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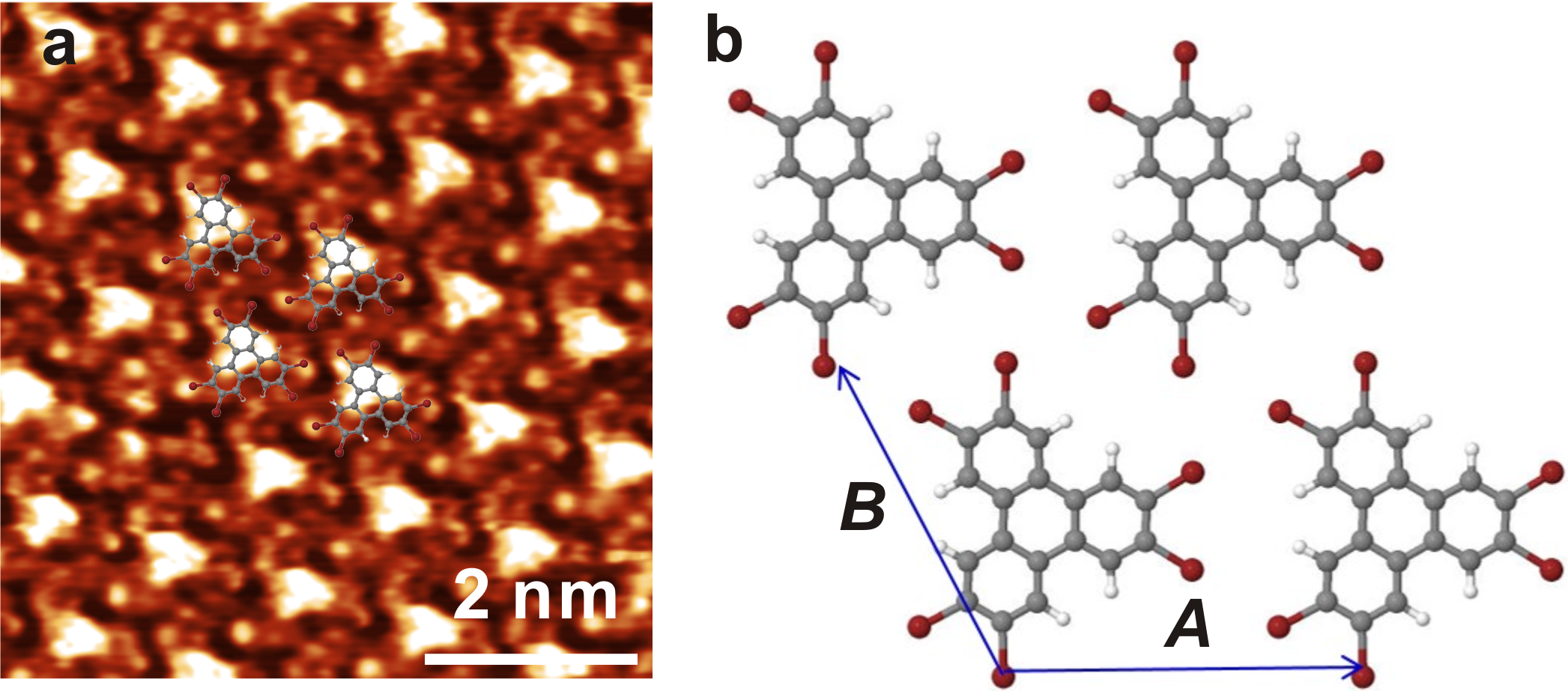
**Thermodynamics of halogen bonded monolayer   
self-assembly at the liquid-solid interface**

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Monolayer self-assembly of a hexabrominated, three-fold symmetric aromatic molecule is studied at the heptanoic acid–graphite interface. Thermodynamical insights are obtained from an adapted Born-Haber cycle that is utilized to derive the overall enthalpy change including solvent effects. Comparison with theoretical entropy estimates suggests a minor influence of solvation.

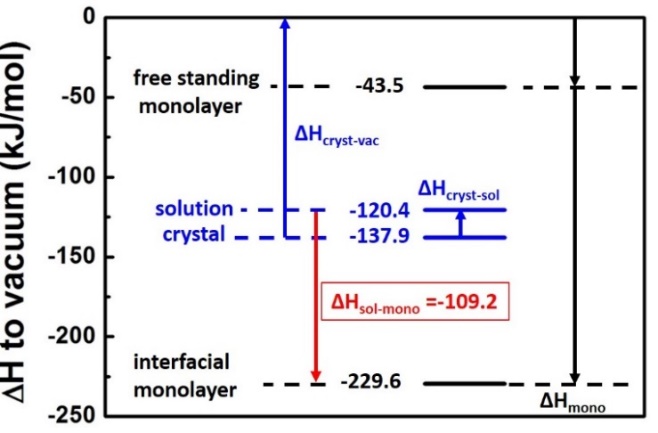
Halogen bonds have gained attention in crystal engineering, supramolecular self-assembly, and even computer aided drug design.1-4 Halogen bonds are viewed as a separate class of donor-acceptor type interactions that originate in the anisotropic charge distribution around halogen substituents, most importantly Br and I. The so called -hole gives rise to a positive electrophilic cap at the opposing pole of the -bond that is balanced by a nucleophilic equatorial ring of negative charge.5-7 A halogen bond is formed through a net attraction between the electrophilic cap of the halogen with either the nucleophilic part of a different entity, as for instance the nitrogen electron lone pair, or the nucleophilic ring of another halogen. This characteristics results in directionality and selectivity, whereas the halogen bond strength decreases as the halogen electronegativity increases. Moreover, a single halogen substituent can simultaneously act as both halogen bond acceptor and donor – an important distinction from the in other aspects quite comparable hydrogen bonds. A triangular arrangement of three halogen atoms with three cyclic halogen bonds is a frequently encountered motif both in bulk crystals and on surfaces,8, 9 but alternative configurations were similarly reported.10

Supramolecular self-assembly of halogen bonded systems on surfaces was recently studied by scanning tunneling microscopy (STM) both under ultra-high vacuum9, 11, 12 and ambient conditions.10, 13-15 Accompanying density functional theory simulations based on experimental structures provided insights into the energetics.10, 12, 16 While this is an important first step, a fundamental understanding of self-assembly from solution requires a full thermodynamical assessment, including the role of the solvent. As a model system hexabromotriphenylene (HBTP, cf. Fig. 1) is studied at the heptanoic acid (7A)–graphite interface with the aim to quantify all relevant contributions to G. At the liquid-solid interface, the adsorption energy becomes significantly lowered as compared to vacuum by the supernatant liquid phase. This important solvent influence is quantified for the proposed model system by applying an adapted Born-Haber cycle derived from sublimation, dissolution, and monolayer binding enthalpy in vacuum.17



**Fig. 1** (a) High resolution STM image of a HBTP monolayer at the 7A-graphite interface. Vsample= -0.40 V, I= 80 pA, (b) DFT optimized geometry of a free standing HBTP monolayer (grey: carbon, red: bromine, white: hydrogen)

At the 7A–graphite interface HBTP self-assembles into a densely packed structure, a high resolution STM image is presented in Fig. 1a. The lattices of HBTP and graphite are aligned (cf. ESI) and the experimental lattice parameters of A=B= (12.5±0.2) Å, γ= 59°±2° match well with a 55 graphite superstructure. Moreover, the absence of a Moiré pattern in these images with pronounced submolecular contrast strongly indicates commensurability. To obtain a detailed structural model, dispersion-corrected DFT simulations of a free-standing HBTP monolayer constrained to a hexagonal lattice were carried out using both empirical dispersion correction (PBE+D) and a van der Waals functional (vdW-DF) (cf. ESI for details). The optimized structure is shown in Fig. 1b. Simulations with unconstrained length of the lattice parameter yield an optimized value of 12.5 Å, i.e. only 1.5 % larger than 12.3 Å of the 55 graphite superstructure. The structure belongs to planar space group p31m, and all bromines take part in the prototypical triangular cyclic arrangement of halogen bonds with a Br-Br distance of 3.70 Å, in good agreement with literature.18 Interestingly, the symmetry of high resolution STM images is significantly lower than that of the DFT structure and does not even reflect the three-fold symmetry of HBTP. This clearly indicates an adsorption site of HBTP with lower local symmetry. Nevertheless, in an overlay STM protrusions can be matched with bromine positions (cf. Fig. 1a). The vdW-DF derived monolayer binding energy per molecule with respect to vacuum of -48.3 kJ/mol is rather small. Compressing the structure to the experimental 55 graphite superstructure reduces the binding energy to -43.5 kJ/mol. A very similar binding energy was obtained using PBE+D: -45.5 kJ/mol for the unconstrained free-standing structure and -42.7 kJ/mol for the constrained 55 superstructure; VdW-DF was also employed to calculate the total binding energy comprised of molecule-molecule and molecule-surface interaction of HBTP in the adsorbed monolayer on graphite and results in Hmono=-229.6 kJ/mol (cf. ESI). Similarly, these calculations yield an adsorption energy of HBTP on graphite of -186.1 kJ/mol per molecule, i.e. the molecule-surface interactions are approximately a factor of 4 larger than the intermolecular interactions.



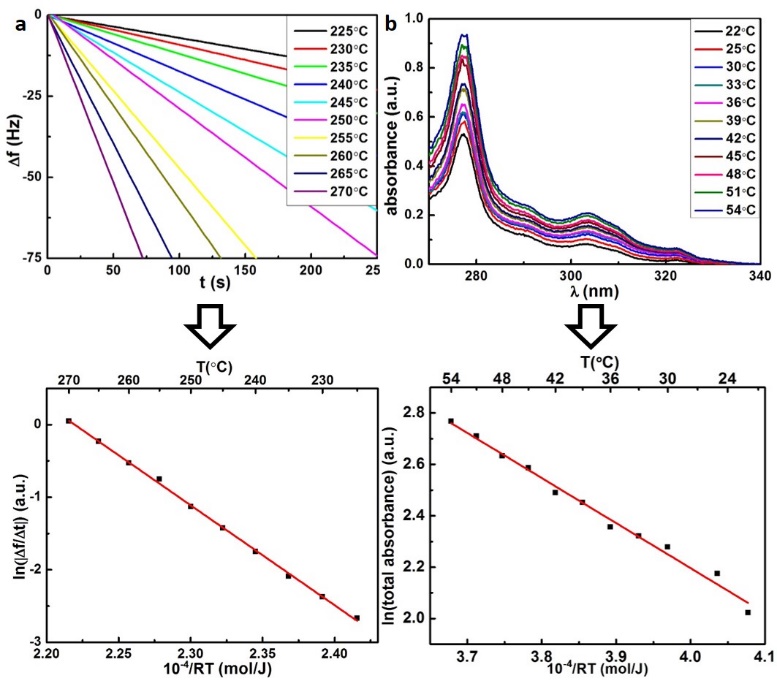
**Fig. 2** Enthalpy diagram of the HBTP–7A–graphite system. Black and blue depicted values and arrows correspond to computational and experimental results, respectively. Vertical arrows denote the respective enthalpies.

In the actual experimental situation at the liquid-solid interface HBTP adsorbs from solution, whereby the effective enthalpy difference Hsol🡪mono is significantly lowered due to solvent interactions. Since a direct measurement is intricate, Hsol🡪mono is indirectly evaluated by combining Hmono with experimentally determined sublimation (Hcryst🡪vac) and dissolution enthalpy (Hcryst🡪sol) as illustrated in Fig. 2.17, 19, 20

Hcryst🡪vac was determined from the temperature dependence of the effusion rate from a Knudsen cell as measured with a Quartz Crystal Microbalance (cf. ESI for details).19 The eigenfrequency decline f vs. time t traces are shown in Fig. 3a for crucible temperatures between 225 °C and 270 °C. The rate f/t is proportional to the vapour pressure at the respective temperature. The data yield a perfectly linear Van’t Hoff plot with a slope corresponding toHcryst🡪vac= +137.9±1.6 kJ/mol.

Hcryst🡪sol was determined from the temperature dependence of the solubility as measured by UV-Vis absorption spectroscopy of saturated solutions (cf. ESI for details). Individual absorption spectra are shown in Fig. 3b for temperatures between 22 °C and 54 °C. The absorbance, hence the solubility, increases with temperature indicating endothermic dissolution. The total amount of dissolved HBTP molecules was estimated by integrating over the absorption band from λ = (270..340) nm. The data yield a perfectly linear Van’t Hoff plot with a slope corresponding to Hcryst🡪sol=+17.5±0.6 kJ/mol.

The theoretical (Hmono) and both experimental (Hcryst🡪vac and Hcryst🡪sol) enthalpies are compiled in Fig 2 with isolated molecules in vacuum as a common reference state. From this Hsol🡪mono of -109.2 kJ/mol can be inferred. In principle, this enthalpy difference between solution and monolayer may be overestimated due to a possible contribution from dewetting.17, 20 Self-assembly of a solute monolayer can require desorption of an initially assembled solvent monolayer, resulting in an enthalpy cost and an entropy gain for dewetting. However, an ordered quasi-static monolayer of 7A has never been observed at room temperature, suggesting that it is thermodynamically unstable (G>0). Based on the assumption that the differences in interfacial tension between 7A–graphite and 7A–HBTP monolayer are small in comparison to Hsol🡪mono contributions of solvent dewetting are not further considered, as they do not significantly affect G.



**Fig. 3** Experimental determination of Hcryst🡪vac and Hcryst🡪sol (a) f vs. t traces for different crucible temperatures; (b) UV/Vis absorption spectra of saturated solutions for different solution temperatures; the lower panels depict the corresponding Van’t Hoff plots; the slopes yield Hcryst🡪vac = (+137.9±1.6) kJ/mol and Hcryst🡪sol = (+17.5±0.6) kJ/mol.

For a complete thermodynamical understanding consideration of entropy contributions is inevitable. Since a direct measurement of S is not possible, we propose an indirect evaluation via the critical concentration ccrit, i.e. the lowest solute concentration where self-assembled monolayers are still thermodynamically stable. Since G becomes zero at ccrit, the overall entropy change can be determined from: S=H/T. For HBTP monolayers at the 7A–graphite interface dilution experiments result in ccrit= 28.8±3.7 μmol/L (cf. ESI). Generally, S has several contributions, yet for a rigid molecule such as HBTP translational (Strans) and rotational (Srot) entropy dominate. For molecules within the monolayer Srot and Strans are negligible as compared to the dissolved state, accordingly S can be estimated as the total loss of Srot and Strans. Whitesides et al. propose to estimate Srot with the rigid rotator model, and Strans with the Sackur-Tetrode equation.21 Since the latter was originally derived for gases, the solvent influence has to be implicitly taken into account, e.g. by free volume corrections. Table 1 summarizes Srot and Strans contributions evaluated according to this approach (cf. ESI for details). Interestingly, the entropic contribution -TS=+111.2 kJ/mol is almost similar to the absolute value of |Hsol🡪mono|=109.2 kJ/mol. This perfect match implies that for HBTP self-assembly no further relevant thermodynamic contributions arise from desolvation. These results stand in vast contrast to interfacial monolayer self-assembly of 1,4-benzenedicarboxylic acid and stilbene dicarboxylic acid likewise from fatty acid solution.17, 20 In both cases |Hsol🡪mono| was significantly smaller than -TS for the adsorption of unsolvated solute molecules. Therefore, it was only possible to explain spontaneous self-assembly by including a favourable entropy contribution from desolvation upon solute adsorption, whereby the released solvent molecules regain Strans and Srot. Dicarboxylic acid solute molecules strongly interact with fatty acid solvent molecules through two-fold hydrogen bonds. Accordingly, solute-solvent and solvent-solvent interactions are of comparable strength in these systems. Owing to the high strengths of carboxylic acid hydrogen bonds, formation of a quasi-static solvation shell can be anticipated for dicarboxylic acids dissolved in fatty acids.

Yet, an entirely different type of solvation can be expected for HBTP in 7A. DFT simulations of HBTP + 7A aggregates provide semi-quantitative values of the solvent-solute interaction strength (cf. ESI). Two different configurations were compared: a structure where simultaneously the 7A hydroxyl forms a hydrogen bond with bromine and the carbonyl oxygen a weak hydrogen bond with a phenyl-hydrogen has a bond strength of -34.1 kJ/mol; an alternative structure with 7A on top of HBTP bound only by dispersion forces has a higher bond strength of -49.8 kJ/mol; The bond strength for both configurations is significantly lower than -67.8 kJ/mol for the hydrogen bonds between carboxylic acids.22 Consequently, for HBTP in 7A, solvent-solvent interactions are significantly stronger than solute-solvent interactions, rendering formation of a strongly bound solvation shell unfavourable. These calculations suggest that the predominant interaction for solvation is van der Waals rather than hydrogen bonding. Also, adsorbed HBTP does not need to lose its “on-top” bound 7A molecules and therefore it can partly retain its solvation shell.

|  |  |  |
| --- | --- | --- |
| -TΔStrans(kJ/mol) | -TΔSrot(kJ/mol) | -TΔS(kJ/mol) |
| +66.3 | +44.9 | +111.2 |

Table 1. Estimation of entropy contributions of HBTP in 7A to G. The entropies are evaluated for ccrit at T = 298 K (cf. ESI for details).

In conclusion, HBTP self-assembles into densely packed monolayers at the 7A-graphite interface, where each bromine forms intermolecular halogen bonds in a prototypical triangular cyclic arrangement. Simulations clearly indicate that by far the largest contribution to the stabilization of the monolayer arises from molecule-surface interactions through - bonds between graphite and the aromatic triphenylene core. The overall enthalpy difference for monolayer self-assembly from the supernatant liquid phase was deduced from a Born-Haber cycle. Accordingly, the solvent reduces the binding enthalpy gain to about 50% of the vacuum value, whereas for dicarboxylic acids a significantly more drastic effect was found in previous studies.17, 20 Based on a quantitative comparison of the overall enthalpy and entropy changes, we propose that the contributions from dewetting the substrate and stripping the weakly bound solvation shell cancel each other such that there are no significant net contributions to G. This hypothesis is supported by energetic arguments: 7A-7A solvent-solvent hydrogen bond interactions are markedly stronger than the dispersive 7A-HBTP solvent-solute interactions. The predominant type of solvation is likely to be mediated by the solvent’s interaction with the π-system of HBTP which is not fully lost after adsorption. This profoundly affects G: there is no favourable entropy contribution due to desolvation; since the intermolecular halogen bonds are comparatively weak, strong molecule-surface interactions are required to render self-assembly of halogen bonded monolayers thermodynamically favourable.

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Notes and references

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