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Strong Short-range Cooperativity in Hydrogen-Bond Chains

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Abstract: Chains of hydrogen bonds such as those found in water and proteins are often presumed to be more stable than the sum of the individual H-bonds. However, the energetics of cooperativity are complicated by solvent effects and the dynamics of intermolecular interactions, meaning that such effects are typically derived from theory or indirect structural data. Here, we present direct measurements of energetic cooperativity in an experimental system in which the geometry and number of H-bonds in a chain were systematically controlled. Strikingly, we found that adding a second H-bond donor to form a chain can almost double the strength of the terminal H-bond, while further extension had little effect. The experimental observations add weight to computations which have suggested that strong, but short-range cooperative effects may occur in H-bond chains.

Hydrogen bond chains are prevalent structural motifs in supramolecular and biological systems. H-bonds are widely proposed to exhibit positive cooperativity,[1] which may be manifested by a combination of conformational[5b-6] and electronic effects that may make a chain more stable than the sum of its parts.[2] Such cooperative effects have been shown to influence reactivity,[4] to contribute to the structure, interactions[7] and properties of biomolecules and materials,[8] and to facilitate the communication of chemical information.[9] H-bonded water clusters and chains have been isolated in the solid state[10] and studied experimentally in both liquid and gas phases.[11] Although many nanoscale and bulk properties may be influenced by the cooperativity of H-bonded networks, it is not possible to directly quantify interaction energies from structural or vibrational characteristics. In addition, discussion surrounding the relative contributions of electrostatics, polarization, and covalency in H-bond cooperativity[5b, 9] is further exacerbated by the challenge of considering the influence of the surrounding solvent.

Here, we have employed synthetic molecular balances[10] to directly measure the effect of H-bond chain length on the strength of H-bonding interactions in solution. At the outset of our investigation we identified the phenol, catechol, pyrogallol series (Figure 1B) as a pertinent model system for examining cooperativity in H-bond chains. Indeed, H-bond chains have previously been proposed to contribute to the supramolecular properties of catechol and pyrogallol derivatives.[9b, 11] We reasoned that the pre-organization and proximity of the intramolecular H-bond donors and acceptors in this series of compounds would minimize conformational entropic effects to allow examination of cooperative electronic influences. Initially we measured the experimental complexation free energies of phenol, catechol and pyrogallol with the strong H-bond acceptor, tri-n-butylphosphine oxide using 31P NMR. The binding energies became more favorable as the number of OH groups was increased (Figure 1A). Such a trend could be rationalized by cooperative effects arising from the formation of a linear intramolecular H-bond network between the OH groups (Figure 1B).[11b, 11c] However, the experimental energetic trend shown in Figure 1A was not reproduced in DFT energy calculations for the linear binding mode (Figure 1A cf. solid bars in Figure 3A). Furthermore, experimental evidence obtained in solution and the solid state indicates that catechol-derivatives may bind acceptors in alternative binding modes such as those shown in Figure 1C.[11e, 11f] Thus, we side-stepped this conformational ambiguity by designing a constrained intramolecular system that enabled H-bond energies to be measured specifically at the end of a chain (Figure 2A).

The strength of intramolecular interactions can be assessed using conformational reporters that act as molecular balances.[10] The molecular balances employed in the present study were based on previous designs that enable the measurement of solvent and substituent effects on intramolecular interactions (Figure 2A).[14] The position of the conformational equilibrium in these new balances enables measurement of the energy of the H-bond at the end of a linear chain containing one, two or three H-bonds. These molecular balances were synthesized and found to exist in two conformational states on the NMR timescale at room temperature (see SI for NMR and minimized structures). Conformers were assigned using 2D NMR spectroscopy and the equilibrium constant K determined by integration of the 19F NMR peaks corresponding to each conformer. The difference in the free energy between the conformers was determined using ΔG = -RT lnK. Balance 1H was found to have a strong preference for the conformation in which the C=O – HO interaction was present in CDCl3 (Figure 2B). Strikingly, adding a second H-bond to form a chain (i.e. going from 1H to 2H) approximately doubled the measured ΔG from -4.2 to -8.1 kJ mol⁻¹. However, adding a further H-bond to the chain (2H to 3H) slightly decreased the preference for the H-bonded conformer. This unexpected trend was seen to persist in CD3CN solutions containing up to 10% (v/v) CD3CN (Figure 2B). At higher concentrations of CD3CN the conformations free energies tended to zero due to disruption of the intramolecular H-bonds (Table S3).

Figure 1. Supramolecular complexation energies of phenol derivatives. Experimental free energies for the complexation of tri-n-butylphosphine oxide with phenol, catechol and pyrogallol in CDCl3 and CD3CN. Errors were found to be <1 kJ mol⁻¹ based on titrations performed in duplicate. Data and additional binding experiments with other phenol derivatives are provided in Table S1.

Figure 2. Interaction energy between the OH groups (Figure 1B).[11b, 11c] However, the experimental energetic trend shown in Figure 1A was not reproduced in DFT energy calculations for the linear binding mode (Figure 1A cf. solid bars in Figure 3A). Furthermore, experimental evidence obtained in solution and the solid state indicates that catechol-derivatives may bind acceptors in alternative binding modes such as those shown in Figure 1C.[11e, 11f] Thus, we side-stepped this conformational ambiguity by designing a constrained intramolecular system that enabled H-bond energies to be measured specifically at the end of a chain (Figure 2A).

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Supporting information for this article is given via a link at the end of the document.
Figure 2. (A) Molecular balances and (B) conformational free energies ($\Delta G$) measured in solution at 300 K. (C) Molecular balances used in the (D) Hammett analysis of substituent effects in H-bond chains in CDCl$_3$. Hammett constants were defined relative to the amide, with ortho-OH groups being approximated by $\sigma_p$ (Table S6). Error bars omitted for clarity (Figure S16 shows error bars). $\Delta G_{1xHB}$, $\Delta G_{2xHB}$, $\Delta G_{3xHB}$ approximate the energies associated with chains containing one, two and three H-bonds, respectively.

The data are indicative of a large positive cooperative effect on forming a chain of two H-bonds compared to a single H-bond, while there is little additional change on further increasing the length of the chain. However, the conformational equilibrium shown in Figure 2A may be influenced by secondary substituent effects$^{[14]}$ in addition to the C=O … HO interaction of interest$^{[15]}$. These secondary substituent effects were controlled for using the 0X and 1X series of compounds (Figure 2C) by plotting the sum of the Hammett constants of the X-substituents against the experimental free energies (Figure 2D). The 0X and 1X series formed separate correlations, with the offset approximating the free energy contribution of a single C=O … HO interaction. The steeper gradient of the 1X versus 0X data indicates the sensitivity of the C=O … HO interaction to the electronic effects of the X-substituents (the more electron-withdrawing the substituent, the stronger the H-bond). The free energies for compounds 2H and 3H (blue and purple circles) are vertically displaced from the 0X correlation in Figure 2D by similar amounts ($\Delta G_{2xHB}$ and $\Delta G_{3xHB}$), confirming the minimal energetic effect of extending a H-bond chain beyond two H-bonds, even when background substituent effects are taken into account.

We originally envisaged extending the investigation to include 1,2,3,4-tetrahydroxybenzene derivatives capable of forming a four-membered H-bond chain. However, we found that 1,2,3,4-tetrahydroxybenzene possessed insufficient stability and solubility to facilitate NMR titrations, or the onward synthesis of molecular balances. Instead, we established that B3LYP/6-311G* calculated conformational energies ($\Delta E$) correlated strongly with experimental $\Delta G$ values for all of the balances shown in Figure 2 (Figure S18, $R^2 = 0.99$). Thus, we confirmed that computations provided the opportunity to probe situations that could not be examined experimentally to offer insights into the physicochemical origins of the observed short-range cooperativity. Calculations performed on both the phosphine oxide complexes (Figure 3A) and balances (Figure 3B) exhibited a binary energetic pattern in which there was either one, or more than one, H-bond in the linear chain. The calculations also allowed H-bonds to be deliberately flipped to deliberately break the continuity of the H-bond chain (hashed bars in Figure 3). The dependence of the energies on the number of H-bonds in the chain, rather than the number of OH groups confirmed that the observed cooperative effects originated from the formation of an intramolecular H-bond network, while also ruling out significant contributions from through-bond substituent effects. Furthermore, entropic and conformational differences across the compound series could not account for the observed binary trend observed in both the experiments and computations (Tables S4-5, Figures S13-15,S19-S20). Additional calculations in which an external...
The hydrogen-bond donor could bind in an ideal geometry to the back of the H-bond chains had similar energies (Figure 3C) to the intramolecular cases (Figure 3B). This result confirmed that intramolecular geometric constraints did not account for the lack of additional energetic cooperativity on adding a third or fourth H-bond to chain.

A key finding from our experiments (Figure 2D) and computations (Figure 3) is that adding a second H-bond can, depending on context, almost double the strength of the terminal H-bond interaction. Such doubling of the energy cannot arise exclusively from additive electrostatic field effects since the H-bond interaction. Such doubling of the energy cannot arise exclusively from additive electrostatic field effects since the ends versus the middle of H-bonded chains can be rationalised thus: if similar length-dependent cooperative effects converge much more rapidly at the ends of H-bonded chains than at the chain centre, the ends versus the middle of H-bonded chains can be rationalised thus: if similar length-dependent cooperative effects converge much more rapidly at the ends of H-bonded chains.
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[15] Secondary interactions such as formyl proton to phenol oxygen H-bonds are disfavored by tens of kJ mol$^{-1}$ in calculations and therefore not significantly populated. Similarly, the B3LYP/6-311G* electrostatic surface potentials of the ring edges vary little across the phenol to tetrahydroxybenzene series (111 ± 3 kJ mol$^{-1}$).
Unchained: Experiments and calculations show that H-bond energies can double upon formation of an H-bond chain, but further extension of the chain results in a surprisingly negligible additional cooperative effect.

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