrier system for doxorubicin, a model anticancer drug, is considered. We focus on the microscopic description of CR and CR/DOX clusters, including interactions responsible for dimer stabilization, the influence of solvent (water), and the mechanism of the clustering process in aqueous solution.

**COMPUTATIONAL DETAILS**

All quantum-chemical calculations were carried out using the Gaussian 09 suite of programs. The structures of DOX and CR (see Figure 1) were located at the density functional level of theory, i.e., the hybrid B3LYP functional with a 6-311G(d,p) basis set. The Grimme dispersion correction was taken into account. The located optimum structures of the monomers were used to prepare the initial structures of CR/CR and CR/DOX dimers. These structures were further optimized at the B3LYP/6-311G(d,p) level of theory. All located minima on the energy surface were verified by vibration frequency analysis. The nature of interactions was characterized by a self-consistent charge and configuration method for subsystems (SCCCMS). The SCCCMS scheme has been described elsewhere. The scheme is based on a simple model of mutual interactions among reactants. The molecular system M = AB is partitioned into reactants A and B. The calculations for reactant A (B) are performed in the presence of the reaction partner B (A) emulated by its charge distribution. The charges fitted to the molecular electrostatic potential were employed (CHelp scheme). Calculations are terminated after the convergence in electronic structures and charge distributions is achieved. The SCCCMS calculations are performed for different oxidation states of A and B. In such a way, the energy surface spanning the population space (N_A, N_B) is constructed. The idea of SCCCMS is shown in Figure 2 that illustrates the cut of the energy surface along the isoelectronic line, E_M = E_M(N_{CT}), together with hypothetical stages of charge reorganization within the SCCCMS scheme, marked by closed circles. ES, P, and CT denote electrostatic, polarization, and charge transfer energies, respectively.

**Figure 1.** Structure of (a) Congo red and (b) doxorubicin.

**Figure 2.** Cut of the energy surface along the isoelectronic line, E_M = E_M(N_{CT}), together with hypothetical stages of charge reorganization within the SCCCMS scheme, marked by closed circles. ES, P, and CT denote electrostatic, polarization, and charge transfer energies, respectively.
\( \eta_{AB} - 2\eta_{AA} > 0 \). The reactant chemical potentials \((\mu_A \text{ and } \mu_B)\) and hardness values \((\eta_{AA}, \eta_{BB} \text{ and } \eta_{AB})\) are given by the following partial derivatives: 
\[ \mu_A = (\partial E_M/\partial N_A)_N, \quad \mu_B = (\partial E_M/\partial N_B)_N, \quad \eta_{AA} = (\partial^2 E_M/\partial N_A^2)_N, \quad \eta_{BB} = (\partial^2 E_M/\partial N_B^2)_N, \quad \eta_{AB} = (\partial^2 E_M/\partial N_A \partial N_B)_N \] 
and 
\[ \eta_{AB} = (\partial^2 E_M/\partial N_B \partial N_A)_N \] respectively. The additional constraint imposed on derivatives, \( v = v(\vec{\eta}) \), is the external potential due to nuclei. Here, we do not consider derivatives with respect to \( v(\vec{\eta}) \). In other words, the positions of nuclei were fixed. Notice that the first (electron density) and second-order energy derivatives (linear response function) with respect to the external potential as well as the mixed second-order derivatives (Fukui function) are of special importance in the theory of chemical reactivity. For details, see the excellent works in the field of conceptual density functional theory.\(^{18-20}\) The SCCCMC scheme utilizes the polarization approximation. The energy difference among hypothetical stages defines electrostatic (ES), polarization (P), and charge-transfer (CT) energies. Due to polarization approximation, the minimum of the parabola does not correspond to energy of the supermolecule. The difference, depending on the level of theory, includes exchange (Hartree–Fock method) or exchange and correlation (Kohn–Sham and post-Hartree–Fock methods) contributions. In addition, it contains Grimme dispersion corrections. For this reason, it will be called van der Waals (VdW) energy. Notice that only two points shown in Figure 2 have a clear physical meaning. The first point corresponds to the intra-reactant equilibrium \([\text{point of the parabola at } E_M(N_{CT} = 0) = E_M^0\text{ (polarized reactants before CT, } M = (A \B)), \mu_A = \mu_A(\vec{r}) \neq \mu_B = \mu_B(\vec{r})\])\), and the second point corresponds to the global equilibrium \([\text{the minimum of the parabola, open reactants in supermolecule } M = (A; B)), \mu_A = \mu_A(\vec{r}) = \mu_B = \mu_B(\vec{r}) = \mu_M\text{. The information from the energy surface can be utilized to compute for the charge reorganization accompanying P and CT processes as well as the CT Fukui function and its components.}\(^{21}\)

Additional information regarding affinity of CR and DOX toward self-organization was derived from molecular dynamics (MD) simulations. The CR/DOX systems were previously discussed by Jagusiak and Panczyk;\(^{22}\) however, the AMBER force-field parameterization of CR did not support self-organization of these molecules, which was observed experimentally.\(^{23,24}\) This divergence can be attributed to the charge distribution that in the AMBER force field is derived by fitting to the molecular electrostatic potential of the isolated molecule. For this reason, CR and DOX molecules were parameterized in the CHARMM force field using a force field toolkit of VMD graphical software.\(^{24}\) In contrast to the AMBER charges, the CHARMM charges are derived by reproducing the interaction energies of hydrogen donor and hydrogen acceptor atoms of the CR or DOX molecule with water molecules. It can be expected that this will improve the description of CR and CR/DOX aggregates.

The MD calculations were carried out using the NAMD package with periodic boundary conditions imposed on the system. An all-atom CHARMM force field\(^{25,26}\) was used. The TIP3P model\(^{28}\) was adopted for water. The VMD software was applied to visualize the trajectories and compute selected quantities. Density-based cluster analysis of the trajectories was performed with the dbscan function of the R package.\(^{29}\)

In all calculations, the number of atoms \((N)\) in the simulation box was greater than 200,000. The initial configuration was obtained by placing the target molecules on a cube grid. A random orientation of each molecule was used. The CR anions were neutralized by sodium cations. Simulations at constant temperature \((T = 293 K)\) and pressure \((p = 1 \text{ bar})\) were employed. These simulations were preceded by energy minimization and 10 ns canonical \((NVT, \text{ number of particles, volume, and temperature})\) ensemble simulations. The \((NpT)\) ensemble was applied for 120 ns. The ratio of the \(x\) and \(y\) box sizes was constant. Five independent trajectories were computed for each molecular system, and their average is reported as the final result.

### RESULTS

In Table 1, we have listed the interaction \((\text{INT})\) energies and their components for dimer systems. To the best of our knowledge, CR/CR and CR/DOX dimers were not described previously in the literature. The interaction energy was computed using a supermolecular approach: \( E_{\text{INT}} = E_{AB} - E_A - E_B \) where the energies \( E_A \) and \( E_B \) are calculated for monomers having the same geometry as in the dimer. The basis set superposition error was eliminated using the counterpoise method of Boys and Bernardi.\(^{30}\) The total binding \((b)\) energy should be supplemented by the deformation energy (shown in the last row of Table 1), \( E_{\text{DEF}} = E_A + E_B - E_A^0 - E_B^0 \geq 0 \). This term is positive, and the energies of the reactants in the minimum structures \( E_A^0 \) and \( E_B^0 \) are more negative than in dimer geometries. As one would expect, P and CT energies are stabilizing (negative) while DEF is destabilizing (positive). The highest destabilization contribution to the INT energy in the CR/CR system is the ES term. This behavior was not surprising because the CR is negatively charged. This energy term in a mixed CR/DOX system is negative; therefore, the ES term should enhance the self-organization of molecules or at least promote the incorporation of DOX molecules into ribbon-like supramolecular structures of CR. The polarization energy is the most stabilizing effect in the CR/CR dimer. The polarization contribution in CR/CR is almost three times higher than in the CR/DOX dimer. The second highest is VdW energy. Except DEF energy, the remaining contribution to CR/DOX stabilizes the dimer formation.

Dimerization enthalpies and Gibbs free energies are summarized in Table 2. Data corresponding to vacuum and the implicit solvent model, namely, the polarizable continuum model (PCM),\(^{31}\) are reported. The temperature and entropy effects did not change the trends observed in INT (b) energies. Binding enthalpy and Gibbs free energies collected in Table 2 favor the formation of a CR/DOX dimer. This picture changes for the PCM water model. The formation of a CR/CR dimer is

### Table 1. Interaction Energies and Their Components Computed at the B3LYP/6-311G(d,p) Level of Theory for CR/CR (Second Column) and CR/DOX (Third Column) Dimers; All Values Are in kcal/mol

| Energy   | CR/CR     | CR/DOX   |
|----------|-----------|----------|
| INT      | 19.2      | −60.5    |
| ES       | 73.9      | −25.0    |
| P        | −31.5     | −11.8    |
| CT       | 0.0       | −0.4     |
| VdW      | −23.2     | −23.3    |
| DEF      | 16.8      | 11.4     |
| b        | 36.0      | −49.1    |

\[ \text{https://dx.doi.org/10.1021/acsomega.0c01095} \]
Table 2. Binding Enthalpy and Gibbs Free Energy in Vacuum and the Implicit Solvent Model (PCM) Computed at the B3LYP/6-311G(d,p) Level of Theory for T = 293 K and p = 1 bar; All Energies Are in kcal/mol

| molecular system | vacuum |PCM model |
|------------------|--------|----------|
| ΔH  | ΔG    | ΔH     | ΔG     |
| CR/CR | 19.3   | 44.8    | -56.1  | -30.6  |
| CR/DOX| -61.0  | -40.2   | 38.6   | -17.3  |

now preferred. This observation supports the self-organizing ability of CR in water. To check whether these conclusions do not depend on the functional type, the binding enthalpy and binding free enthalpy were calculated at the M062X/6-311(d,p) level of theory. The calculated data are collected in Table S1. The same conclusions follow from reported data, namely, the reverse of stability of CR/DOX and CR/CR dimers when going from vacuum to the implicit water model. Additional support for the observed reverse in stability of CR/DOX and CR/CR dimers when going from vacuum to the implicit water model results from polarization (Table 1). The stabilization due to polarization energy in CR/CR dimers is three times greater than in the CR/DOX dimer. The self-consistency in PCM calculations, namely, the polarization of solute and backpolarization of dielectric medium, will depend on this particular feature of charge distribution.

To understand the nature of stabilization in dimers, quantum theory of atoms in molecules (QTAIM) was applied. Topological properties of electron density have been shown to correlate with the strength of intermolecular interactions. Here, bond critical points (BCPs) were used to detect the bonding pattern. The existence of bond paths between the atoms of different monomers is a clear indicator of bonding contacts. Further classification can be done by the value of the electron density at the BCP, \( \rho_{BCP} \). In the CR/CR system, there are 28 BCPs between CR monomers. The value of the electron density at BCPs ranges from 0.004 to 0.031 a.u. In this variety are 28 BCPs between CR monomers. The value of the electron density at the remaining contacts is incorporated into the ribbon while the sugar part protrudes from the clustering process; the aromatic part of the DOX molecule protruding to the surface and the aromatic moiety built into the ribbon. Sample clusters, corresponding to the mixed CR/DOX systems are shown in Figures S2 and S3, observation also supports the reverse of stability of CR/DOX dimers when going from vacuum to the implicit water model.

Molecular Dynamics Simulations. Vacuum and PCM calculations performed for both dimers indicate the increasing ability of CR to form organized structures in water. Such findings are qualitative since no explicit water molecules were taken into account. This was the main reason for using molecular dynamics simulations. For both systems (CR/CR and CR/DOX), the initial configuration was random without any self-organizing pattern. Practically from the beginning of simulations, the formation of dimers could be observed, which subsequently aggregated into larger clusters. Clustering patterns for the last frames of selected CR/CR and CR/DOX trajectories are shown in Figure 4. Panel a corresponds to the pure CR system, while panel b corresponds to the mixed equimolar CR/DOX molecular system. Clusters of different sizes can be observed. Sample clusters, corresponding to the sizes of 8 and 13 molecules, are highlighted with red surfaces and drawn in detail in panels c and d. In the case of the CR system, the molecules form two ribbon-like substructures that are glued by hydrogen bonds. A qualitatively similar picture is observed in a mixed CR/DOX system. Ribbon-like structures are also present with the sugar moiety of DOX molecules protruding to the surface and the aromatic moiety built into CR ribbons. The clustering patterns for the remaining CR and CR/DOX systems are shown in Figures S2 and S3, respectively.

Changes in the average number of monomers forming a cluster during the courses of both types of simulations are presented in Figure 5. The resultant curves were averaged over five trajectories. Deviations from the average values are shown in the Supporting Information (Figures S4 and S5). Two approaches were employed to define the center of mass (CM) of DOX molecules in the CR/DOX system. In the first one (red line in Figure 5), the CM was computed as the center of mass of the whole DOX molecule. In the second case (green line), only the aromatic part of a DOX molecule was taken into account. The differences are associated with the mechanism of the clustering process; the aromatic part of the DOX molecule is incorporated into the ribbon while the sugar part protrudes.
from the ribbon surface. For CR molecules, both approaches give the same result. In both systems, a saturation effect can be observed. After saturation, in the CR system, there are on average five CR molecules in the ribbon. For a mixed CR/DOX system, the average number of molecules is higher and amounts to 7 or 8 molecules depending on the approach employed to calculate the center of mass. For this system, larger fluctuations in the average cluster size can be observed. They are connected with greater mobility of DOX molecules associated with the surface of larger clusters whose aromatic parts are not incorporated into the ribbons. Fluctuations observed between the successive points indicate that the monomer can detach from the aggregate. However, eye examination of the trajectories reveals that this detachment is only temporary and the displaced molecules quickly return to the clusters. Such temporary dissociation is more frequent for the mixed CR/DOX system than for the CR one due to the presence of a hydrophilic sugar group in DOX and a more disordered structure of CR/DOX clusters.

The clustering process is summarized in Figure 6. The histogram illustrates the formation of clusters of various sizes.

Data is acquired from the terminal parts (after saturation, Figure 5 and Figures S4 and S5) of all five trajectories obtained for each system. In summary, ca. 1500 frames were gathered for both CR and CR/DOX systems. Histograms for individual runs are shown in Figures S6 and S7 of the Supporting Information. Again, two approaches were used to calculate CM of DOX molecules. In the case of the CR system, clusters composed of 2 to 17 molecules can be observed. The distribution of cluster sizes is relatively narrow and exhibits a clear maximum at the value of four molecules in a cluster. In this system, free monomers are practically not observed. In contrast, the cluster size distribution in the CR/DOX system is broader and extends toward larger clusters. The largest cluster observed in the simulations consisted of as many as 31 molecules. Surprisingly, free monomers also occur in this system. In the calculations employing CMs of whole DOX molecules, they are the most abundant entity in the system. However, eye examination of the trajectories reveals that these are not free monomers but molecules (mostly DOX) loosely associated with the surface of larger clusters. Among aggregates, the most populated number of molecules in a cluster is 4. However, the maximum is less pronounced than in the CR system and larger clusters are also present. Two approaches used in cluster analysis give qualitatively similar predictions. A higher green bar than a blue one indicates that whole DOX molecules cover the ribbon and the aromatic part

Figure 4. Visualization of the last MD simulation frames for (a) pure CR and (b) mixed CR/DOX molecular systems. Example clusters, marked in red color, are depicted in panels (c) and (d) together with all water molecules (cyan color) and Na⁺ cations (green balls) within a 10 Å distance from the cluster. In panels (a) and (b), CR molecules are drawn in gray and DOX molecules in black. In panels (c) and (d), CR molecules are colored according to atom names (C - black, O - red, N - blue, S - yellow, and H - gray). In DOX molecules, the aromatic part is drawn in orange and the sugar part in magenta.

Figure 5. Average number of monomers in clusters formed during MD simulations. The black line corresponds to the pure CR system, while green and red lines correspond to the mixed CR/DOX system with two different definitions of CM (see main text for details).

Figure 6. Histogram illustrating the cluster size distribution for pure CR (red), mixed CR/DOX based on the center of mass (CM) of whole DOX molecules (green), and mixed CR/DOX based on the CM of aromatic rings (blue).
is completely immersed in water. This can be considered as the first step in incorporating DOX molecules into CR ribbons.

In cluster analysis, information about the source of stabilization is lost. To get some information regarding the nature of stabilization (bonding contacts) in clusters, the radial distribution function (RDF) was plotted. The RDFs between the center of mass in pure and mixed systems are shown in Figure 7a. In this graph, the CM of the aromatic part of DOX molecules was taken into account as the plots based on the CM of the whole molecule do not provide an accurate picture of the interactions between the aromatic parts of the molecules. Panel b in Figure 7 presents the RDF describing the ability of the formation of hydrogen bonds between the oxygen atoms from the SO3 group and DOX oxygen and nitrogen atoms. In Figure 7a, an oscillatory pattern is observed. This indicates the formation of ribbon-like structures. Such stacked structures are stabilized by van der Waals interactions as demonstrated by the bond critical paths shown in Figure 3 for CR/CR and CR/DOX dimers. This type of interaction occurs in pure CR (dashed red line) as well as in a mixed CR/DOX system. In a mixed system, the RDF for CMs of CR/CR, CR/DOX, and DOX/DOX molecules is plotted. Hydrogen bonds are another important source of stability between CR and DOX molecules. Panel b in Figure 7 indicates the formation of hydrogen bonds between O(SO3)−H−O and O(SO3)−H−N. The former hydrogen bonds are particularly strong due to the flexibility of the DOX sugar unit. The BCP in the CR/DOX dimer has the highest value of electron density (0.04). The later hydrogen bonds are weaker with a ρ_{BCP} close to 0.01.

**CONCLUSIONS**

It was demonstrated that Congo red molecules self-assemble in water in accordance with experimental data. This indicates that proper force-field parameters for CR were obtained. Ribbon-like structures of varying lengths are formed. These flat stacked structures are stabilized by van der Waals contacts. Hydrogen bonds are responsible for gluing short ribbons of CR. In the mixed CR/DOX system, doxorubicin is easily incorporated into the CR ribbons. The aromatic parts of DOX penetrate the CR ribbon, while the sugar part covers the surface of the ribbon. Hydrogen bonds are responsible for covering the CR ribbon. In the mixed CR/DOX system, larger clusters are formed than in a pure CR system. Weak intermolecular interactions are responsible for complexation. Self-assembled CR structures can be used as carriers for DOX molecules.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.0c01095.

Table S1: binding enthalpies and Gibbs free energies computed at the M062X/6-311G(d,p) level of theory; Figure S1: isosurfaces illustrating the polarization process in CR/CR and CR/DOX dimers; Figure S2: clustering patterns for the remaining four CR trajectories; Figure S3: clustering patterns for the remaining four CR/DOX trajectories; Figure S4: average number of monomers in CR clusters; Figure S5: the average number of monomers in CR/DOX clusters; Figure S6: cluster size distribution in CR systems; and Figure S7: cluster size distribution in CR/DOX systems (PDF)

**AUTHOR INFORMATION**

Corresponding Author

Jacek Korchowiec — Faculty of Chemistry, K. Gumiński Department of Theoretical Chemistry, Jagiellonian University, 30-387 Kraków, Poland; orcid.org/0000-0002-9698-7369; Phone: +48 (12) 6862379; Email: korchow@chemia.uj.edu.pl

Authors

Klaudia Kwiecińska — Faculty of Chemistry, K. Gumiński Department of Theoretical Chemistry, Jagiellonian University, 30-387 Kraków, Poland
Anna Stachowicz-Kuśnierz — Faculty of Chemistry, K. Gumiński Department of Theoretical Chemistry, Jagiellonian University, 30-387 Kraków, Poland
Anna Jagusiak — Medical College, Chair of Medical Biochemistry, Jagiellonian University, 31-034 Kraków, Poland
Irena Roterman — Medical College, Department of Bioinformatics and Telemedicine, Jagiellonian University, 31-530 Kraków, Poland

Complete contact information is available at: https://pubs.acs.org/10.1021/acsomega.0c01095

Author Contributions
A.J. and I.R. conceived the work. J.K. planned all the calculations. K.K. performed quantum-chemical calculations and part of molecular dynamics simulations. A.S.-K. performed part of molecular dynamics calculations and conducted all the works connected with analyzing MD trajectories. K.K., A.S.-K., and J.K. wrote the manuscript draft. All authors reviewed and discussed the manuscript.

Notes
The authors declare no competing financial interest.

ACKNOWLEDGMENTS
The calculations were carried out at the Faculty of Chemistry of the Jagiellonian University and on the PL-Grid infrastructure at ACK CYFRONET with the support of the project "HPC Infrastructure for Grand Challenges of Science and Engineering". A.J. acknowledges the support from the Polish National Science Centre, project no. 2016/21/D/NZ1/02763. K.K. acknowledges the support of PhD InterDokMed Project no. POW.03.02.00-00-1013/16.

REFERENCES
(1) Ferreira, Q.; Delfino, C.; Morgado, J.; Alcâcer, L. Bottom-up Self-Assembled Supramolecular Structures Built by Stm at the Solid/Liquid Interface. Materials 2019, 12, 382.
(2) Sathyam, P.; Pugazhendhi, S.; Gopalakrishnan, R. Self-Assembled Supramolecular Structure of 4-Dimethylaminopyridinium P-Hydroxy Benzoate Pentahydrate: Synthesis, Growth Optical and Biological Properties. RSC Adv. 2016, 6, 44588–44598.
(3) Tu, Y.; Peng, F.; Adawy, A.; Men, Y.; Abdelmohsen, L. K. E. A.; Wilson, D. A. Mimicking the Cell: Bio-Inspired Functions of Supramolecular Assemblies. Chem. Rev. 2016, 116, 2023–2078.
(4) Rybarska, J.; Piekar ska, B.; Stopa, B.; Zemanek, G.; Konieczny, L.; Roterman, I. Supramolecular Systems as Protein Ligands. In New Kind of Protein Ligand of Antibodies Engaged. In Immune Complex in Self-Assembled Molecules – New Kind of Protein Ligands, Roterman, I.; Konieczny, L. Eds; Springer, 2018.
(5) Piekar ska, B.; Konieczny, L.; Rybarska, J.; Stopa, B.; Zemanek, G.; Szne le, E.; Król, M.; Nowak, M.; Roterman, I. Heat-Induced Formation of a Specific Binding Site for Self-Assembled Congo Red in the V Domain of Immunoglobulin L Chain Lambda. Biopolymers 2001, 59, 446–456.
(6) Konieczny, L.; Piekar ska, B.; Rybarska, J.; Stopa, B.; Krzywka, B.; Noworolski, J.; Pawlicki, R.; Roterman, I. Bis Azo Dye Liquid Crystalline Micelles as Possible Drug Carriers in Immunotargeting Technique. J. Physiol. Pharmacol. 1994, 45, 441–454.
(7) Gaussian 09, R. A. Gaussian Inc.: Wallingford, CT, 2009.
(8) Grimme, S. Semiempirical Gga-Type Density Functional Constructed with a Long-Range Dispersion Correction. J. Comput. Chem. 2006, 27, 1787–1799.
(9) Anderson, J. S. M.; Melin, J.; Ayers, P. W. Conceptual Density-Functional Theory for General Chemical Reactions, Including Those That Are Neither Charge- nor Frontier-Orbital-Controlled. I. Theory and Derivation of a General-Purpose Reactivity Indicator. J. Chem. Theory Comput. 2007, 3, 358–374.
(10) Korchowiec, J.; Uchimaru, T. New Energy Partitioning Scheme Based on the Self-Consistent Charge and Configuration Method for Subsystems: Application to Water Dimer System. J. Chem. Phys. 2000, 112, 1623–1633.
(11) Korchowiec, J.; Uchimaru, T. Mechanism of Addition of Fluoromethyl Radicals to Fluoroethylenes. J. Phys. Chem. A 1998, 102, 6682–6689.
(12) Geerlings, P.; De Proft, F.; Langenaeker, W. Conceptual Density Functional Theory. Chem. Rev. 2003, 103, 1793–1874.
(13) Geerlings, P.; Fias, S.; Boisdevieng, Z.; De Proft, F. Conceptual Dft: Chemistry from the Linear Response Function. Chem. Soc. Rev. 2014, 43, 4989–5008.
(14) Anderson, J. S. M.; Melin, J.; Ayers, P. W. Conceptual Density-Functional Theory for General Chemical Reactions, Including Those That Are Neither Charge- nor Frontier-Orbital-Controlled. I. Theory and Derivation of a General-Purpose Reactivity Indicator. J. Chem. Theory Comput. 2007, 3, 358–374.
(15) Korchowiec, J.; Uchimaru, T. The Charge Transfer Fukui Function: Extension of the Finite-Difference Approach to Reactive Systems. J. Phys. Chem. A 1998, 102, 10167–10172.
(16) Jagusiak, A.; Pańczyk, T. Interaction of Congo Red, Evans Blue and Titan Yellow with Doxorubicin in Aqueous Solutions. A Molecular Dynamics Study. J. Mol. Liq. 2019, 279, 640–648.
(17) Al-Thabaiti, S. A.; Aazam, E. S.; Khan, Z.; Bashir, O. Aggregation of Congo Red with Surfactants and Ag-Nanoparticles in an Aqueous Solution. Spectrochim. Acta, Part A 2016, 156, 28–35.
(18) Humphrey, W.; Dalke, A.; Schulten, K. Vmd: Visual Molecular Dynamics. J. Mol. Graphics 1996, 14, 33–38.
(19) Phillips, J. C.; Braun, R.; Wang, W.; Gumbart, J.; Tajkhorshid, E.; Villa, E.; Chipot, C.; Skeel, R. D.; Kalé, L.; Schulten, K. Scalable Molecular Dynamics with Namd. J. Comput. Chem 2005, 26, 1871–1892.
(20) Klauda, J. B.; Venable, R. M.; Freites, J. A.; O’Boyle, N. M.; Zheng, C.; Hornak, V.; Mark, A. E.; Todd, J. A.; Darden, T. A.; Merz, K. M.; et al. Open Babel: An Open Source, Cheminformatics Toolkit. J. Cheminf 2012, 4, 36.
(21) Jorgensen, W. L.; Chandrasekhar, J.; Madura, J. D.; Impey, R. M.; Skolnick, J. Comparison of Simple Potential Functions for Simulating Liquid Water. J. Chem. Phys. 1983, 79, 926–935.
(22) Jorgensen, W. L.; Chalasani, K.; Madura, J. D.; Impey, R. W.; Klein, M. L. Comparison of Simple Potential Functions for Simulating Liquid Water. J. Chem. Phys. 1983, 79, 926–935.
(30) Boys, S. F.; Bernardi, F. The Calculation of Small Molecular Interactions by the Differences of Separate Total Energies. Some Procedures with Reduced Errors. *Mol. Phys.* **1970**, *19*, 553–566.

(31) Tomasi, J.; Mennucci, B.; Cammi, R. Quantum Mechanical Continuum Solvation Models.*Chem. Rev.* **2005**, *105*, 2999–3094.

(32) Grabowski, S. J. Ab Initio Calculations on Conventional and Unconventional Hydrogen Bonds: Study of the Hydrogen Bond Strength. *J. Phys. Chem. A* **2001**, *105*, 10739–10746.

(33) Koch, U.; Popelier, P. L. A. Characterization of C-H-O Hydrogen-Bonds on the Basis of the Charge-Density. *J. Phys. Chem.* **1995**, *99*, 9747–9754.