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Prediction of self-assembly of adenosine analogues in solution: a computational approach validated by isothermal titration calorimetry

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The recent discovery of the role of adenosine-analogues as neuroprotectants and cognitive enhancers has sparked an interest in these molecules as new therapeutic drugs. Understanding the behavior of these molecules in solution and predicting their ability to self-assemble will accelerate new discoveries. We propose a computational approach based on density functional theory, a polarizable continuum solvation description of the aqueous environment, and an efficient search procedure to probe the potential energy surface, to determine the structure and thermodynamic stability of molecular clusters of adenosine analogues in solution, using caffeine as a model. The method was validated as a tool for the prediction of the impact of small structural variations on self-assembly using paraxanthine. The computational results were supported by isothermal titration calorimetry experiments. The thermodynamic parameters enabled the quantification of the actual percentage of dimer present in solution as a function of concentration. The data suggest that both caffeine and paraxanthine are present at concentrations comparable to the ones found in biological samples at concentrations comparable to the ones found in biological samples.

Introduction

Adenosine is a naturally occurring purine nucleoside present in human cells and known to play a key role in a number of pathologies, including neurological disorders, cardiovascular and autoimmune diseases.^{1–3} The role of adenosine and its interactions with the different receptors have been studied extensively^{4–6} with consequent interests in adenosine analogues with agonist and antagonist activity⁷ and methods that allow fast screening.⁸ Among the different adenosine analogues, caffeine (**Figure 1A**) has been the most extensively studied compound. Research carried out on the health implications of caffeine intake has established important beneficial links to a number of illnesses, including depression,^{9,10} type 2 diabetes, liver disease, Parkinson's disease¹¹ and stroke risk.¹² Caffeine is rapidly absorbed in the stomach and it is metabolised to three primary compounds: paraxanthine (82%), theophylline (11%) and theobromine

(4%).^{13–16} Studies have demonstrated that the key physiological effects of caffeine¹⁷ are the result of interactions effect on the A₁ and A₂ subtypes of the adenosine receptor.¹⁸ The role of caffeine as a cognitive enhancer¹⁹ and also as a neuroprotectant²⁰ have also been reported recently. Caffeine used in combination with melatonin was shown to have antiamyloidogenic action, for the treatment of neurodegenerative diseases.²¹ Borota *et al.* have demonstrated recently that caffeine enhances consolidation of long-term memory in humans.²² There is considerable interest in understanding the mechanism by which caffeine and more generally adenosine analogues can interact with the different receptors, as this may subsequently lead to the identification of new compounds with drug-like activity. In this context, information on a new molecule's behaviour in solution and its potential for self-assembly, particularly its degree of complexation (*i.e.* dimers, trimers, tetramers) and relative distribution, is a key requirement. The self-association of caffeine has been studied experimentally using a variety of techniques including NMR^{23–25} and FTIR spectroscopies,²⁶ dynamic light scattering (DLS),^{27,28} osmometry,²⁹ fluorescence,³⁰ thermal analysis and differential scanning calorimetry (DSC).^{31,32} These methods provide some characterisation although the analysis is often model-dependent. Computational approaches have also been explored to provide a useful tool to understand the behaviour of caffeine. Recent molecular dynamics (MD) simulations of aqueous caffeine solutions by Tavagnacco *et al.* showed extensive aggregation, with caffeine molecules 'stacking their flat faces against one another like coins'.^{33–35}

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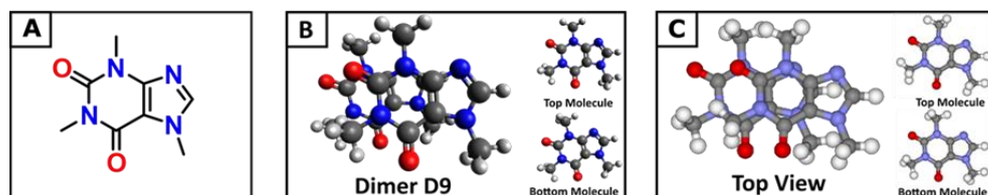


Fig. 1 A) The skeletal chemical structure of caffeine. B) The most stable dimer structure (D9) in water as calculated using the proposed computational approach. C) The dimer structure of caffeine as found in the solid state.^{67,68} For clarity the reciprocal orientations of the top and bottom molecules in the dimer in (B) and (C) are reported separately.

These simulations were, however, conducted at the limit of solubility of caffeine in water (about 16 mg.mL⁻¹) which is approximately five orders of magnitude higher than the concentration range at which caffeine is present in biological systems (0.002 mg.mL⁻¹ in plasma).^{36–38} Such low concentrations would require simulation at the limit of MD techniques, containing hundreds of millions of solvent molecules.

The continuum solvation method is an alternative technique for the determination of the structure and thermodynamic stability of molecular aggregates in solution,³⁹ where the solute (monomer, dimer, trimer, etc.) is treated at the density functional theory (DFT) level and the solvent is described using a polarizable continuum solvation model. Using this approach Senthilnithy *et al.*⁴⁰ and Bradley and Henderson⁴¹ report highly positive (+12 kcal.mol⁻¹) dimerization free energies. However, the B3LYP density functional employed in these studies does not allow dispersion interactions to be described fully,⁴² which can affect accuracy. The availability of a computational approach that can accurately predict changes in free energies, because of small structural alterations, would accelerate the identification of new candidate drugs. In this work, we present a quantum mechanical continuum solvation approach as a computational tool that allows to predict the behaviour in solution of adenosine-analogues (AA) by determining the structure and thermodynamic stability of their molecular clusters (AA)_n (*n* = 2–4), in solution, at concentrations comparable to the ones found in biological samples. Caffeine was initially used as model molecule; the same tool was then employed to predict the effect of small structural variations in the thermodynamics of molecular aggregation by studying paraxanthine (1,7-dimethylxanthine). The computational approach was then validated experimentally by using isothermal titration calorimetry, (ITC) to determine the association enthalpies for both molecules. The thermodynamic data were then used to assess the presence and distribution of potential clusters.

Computational methods

Density functional theory methods

Electronic structure calculations were carried out with the NWChem (version 6.3)⁴³ and Gaussian09⁴⁴ codes. The B97 density functional together augmented with the D2 Grimme's dispersion correction (B97-D2)⁴⁵ and the 6-31+G(d,p) basis set⁴⁶ was chosen to conduct geometry optimization and compute

thermal contributions to the gas-phase free energy. The B97-D2 functional was chosen because it is a suitable method to describe molecular complexes with predominant hydrogen-bonding and dispersion contribution.^{47–50} The 6-31+G(d,p) basis set was chosen because when evaluating binding energies, the addition of diffuse functions on non-hydrogenic atoms to a doubly polarized valence double- ζ basis set provides a good compromise between accuracy and computational cost.^{51–53} The B97-D2/6-31+G(d,p) method, together with the combination of other functionals and basis sets, was also tested in terms of its ability to locate the “face-to-face” and “face-to-back” dimeric motifs found in the crystallographic structure of caffeine (Section S.1).

The effect of the basis set superposition error (BSSE) was not considered in our simulations. However, the effect of increasing the size of basis set from split-valence double-zeta (6-31+G(d,p)) to split-valence triple-zeta (6-311+G(2df,2p)) on the gas-phase binding energy is only 0.2 kcal.mol⁻¹ (Section S.2). In addition, Di Tommaso and Watson previously demonstrated, using a similar methodology, that the contribution of BSSE to the dimerization constant of carboxylic acids in solution is less than 0.2 unit, when evaluated using the counterpoise correction (CPC) method of Boys and Bernardi,⁵⁵ to the dimerization constant (*pK_D*) of carboxylic acids in solution is less than 0.2 unit. Given that the cost of calculating the CPC is higher than that of the original gas-phase interaction energy,⁵⁶ we have not included this term in the evaluation of the gas-phase energy of caffeine and paraxanthine clusters.

The universal Solvation Model based on solute electron density (SMD)⁵⁷ together with the M06-2X density functional⁵² and the 6-31+G(d,p) basis set were employed to compute the hydration free energies.⁵⁷ Originally tested over a set of 2892 solvation free energies and transfer free energies for both neutral and ionic species in water and non-aqueous solutions, the mean unsigned error obtained with the SMD model over 26 combinations of various basis sets and density functionals was 0.8 kcal mol⁻¹ for neutral solutes.⁵⁸

Overall, we expect that the computed Gibbs free energies of reaction have an uncertainty of ± 1.1 kcal mol⁻¹ (Section S.2). Further calculations to assess the methodology and conduct electron density analysis were also performed using the plane-waves Quantum ESPRESSO⁵⁹ software suite with the PBE functional with and without dispersion correction and plane-wave basis set cut offs for the smooth part of the wave functions and the augmented density of 28 and 300 Ry respectively.

Free energies of association in solution

The free energy of formation of caffeine self-associates, (CAF)_n (*n* = 2–4) were computed according to this equation:

$$\Delta G_{\text{ass}}^* = G_{\text{AB}}^* - G_{\text{A}}^* - G_{\text{B}}^* \quad (1)$$

where G_{X}^* is the total Gibbs free energy of caffeine species X (X = monomer, dimer, trimer or tetramer) in the liquid at 298 K. For example, the dimerization free energy (ΔG_{D}^*) was computed as:

$$\Delta G_{\text{D}}^* = G_{\text{dimer}}^* - 2G_{\text{monomer}}^* \quad (2)$$

The term G^* was determined as the sum of the following terms

$$G_{\text{X}}^* = E_{\text{e,gas}} + \delta G_{\text{VRT,gas}}^{\circ} + \Delta G_{\text{sol}}^* + RT \ln[\bar{R}T] \quad (3)$$

In Eq. 3 $E_{\text{e,gas}}$ is the gas-phase total electronic energy of the solute, $\delta G_{\text{VRT,gas}}^{\circ}$ is the vibrational-rotational-translational contribution to the gas-phase Gibbs free energy at *T* = 298 K under a standard-state partial pressure of 1 atm. ΔG_{sol}^* is the solvation free energy of the solute corresponding to transfer from an ideal gas at a concentration of 1 mol.L⁻¹ to an ideal solution at a liquid-phase concentration of 1 mol.L⁻¹, $RT \ln[\bar{R}T]$ is the free energy change of 1 mol of an ideal gas from 1 atm to 1 mol.L⁻¹ ($RT \ln[\bar{R}T] = 1.89 \text{ kcal.mol}^{-1}$ at *T* = 298.15 K and $\bar{R} = 0.082052 \text{ K}^{-1}$).

The terms $E_{\text{e,gas}}$ and $\delta G_{\text{VRT,gas}}^{\circ}$ were computed at the B97-D2/6-31+G(d,p) level of theory, whereas the hydration free energy component, ΔG_{sol}^* , was determined using the SMD/M06-2X/6-31+G(d,p) solvation model using the gas-phase B97-D2/6-31+G(d,p) optimised geometries. Although an approach that includes nuclear relaxation in solution of the solute is supposed to improve a method that uses gas-phase geometries, the hydration free energy (ΔG_{sol}^*) of caffeine calculated as the difference between the solute molecule's energy in solution (SMD/M06-2X/6-31+G(d,p)) at the geometry optimized in gas-phase (B97-D2/6-31+G(d,p)) and the gas phase energy (M06-2X/6-31+G(d,p)) at the geometry optimized in the gas phase ($\Delta G_{\text{sol}}^* = -12.8 \text{ kcal.mol}^{-1}$) is in excellent agreement with the experimental value of $-12.1 \text{ kcal.mol}^{-1}$ (Table S.3). The value of ΔG_{sol}^* of caffeine changes only slightly if the hydration free energy is calculated as a difference between the caffeine's energy in solution at the geometry optimized in solution and the gas phase free energy at the geometry optimized in the gas phase ($\Delta G_{\text{sol}}^* = -13.2 \text{ kcal.mol}^{-1}$) (Table S.3). This approach also follows the recommendation by Ho et al. that free energies in solution should be obtained from separate gas- and solution-phase calculations,⁶⁰ because well-established equations based on the statistical thermodynamics of an ideal gas, under the harmonic oscillator rigid rotor approximation, have been developed to calculate gas-phase partition functions and associated thermodynamic functions (entropy, enthalpy, free energy) from the geometries, frequencies, and energies derived from standard ab initio calculations. If a structure is optimised in the gas-phase, then the thermal contribution is obtained by applying the same ideal gas partition functions to the frequencies calculated in a dielectric continuum. This approach

is questionable because the translational and rotational contributions, are unlikely to be valid in solution, and it should only be when stationary points in the solution do not correspond to stationary points in the gas-phase.⁶¹

Localisation of energy minima on the PES of dimers

The potential energy surfaces (PES) of weakly interacting molecular complexes is characterised by a large number of minima. The most stable isomers of the caffeine dimer, (CAF)₂, were located using modified version of the computational protocol developed by Di Tommaso and Watson:⁵⁴ (i) Hundreds of thousands of caffeine dimeric configurations were initially generated using the Granada software.⁶² (ii) A first selection of the caffeine dimers was based on geometrical considerations: only those configurations with caffeine molecules at a distance of less than 4.0 Å were considered for subsequent calculations. (iii) The energies of the configurations generated at step (ii) were computed at the B97-D2/6-31+G(d,p) level of theory and used to calculate the Boltzmann factor (f_i):

$$f_i = \frac{e^{-(E_i - E_0)/RT}}{\sum_j e^{-(E_j - E_0)/RT}} \quad (4)$$

where E_i is the energy of the *i*th candidate dimeric structure, E_0 is the energy of the most stable generated dimer, the index *j* runs over all selected configurations, *R* is the universal gas constant and *T* is the absolute temperature, which was set at 300 K. (iv) The 110 candidate structures with a Boltzmann factor $f_i \geq 0.01$ were subject to full geometry optimization, and calculation of thermochemical properties and solvation energies. (v) The Boltzmann averaged free energy of association was then calculated according to the following equation:

$$\Delta G_{\text{D}}^* = \sum_i f_i \cdot \Delta G_{\text{D}_i}^* \quad (5)$$

where f_i is the Boltzmann factor calculated according to eq. (4) and $\Delta G_{\text{D}_i}^*$ is the association free energy for the *i*th dimeric configuration. The sum runs over all the selected dimers.

The procedure to localise minima on the PES of (CAF)₂ involved 13,000 single point energy calculations (~154k CPU hours using NWChem and 288 cores), followed by 95 structure optimizations (~13k CPU hours using Gaussian09 and 12 cores), frequency calculation (~8k CPU hours using Gaussian and 12 cores) and solvation energy evaluation (~300 CPU hours using Gaussian09 and 12 cores). The total number of CPU hours for the caffeine dimer at the B97-D2/6-31+G(d,p) level of theory was more than 175k CPU hours.

Experimental methods

Caffeine (1,3,7-Trimethylxanthine), paraxanthine (1,7-dimethylxanthine) were purchased from Sigma-Aldrich as anhydrous powder. GC grade methanol (Sigma-Aldrich) was employed for recrystallisation. Aqueous solutions were prepared in distilled water. For ITC experiments the solutions were sonicated, using an ultrasound bath (Sonorex Super

by BANDELIN electronic GmbH & Co. KG) for 5 min followed by degassing with ThermoVac (MicroCal, Malvern Instruments).

X-Ray Diffraction Data

Paraxanthine was recrystallized from methanol by slow evaporation at room temperature. A translucent colourless block-like crystal, approximate dimension 0.080 x 0.150 x 0.300 mm, was used for X-Ray crystallographic analysis. A total of 1088 frames were collected. The total exposure time was 3.02 hours. The frames were integrated with the Bruker SAINT software package using a narrow-frame algorithm. Crystal data at 100(2)K: C₇H₈N₄O₂, Mr=180.17, monoclinic crystal system. The final cell constants of a=7.376(7) Å, b=13.329(13) Å, c=8.193(8) Å, β=113.50(2)°, Volume=738.7(13) Å³, space group P2₁/n, Z=4, calculated=1.620 g cm⁻³, μ=0.124mm⁻¹, (R_{int}=0.0273) reflections collected within the θ range 3°–28°. The Final anisotropic full-matrix least-squares refinement on F₂, based on 7147 with I>2σ(I) and 120 variables converged at R₁=0.0330 for the observed data, wR₂=0.0914 for all data and Goodness-of-fit=1.032.

Dissociation experiments by isothermal titration calorimetry

Isothermal titration calorimetry (ITC) measurements were carried out on a VP-ITC Microcalorimeter (MicroCal, Malvern Instruments) at 25°C. A solution of caffeine (82.4mM) or paraxanthine (5.55 mM) was loaded in the syringe and stepwise injected in the cell, in degassed H₂O. The dilution effect when gradually injecting the concentrated solution of caffeine or paraxanthine in H₂O made the aggregates to dissociate and the heat generated was measured. The reference cell was also filled with distilled H₂O. The reference power was set to 4 μcal/sec; initial delay 60 sec; stirring speed 400 rpm; equilibration options were set to fast equilibration; feedback mode high. For both experiments the initial injection was 5 μL and the rest aliquots were of 15 μL with 300 s time spacing between each injection to assure signal returning to the baseline. The filter period was 2 s. The duration of the first injection was 10 s and 30 s for the following ones. The total number of injections was 30. The enthalpy (ΔH_{diss}) and equilibrium constant (K_{diss}) of the dissociation in solutions containing either caffeine or paraxanthine were obtained by integrating each heat peak in the raw calorimetric data (Figures 7 and 9) using a dissociation curve fitting model. The integrated heat plots were acquired (Figures S89 and S101) and the values of ΔH_{diss} and K_{diss} were obtained together with their associated errors. The association constant (K_{ass}), the Gibbs free energy (ΔG_{diss}) and the entropy (ΔH_{diss}) changes were calculated according to the equation $K_{ass} = 1/K_{diss}$, $\Delta G_{diss} = -RT \ln K_{diss}$, and $\Delta G_{diss} = \Delta H_{diss} - T\Delta S_{diss}$, where $R = 1.987 \text{ cal}\cdot\text{mol}^{-1}\cdot\text{K}^{-1}$ (gas constant) and $T = 298 \text{ K}$.

Results and Discussion

Caffeine dimers

Dimers are the first aggregates that can form during the process of self-association and their structure often corresponds to the

building unit of the crystal that grows from solution.^{63–66} The chemical nature of the intermolecular interactions in the caffeine dimer (CAF)₂ can therefore be expected to control its aggregation. The rearrangement of the electronic charge of the face-to-face and face-to-back dimeric configurations found in the X-ray crystalline structure of caffeine^{67,68} (Figures S.4.b and S.4.b') confirms the presence of six C-H...O and C-H...N intermolecular hydrogen bonds between the two caffeine molecules.^{69–72} Although the difference in electronegativity between C and H is small, the weak hydrogen bonds between caffeine molecules play an important role, together with the π-π stacking of the aromatic rings, in controlling the self-association process.⁷³ To evaluate all possible interactions, a computational approach based on probing the potential energy surface of caffeine dimers was applied. Nine low energy configurations for the caffeine dimer (D1-9) were identified using this approach (Figure S5), five of which had never been reported before.^{40,69} All these structures present the CH₃...X (X = N, O) interactions between the two caffeine molecules, therefore confirming their key role. Overall, the gas-phases energy ($\Delta E_{e, gas}$) and Gibbs free energy (ΔG_D^*) of dimerization were found to be negative. However, in aqueous solution the formation of dimers D1-8 is thermodynamically unfavourable (Table 1). The exception is represented by dimer D9 (Figure 1B), whose liquid-phase reaction is slightly exergonic ($\Delta G_D^* = -0.4 \text{ kcal}\cdot\text{mol}^{-1}$). This therefore suggests that dimer D9 can exist in aqueous solutions. There is much debate on the structural relationship between the most populated dimers in solution and the structural motifs found in solid state forms.^{53,64,72} Figures 1B and 1C highlight how the structure of the most stable caffeine dimer in water (D9), obtained computationally, correlates very well with the structural synthon found in the X-ray crystal structure (Figure 1C). On this basis, it can be affirmed that the aggregation of caffeine obeys the so-called "link hypothesis",⁷² according to which the transfer of structural information from the liquid- to the solid state-phase is linked to the presence in solution of stable "face-to-back" dimers.

For dimer D9 to have the lowest free energy in solution, Table 1 suggests that it has a contribution of approximately $-2 \text{ kcal}\cdot\text{mol}^{-1}$ from the hydration free energy. This contrasts with dimers D1–D8, whose solvation correction are all positive, with a range of 1.6 to 4.5 kcal.mol⁻¹. Assuming a continuum representation of the solvent, the solvation free energy (ΔG_{solv}^*) is given by two components:⁵⁷ the electronic polarization term (ΔG_{EP}^*), associated with the electrostatic interaction between the solute and the solvent; the free energy change contribution (G_{CDs}^*) from solvent cavitation (C), changes in dispersion (D) energy, and possible changes in local solvent structure (S). For all dimers, the G_{CDs}^* contribution to the solvation energy is between 8.7 and 9.2 kcal.mol⁻¹. Consequently, the extra-stabilization in solution of D9 is due to the electronic polarization term. In this configuration, the monomers are almost superimposed (Figure 1A) and this configuration does not maximize the attractive interaction between electropositive and electronegative parts of the two caffeine molecule. The gas-phase interaction between the two caffeine molecules ($\Delta E_{e, gas} = -12.87 \text{ kcal}\cdot\text{mol}^{-1}$) is lower than in dimers D1-D8 (-17

$\text{kcal.mol}^{-1} < \Delta E_{e, aq} < -19 \text{ kcal.mol}^{-1}$) and the extra-stabilization due to the higher electrostatic interaction between **D9** and the polarizable solvation model.

The effect of changes to the solvation environment on caffeine dimerization was also investigated by computing the formation of $(\text{CAF})_2$ in three different systems: (i) methanol, a polar protic solvent with a static dielectric constant (ϵ) lower than water; (ii) acetone, a polar aprotic solvent, and (iii) chloroform, an non polar aprotic solvent. The Boltzmann averaged Gibbs free energy of dimerization obtained for all three systems (Table S.5) shows an inverse correlation with the dielectric constants.

Table 1. Energetics for the formation of the nine low-energy minima dimer configurations in the gas-phase and aqueous solution as computed at the B97-D2/6-31+G(d,p) theory level. Free energies of hydration computed using the SMD/M06-2X/6-31+G(d,p) solvation model. Values in kcal.mol^{-1} .

Dimer	$\Delta E_{e, aq}$	ΔG_D^g	ΔG_D^a
D1	-18.89	-2.03	2.48
D2	-18.74	-1.96	2.45
D3	-17.57	-1.93	1.09
D4	-18.51	-1.60	3.15
D5	-18.53	-1.52	2.96
D6	-18.91	-1.52	2.80
D7	-17.21	-0.79	2.15
D8	-17.31	-0.57	1.04
D9	-12.87	1.63	-0.39

Caffeine clusters

Given the existing literature data on higher order clusters for caffeine, the next step focused on the stability of these systems.^{28,35} The configuration of the tetramer was obtained by stacking two dimers of type **D9** (the most stable in water) on top of each other. The configuration of the trimer was then obtained by removing one caffeine molecule from the $(\text{CAF})_4$. The optimised structures of the higher order clusters $(\text{CAF})_3$, and $(\text{CAF})_4$ are reported in Figure 2.

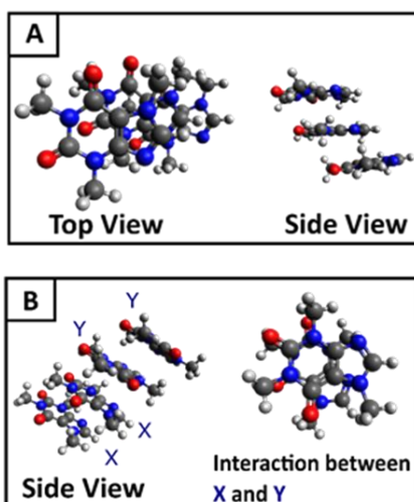


Fig. 2. Optimized structures of trimer (A) and tetramer (B) of caffeine. For the trimer a top and a side view are reported, which show that the reciprocal orientation of the molecules is retained. In the case of the tetramer, the first two neighbor molecules have an identical orientation while the second two caffeine are rotated, leading to a final geometry of the type (XX-YY). The interaction geometry between X and Y is reported.

Two reaction schemes were used to compute the thermodynamic stability of the trimer in solution: i) the condensation of **D9** with an additional caffeine molecule, $(\text{CAF})_{D9} + \text{CAF} \rightarrow (\text{CAF})_3$; ii) the aggregation of three isolated CAF molecules, $3\text{CAF} \rightarrow (\text{CAF})_3$. For the tetramer $(\text{CAF})_4$, we considered the condensation of two **D9** dimers, $2 \text{CAF}_{D9} \rightarrow (\text{CAF})_4$, as well as the association of one caffeine molecules to the trimer structure $(\text{CAF})_3 + \text{CAF} \rightarrow (\text{CAF})_4$. The energetics of formation of the caffeine trimer and tetramer are listed in Table 2 and show that the free energies of formation of these two species are approximately $+2 \text{ kcal.mol}^{-1}$. Trimers and tetramers are unstable in water, compared to both monomer and dimer **D9**, and unlikely to be present in caffeine aqueous solutions at the concentrations found in biological samples.

Table 2. Energetics [kcal/mol] for the formation of caffeine trimer and tetramer forms in the gas-phase and aqueous solution as computed at the B97-D2/6-31+G(d,p) theory level. Free energies of hydration computed using the SMD/M06-2X/6-31+G(d,p) solvation model. Values in kcal.mol^{-1} .

Species	Reaction	$\Delta E_{e, aq}$	ΔG_{ass}^g	ΔG_{ass}^a
Trimer	$3 \text{CAF} \rightarrow (\text{CAF})_3$	-26.77	4.83	2.14
	$(\text{CAF})_2 + \text{CAF} \rightarrow (\text{CAF})_3$	-13.90	3.20	2.52
Tetramer	$(\text{CAF})_2 + (\text{CAF})_2 \rightarrow (\text{CAF})_4$	-19.99	-0.24	2.04
	$(\text{CAF})_3 + 2\text{CAF} \rightarrow (\text{CAF})_4$	-32.86	1.39	1.66

Paraxanthine dimers

Understanding the molecular structure of the analogues and their self-association behaviour is an important requirement for the evaluation of interactions with biological receptors.⁷⁴

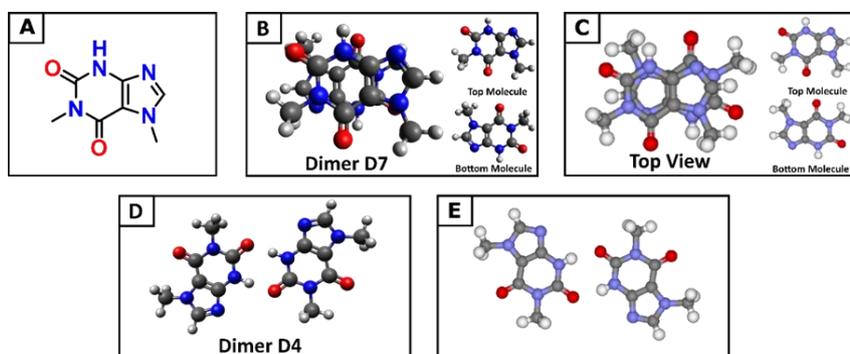


Fig. 3 A) The skeletal chemical structure of paraxanthine. B) Stable dimer structure in gas phase as calculated using the proposed computational approach. C) The π - π stacking dimer synthon of paraxanthine as found in X-Ray structure. For the dimer structures (B, C) the top and bottom molecules are reported separately to clarify the reciprocal orientation. D) Stable dimer structure linked by H-bond interactions as calculated using the proposed computational approach. E) The later dimer synthon of paraxanthine as found in X-Ray structure.

Paraxanthine (1,7-dimethylxanthine) is the major primary metabolite of caffeine, and although the two molecules are very similar in structure, the removal of one methyl group results in a tenfold decrease in the limit of solubility in water.⁷⁵ Compared with the nine low energy dimeric configurations of caffeine, the DFT tool identified 20 stable paraxanthine dimers (Figure S.6), showing the significant effect of small structural changes in caffeine analogues to the process of self-association. Two types of structures can be identified: (i) dimers **D1**, **D2**, **D4** and **D13**, where the two paraxanthine units are linked by two lateral H-bonds; (ii) all remaining (PX)₂ structures, where the interaction is controlled by π - π stacking. Dimers **D4** and **D7** correspond to the synthon in the crystallographic structure of paraxanthine (Figure 3). The agreement between the simulated building units of paraxanthine in solution and its solid-state structure confirms the efficiency of the computational tool here adopted to identify possible dimeric structures of caffeine-analogues. In the gas phase most dimers are thermodynamically stable but in aqueous solution only the configurations forming π - π stacking interactions have a negative free energy of formation ($-1.0 < \Delta G_p^* < -0.1$ kcal.mol⁻¹) (Table S8). In water the Boltzmann averaged value for the dimerization of paraxanthine is -0.11 kcal.mol⁻¹. The comparison of the computed thermodynamic stability of the dimers of caffeine and paraxanthine shows that, despite their very small structural difference, the free energy of formation of (PX)₂ is 1.1 kcal.mol⁻¹ lower than (CAF)₂.

Experimental ITC verification and solution speciation

ITC experiments were conducted to obtain the enthalpy and equilibrium constant of dissociation of caffeine and paraxanthine in aqueous solution (Table 3). The raw calorimetric data are characterised by positive heat peaks (Figures S8 and S10) indicating that the dissociation for both caffeine and paraxanthine is an endothermic process ($\Delta H_{\text{diss}} > 0$). Fitting of these data to a dimer dissociation model provides constants for both caffeine ($K_{\text{diss}} = 0.112 \pm 3.3 \times 10^{-3}$ M) and paraxanthine ($K_{\text{diss}} = 0.03 \pm 1.8 \times 10^{-3}$ M). In both cases, the Gibbs free energy of association ($\Delta G_{\text{ass}} < 0$) is negative. The ΔG_{ass} value obtained using ITC measurements (-1.3 kcal.mol⁻¹) is in

excellent agreement with a previously reported DSC study ($\Delta G_{\text{ass}} = -1.5$ kcal.mol⁻¹ at 25 °C).⁷⁶ The free energies of dimerization obtained computationally for caffeine ($\Delta G_{\text{dim}} = +0.92$ kcal.mol⁻¹) and for paraxanthine ($\Delta G_{\text{dim}} = -0.11$ kcal.mol⁻¹) are approximately 2 kcal.mol⁻¹ higher than the experimental values obtained from the ITC experiments (Table 2). However, if the errors associated with the use of the B97-D2/6-31+G(d,p) method together with the SMD/M06-2X/6-31+G(d,p) solvation model [$\epsilon(\Delta G_{\text{dim}}^*) = \epsilon(\Delta G_{\text{ass}}^*) + \epsilon(\Delta \Delta G_{\text{solv}}^*) = 1.1$ kcal.mol⁻¹] are taken into account (Section S.2) there is a good agreement between computational and experimental results. The small values obtained for the free energy of dimerization of both caffeine and paraxanthine (< 2 kcal.mol⁻¹) also confirm that the monomer and dimer can coexist in solution. The results from the computational approach are considerably closer to the experimental values, and represent a significant improvement compared to previously published data, which reported free energies of dimerization ($+12$ kcal.mol⁻¹) considerably higher.^{40,41} Furthermore the difference in Gibbs free energy of association between caffeine and paraxanthine ($\Delta \Delta G_{\text{ass}} = \Delta G_{\text{ass}}^{\text{CAF}} - \Delta G_{\text{ass}}^{\text{PX}}$) obtained by ITC (0.6 kcal.mol⁻¹) and DFT (1.1 kcal.mol⁻¹) is also in excellent accordance, providing strong evidence for the use of our computational approach to predict the self-association behaviour in solution of adenosine-analogues.

Table 3. Comparison of the enthalpy and free energy of association obtained by ITC measurements, and the dimerization free energy obtained using DFT calculations at 25 °C. Values in kcal.mol⁻¹.

Molecule	ΔH_{ass} (ITC)	ΔG_{ass} (ITC)	ΔG_p^* (DFT)
Caffeine	-5.25 ± 0.04	-1.30 ± 0.71	$+0.92 \pm 1.10$
Paraxanthine	-2.83 ± 0.07	-1.89 ± 0.35	-0.11 ± 1.10

Having demonstrated that at low concentrations both caffeine and paraxanthine are potentially present in solution only as a combination of monomers and dimers, the experimental and computational values of ΔG_{ass} were used in the monomer-dimer model to estimate the percentage of dimer in solution as a function of total concentration:⁷⁷

$$[(CAF)_2] = \frac{\{-1 + (1 + 8K_D[C])^{1/2}\}^2}{16K_D} \quad (4)$$

Equation (4), shown for caffeine, was used, where K_D is the dimerization constant, $K_D = [(CAF)_2]/[CAF]^2$, $[C]$ is the stoichiometric concentration of the solute, $[C] = [CAF] + 2[(CAF)_2]$, and $[CAF]$ and $[(CAF)_2]$ are the monomer and dimer concentrations, respectively (mol L^{-1}).

Figure 4 shows the computational and experimental profiles of the dimer percentage in solution for caffeine and paraxanthine in the concentration range $5\mu\text{M}$ – $50\mu\text{M}$, which is close to the one found in biological samples. The computational data consistently suggest a lower percentage of dimer in solution compared to ITC for both paraxanthine and caffeine. This is due to the value of ΔG_D^\ddagger obtained from DFT calculations, for both molecules, being about 2 kcal.mol^{-1} more positive than ΔG_{ass} obtained from ITC measurements. For both approaches, computational and ITC, paraxanthine is found to be present as dimer in higher concentrations compared to caffeine, which may be related to the differences in solubility of the two molecules (PX = 5mM , CAF 82mM). This behaviour is observed not only at very low concentrations, but also towards the limit of solubility (Figure S11). The results therefore indicate that both paraxanthine and caffeine, when evaluated at the concentrations found in biological samples ($10\mu\text{M}^{36}$), are predominantly present as monomers, with less than 0.05% dimer in solution.

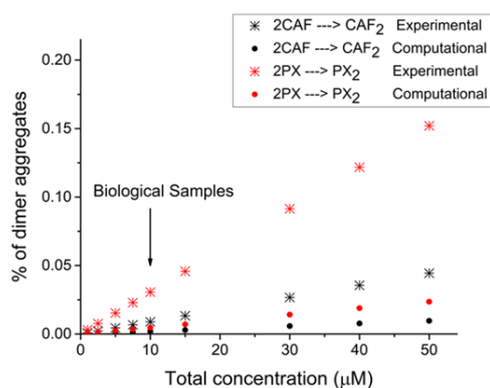


Fig. 4 Comparison between the computational and experimental profiles of the population of dimers in solution at different total concentrations of caffeine and paraxanthine (μM). The computational points were determined using the Gibbs free energy of dimerization in water of the most stable dimers: caffeine (black circles) and paraxanthine (red circles). The experimental points were calculated using the Gibbs free energy of association determined using ITC experiments: caffeine: (black star) and paraxanthine (red star).

Conclusions

We presented a DFT-based computational tool that allowed the study of the self-assembly behaviour in solution of two

adenosine-analogues like caffeine and paraxanthine, and we demonstrated that for both molecules the formation of higher order clusters is thermodynamically disfavoured. The results were validated experimentally by ITC measurements, which provided values for the Gibbs free energy of association consistent with the DFT results. Significantly we showed how the computational approach allows to predict the effect of small structural changes on the chemical structure and thermodynamic stability of clusters resulting from self-assembly. In low concentration solutions, both caffeine and paraxanthine are mainly present as monomers, with less than 0.05% dimer present in $10\mu\text{M}$ solutions. The computational method used here is a valid tool to quantify the speciation of adenosine-analogues and predict accurately their behaviour in solution, which can provide a standardised and faster approach to the evaluation of potential new drug leads.

Conflicts of interest

There are no conflicts to declare.

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