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Hydrogen bond design principles

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Abstract

Hydrogen bonding principles are at the core of supramolecular design. This overview features a discussion relating molecular structure to hydrogen bond strengths, highlighting the following electronic effects on hydrogen bonding: electronegativity, steric effects, electrostatic effects, π -conjugation, and network cooperativity. Historical developments, along with experimental and computational efforts, leading up to the birth of the hydrogen bond concept, the discovery of nonclassical hydrogen bonds (C—H...O, O—H... π , dihydrogen bonding), and the proposal of hydrogen bond design principles (e.g., secondary electrostatic interactions, resonance-assisted hydrogen bonding, and aromaticity effects) are outlined. Applications of hydrogen bond design principles are presented.

Keywords

aromaticity; hydrogen bonding; resonance-assisted hydrogen bonding; secondary electrostatic interactions

1 | INTRODUCTION

"Water...shows tendencies both to add and give up hydrogen, which are nearly balanced. [...] a free pair of electrons on one water molecule might be able to exert sufficient force on a hydrogen held by a pair of electrons on another water molecule to bind the two molecules together. [...] Such an explanation amounts to saying that the hydrogen nucleus held between two octets constitutes a weak 'bond'." (Latimer WM and Rodebush WH, 1920)¹.

Hydrogen bonding interactions stand at the crossroad between weak noncovalent bonding and strong covalent bonding. They can be as weak as less than a kilocalorie per mole, they can be as strong as half the association of a single C—C bond (e.g., the $[F...H...F]^-$ interaction is about 40 kcal/mol), and the directionality of hydrogen bonds gives a clue to how molecules and molecular fragments might arrange in space.² From the varying strengths and the directionality of hydrogen bonds, emerges the opportunity for chemical

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CONFLICT OF INTEREST

The authors declare no potential conflict of interest.

design. Since the 1920 report of Latimer and Rodebush,¹ regularity in hydrogen bonding patterns were recognized, hydrogen bond design principles were developed, and it became possible to explain and imagine the structures and functions of many hydrogen bonding systems.

Leading up to the magnum opus of Latimer and Rodebush's proposal of hydrogen bonding, two events in the early 1900's steered the direction of hydrogen bond research during the first half of the 20th century: (a) G. N. Lewis' theory³ of valence and chemical structures (1916)—so that the idea of a hydrogen bond could be conceived, and (b) the discovery of X-ray (1895)—so that a hydrogen bond, that is, close contact between proton-sharing atoms, could be observed. Recognizing that a hydrogen nucleus shared between two atoms could largely influence the three dimensional structure of molecules and molecular fragments initiated a contentious race among multiple groups toward unveiling the double helix DNA structure (1953, Watson–Crick),⁴ the α -helix structure (1951, Pauling–Corey–Branson)⁵ and the β -sheet structure (1950, Pauling–Corey)⁶ of proteins, along with surveys and developments of hydrogen bonding patterns in organic crystals (1950's–1960's). These events happened at the height of early applications of crystallography in chemistry and set the stage for the discovery of many hydrogen bond design principles (1990's–2000's), which are now routinely applied to areas of recognition, catalysis, and assembly in organic and supramolecular chemistry.

This review presents an overview of hydrogen bond design principles, based on five structural features: electronegativity effects (Section 2), steric effects (Section 3), electrostatic effects (Section 4), π -conjugation effects (Section 5), and cooperativity effects (Section 6). Debates touched on but not discussed in detail include the physical origins of hydrogen bonds and the physical explanations of each design principle. Discussion focuses on how molecular structure affects hydrogen bond strength, and on showcasing selected applications of hydrogen bond design principles.

2 | ELECTRONEGATIVITY EFFECTS

2.1 | The classical hydrogen bond

In 1920, Latimer and Rodebush¹ (along with the unpublished works of Huggins a year earlier, Box 1) first related the idea of electronegativity and bond polarity to the sharing of a hydrogen atom between two atoms. They noted that ammonia readily adds a hydrogen, hydrogen chloride readily loses one, but water could add or lose a hydrogen, and therefore a hydrogen could be shared between two water molecules and bind two molecules together (Figure 2a). They recognized that ammonium hydroxide (Figure 2b) is another example in which the "*union is fairly strong*," explaining that "...*the hydrogen nucleus held between two octets constitutes a weak bond*." Huggins proposed the term "*hydrogen bridges*" to describe the sharing of an H atom between two molecules.⁷ These early depictions of hydrogen bonds were developed from Lewis' theory⁶ for valence and bonding and hinted at the covalent character (i.e., orbital interaction) of hydrogen bonds. A hydrogen bond X—H...Y may be viewed as a donor–acceptor orbital interaction, in which a set of lone pairs on Y donate into the antibonding orbital of X—H.⁸ The covalent character of hydrogen bonding also has been described by Coulson as "covalent-ionic resonance"⁹ and by several others, as

well as from a valence bond perspective, as "three-center-four-electron" interactions.^{10–14} An alternative view emerged when Pauling developed a scale of electronegativity,¹⁵ proposing an electrostatic explanation (i.e., dipole–dipole interaction) of hydrogen bonding instead, and commenting that, "*the hydrogen bond is largely ionic in character […] formed only between the most electronegative atoms.*"

Debates regarding the nature of hydrogen bonding continued for the remainder of the 20th century, but the important effects of electronegativity on hydrogen bond strengths were commonly recognized. An illustrative example comparing the O—H...N vs. N—H...N hydrogen bond is shown in Figure 3. According to the electronegativities of O (3.5) and N (3.0), O—H...N is considered a stronger hydrogen bond than N—H...N, and this trend can be understood from both the orbital interaction and the dipole–dipole interaction model of hydrogen bonding: (a) Since O is more electronegative than N, the antibonding orbital of O —H will have a larger coefficient on the electropositive H, thereby increasing donor-acceptor orbital interaction (Figure 3a). (b) Since O is more electronegative than N, an H atom attached to O will be more positively charged, thereby increasing the dipole–dipole interaction (Figure 3b).

Electronegativity differences are the simplest ways of explaining hydrogen bond strengths, and these effects were used to rationalize hydrogen bonding patterns in the early days of crystallography. Based on surveys of hydrogen bonding patterns in organic crystals, Donohue observed that all acidic hydrogens available in a molecule will be used in hydrogen bonding in the crystal of that compound.¹⁶ This idea was significantly expanded in Etter's works in the 1980's–1990's, where she applied graph sets to analyze organic crystals and proposed a set of rules, noting that: (a) All good proton donors and acceptors are used in hydrogen bonding, and that (b) the best hydrogen bond donor and the best hydrogen bond acceptor will preferentially form hydrogen bonds to one another.^{17,18}

A closely related hydrogen bond design principle, considering the acidity and basicity of proton donors and acceptors, is the idea of pK_a match.^{19–22} It was proposed that when a hydrogen bond is formed between an acid and its conjugate base, for example, HF...F⁻ (i.e., [F...H...F]⁻),²³ a matching pK_a value can give rise to short, strong, low-barrier hydrogen bonds, in which a proton can readily exchange between two atoms. The low-barrier hydrogen bond hypothesis was originally proposed to explain how enzymes might stabilize charged centers in catalytic reactions and remains a controversial topic.^{24–26}

2.2 | Nonclassical hydrogen bonds

In their authoritative work, "*The Hydrogen Bond* (W. H. Freeman, San Francisco, 1960),"²⁷ Pimentel and McClellan explained that X—H...Y can be considered to be a hydrogen bond if there is evidence of bond formation linking the two groups. Without restricting what atoms or groups X and Y had to be, this much broader definition of the hydrogen bond (progressive for its time!) opened imaginative possibilities to many types of "nonclassical" hydrogen bonds. The most common textbook depiction of a hydrogen bond (X—H...Y) is the attractive force of an H atom between two electronegative atoms (F, O, or N). A bonding interaction forms because a lone pair of the electron rich Y atom donates into the σ -antibonding orbital of X—H (Figure 3a), and also because of attractive electrostatic

interactions between the interacting H and Y atoms ($^{\delta-}X$ —H $^{\delta+}...Y^{\delta-}$) (Figure 3b). Yet, it is increasingly recognized that X and Y do not have to be electronegative atoms. Three types of nonclassical hydrogen bonds are discussed here: C—H...Y interactions, X—H... π interactions, and X—H...H M dihydrogen bonding.

2.2.1 C—**H...O** hydrogen bonding—In her 1962 Nature paper, "The C—H...O Hydrogen Bond in Crystals," Sutor first considered the possibility of attractive C—H...Y hydrogen bonding.²⁸ She found the carbon–oxygen contacts in many crystals to be closer than the combined van der Waals radii for O and H, and suggested that these interactions might be considered as C—H...O hydrogen bonds (Figure 4a).^{28,29} She commented that, "The C—H group may be activated by other atoms or groups of atoms promoting ionization or partial ionization of the hydrogen atom. Under these conditions, it resembles the O—H and N—H groups, etc., and it may form hydrogen bonds."²⁸ Her hypothesis was initially met with strong criticism from Donohue, a prominent crystallographer at the time.³⁰ But studies based on hundreds of neutron diffraction crystal structures 20 years later unveiled even more examples of attractive CH...O, CH...N, and CH...Cl interactions.³¹

Decades later following its initial discovery,^{28,29} the C—H...O hydrogen bond now finds many useful applications in structural chemistry and in supramolecular design.^{32–35} Lippard et al. demonstrated a remarkable F_2C —H...O hydrogen bonding motif³⁶ (Figure 4b), suggesting that C atoms with significant s character can form very strong C—H...O interactions, and proposing that F_2C —H groups can be useful for replacing OH groups in medicinal applications of hydrogen bonding. Anion binding based on C—H...anion interactions have gained increasing popularity,³⁷ and many receptors containing aryl C—H hydrogen bonding interactions have been developed (see Figure 4c for some examples).^{38–40}

2.2.2 | X—H... π hydrogen bonding— π -Bonds are electron rich and can replace F, O, or N atoms as hydrogen bond acceptors, giving rise to X—H... π hydrogen bonding interactions (Figure 5a). π -Hydrogen bonding interactions are weaker than classical hydrogen bonds, but they are prevalent in chemistry and biology, and it is increasingly recognized that these weak interactions are important for interpreting the interactions of aromatic rings, the conformations of organic compounds, chemical and biological recognition, crystallographic data, and the three dimensional structures of proteins.⁴¹

West first observed, based on infrared studies, that addition of olefins to phenol and other alcohols led to the appearance of a broad O—H band at low frequencies, suggesting that O —H groups could interact favorably with olefinic π -bonds (Figure 5b).⁴² Benzene forms hydrogen bonds with water through O—H… π interactions (Figure 5c).⁴³ Many drug– protein interactions feature N—H… π hydrogen bonding involving amine functional groups and aromatic rings.⁴¹ X—H… π interactions also can involve X atoms that are not especially electronegative. For example, C—H… π interactions are mostly the result of dispersion effects.^{44,45} Interestingly, Imamoto et al.⁴⁶ found that loss of a single CH… π interaction, between an alkyl group and π -ring of residues, could significantly alter the stability and photocycle of the photoactive yellow protein. Cremer et al. reported examples of B—H… π interactions in a carborane…benzene complex (Figure 5d), a diborane (B₂H₆)…benzene complex, and an Ir-dimercapto-carborane complex.⁴⁷

2.2.3 | X—H...HM dihydrogen bonding—Hydrogen bonding interactions can form between two hydrogen atoms if one is partial negatively charged and the other is partial positively charged ($^{\delta-}X-H^{\delta+}...^{\delta-}H-M^{\delta+}$) (Figure 6a).⁴⁸⁻⁵⁰ This happens when one H atom is attached to an electronegative center (X), and the other H atom is attached to an electropositive center (M), such as boron, silicon, or transition metals. In essence, the dihydrogen bond can be seen as an attractive proton-hydride interaction. Jackson et al. found that dihydrogen bonds could be used to preorganize colavent organic frameworks and to control the stereoselectivity outcome of organic reactions.^{51,52} Dihydrogen bonds can transform into hydrogen-hydrogen covalent bonds, driving off H₂ and leaving Lewis acidic and basic sites in close proximity, and ready to form strong covalent bonds (Figure 6b).⁵¹ Dihydrogen bonds also were shown to direct the diastereoselective outcomes of borohydride reduction reactions (Figure 6c).⁵² Based on computational and experimental NMR data, a favorable Si-H...H-O interaction was found between trihexylsilane and perfluoro-tertbutanol.⁵³ Besides electrostatically-driven dihydrogen bonding, homopolar, dispersiondriven, dihydrogen bonding interactions, for example, C-H...H-C interactions between dimers of alkanes and polyhedranes, 5^4 and X—H...H—X (X = B, Al, Ga) interactions, 5^5 also have been reported.

3 | STERIC EFFECTS

Steric effects were one of the earliest aspects to be considered in hydrogen bond design. Alder et al.'s 1968 work on the 1,8-bis(dimethylamino)naphthalene (DMAN, the original "proton sponge") first demonstrated that molecular strain can affect the Brønsted basicity of diamines.⁵⁶ In DMAN ($pK_a = 18.6$ in MeCN), two dimethylamino groups are attached to a naphthalene backbone, and two N lone pairs are pointed toward each other, giving rise to repulsive interactions (Figure 7a). Protonation of the diamine relieves lone pair repulsion, forming a low-barrier [N...H...N]⁺ hydrogen bond (Figure 7b).^{57,58} Along with other medium-ring diamine and polyamine structures, DMAN shows remarkable proton accepting ability compared with the typical aliphatic amine.

Following Alder's classic example, many examples of proton sponges with increased molecular strain and enhanced basicities were developed (Figure 8). Two strategies for developing proton sponges include: (a) Adding bulky, electrondonating, substituents to the amino groups (i.e., a "buttressing" effect). For example, guanidinyl-substituted proton sponges such as 1,8-bis(tetramethyl-guanidinyl)naphthalene (TMGN) ($pK_a = 25.1$ in MeCN)⁵⁹ and phosphazene-substituted proton sponges like P₂-TPPN ($pK_a = 42.1$ in MeCN)^{60,61} show increased basicity compared to DMAN. (b) Modifying the aromatic backbone to push the N lone pairs even closer to each other (i.e., a "crowding" effect). Some examples include *t*-Bu–P₂ ($pK_a = 33.4$ in MeCN),⁶² vinamidine ($pK_a = 31.9$ in MeCN),⁶³ and a fluorene-based sponge which readily deprotonates DMAN and displays near linear N…H…N hydrogen bonding.⁶⁴

4 | ELECTROSTATIC EFFECTS

4.1 | Secondary electrostatic interactions

The secondary electrostatic interaction (SEI) model of Jorgensen and Pranata^{65,66} has long been regarded as a textbook guideline for designing multipoint hydrogen bonding arrays, that is, hydrogen bonded complexes with more than one set of hydrogen bonds. They suggested that both primary electrostatic interactions (between the proton donor and acceptor) and secondary electrostatic interactions (between a proton donor or acceptor group and the hydrogen bonding group diagonal to it) could affect the association strengths of arrays.

According to the SEI model, arrays with all proton donors (D) on one fragment and all proton acceptors (A) on the other fragment (e.g., an "AA-DD" array) will exhibit stronger association strengths than arrays with alternating D and A arrangements (e.g., an "AD-DA" array), since the former arrangement maximizes the number of attractive electrostatic interactions. Note two attractive secondary interactions in the AA-DD array, but two repulsive secondary interactions in the AD–DA array (see Figure 9a). Based on the SEI model, the association strengths of triply hydrogen bonded arrays are expected to follow the order: AAA-DDD > AAD-DDA > ADA-DAD (Figure 9b), and those of quadruply hydrogen bonded arrays are expected to follow the order: AAAA-DDDD > ADDA-DAAD \approx ADAA–DADD > ADAD–DADA (Figure 9c). Especially robust multipoint hydrogen bonding arrays have been prepared following the SEI model (Figure 10).^{67,68} Statistical analyses supporting the SEI model have shown that the association strengths of hydrogen bonded arrays could be reproduced by summing up empirical increments^{69,70} or by combining calculated electrostatic forces⁷¹ that take into account the primary and secondary electrostatic interactions in complexes. It was suggested that electrostatic interactions between remote atom pairs in a hydrogen-bonded complex also could affect array association.72

Nevertheless, the SEI model remains a matter of debate and its limitations continue to invite controversy. Lukin and Leszynski argued based on extensive quantum chemical calculations that some ADD–DAA arrays appear to have weaker associations than their analogous AAA–DDD arrays only because of a more solvated ADD and DAA monomer in wet polar solvent. ⁷³ Fonseca Guerra et al. emphasized the importance of donor–acceptor orbital interactions^{74,75} and the effects of Pauli repulsion^{76,77} on the association strengths of arrays. Wu et al. found that arrays with the same SEI patterns can have varying association strengths depending on the aromatic characters of the interacting fragments.^{78–80} Rocha-Rinza et al. suggested that the SEI model might be refined by considering the acid–base properties of the hydrogen bonding groups.^{81,82} Fonseca Guerra et al. examined the SEI model to understand why it predicts binding strengths that are in line with experimental results, even though it oversimplifies the description of hydrogen bonded fragments is the result of both electrostatic interactions and σ -orbital interactions.⁸³

Despite a large body of experimental and theoretical work challenging the SEI model, it remains a useful principle for designing multipoint hydrogen bonding arrays. The SEI model

is chemically intuitive and can be easily applied based on simple "back-of-the-envelope" illustrations of donor and acceptor patterns in compounds.

5 | II-CONJUGATION EFFECTS

5.1 | Resonance-assisted hydrogen bonding

In the late 1980's, Gilli et al. introduced the idea of "resonance-assisted hydrogen bonding" (RAHB)—a simple hydrogen bond concept relating π -electron delocalization to hydrogen bonding in compounds.^{84–87} They noted that β -diketones that formed either: (a) intramolecular hydrogen bonds, or (b) a linear array of intermolecular hydrogen bonds, displayed enhanced π -electron delocalization (Figure 11a).⁸⁴ Resonance-assisted O—H... O=C hydrogen bonds were found to display short O...O distances, downfield shifted ¹H NMR signals, and red-shifted O—H stretching frequencies. Gilli et al. reasoned that partial charges generated by resonance on the O atom of the carbonyl group makes it a better proton acceptor, and as a result, the proton donor and acceptor groups move closer to each other, giving rise to stronger hydrogen bonding. In this way, hydrogen bonding increases π resonance, and π -resonance enhances hydrogen bond strength. As noted in the original paper, a reviewer of Gilli's initial paper suggested an alternative explanation, based on synergy between the σ - and π -framework. When a hydrogen bonding C=O group is π conjugated, π -resonance decreases the electronegativity of the O atom, and this raises the energy of the in-plane lone pair of O, making it a stronger hydrogen bond acceptor. Gilli's work concluded by speculating on the many imaginable implications of RAHB in chemical and biological systems, including hydrogen-bonded dimers, the secondary structures of proteins,⁸⁸ and DNA base pairs (Figure 11b).

In the 30 years following Gilli's proposal of the RAHB idea, opposing views either debating the importance of RAHB or the origin of the effect were put forth, based on a variety of theoretical approaches. Based on energy decomposition analyses of DNA base pairs, Fonseca Guerra et al. reported that even though π -polarization effects can enhance hydrogen bonding, electrostatic and donor-acceptor orbital interactions dominate the total interaction energy.^{74,75} Based on valence bond and atoms-in-molecules computations, Gora et al. attributed the effects of RAHB to charge delocalization—an idea captured in Gilli's explanation of the RAHB effect, describing that partial charges generated by resonance can enhance hydrogen bonding.⁸⁹ Evidence based on computed energy decomposition analyses, ⁹⁰ block-localized wavefunction analyses, ⁹¹ and coupling constants, ⁹² suggested that the effects of RAHB originated from geometric constraints of the σ -framework. In line with these ideas, later works from Fonseca Guerra et al.,⁹⁰ while finding little evidence for σ - vs. π - synergy in RAHB systems, confirmed that RAHB happens because π -resonance moves the donor and acceptor groups closer in proximity—an idea also captured in Gilli's account of the RAHB effect.

Most of the supposed arguments and works put forth to dispute or reexamine the RAHB idea, have reinforced rather than disproved Gilli's original explanation and novel discovery of the RAHB effect. This simple and powerful concept, that is, the connection between π -conjugation and hydrogen bonding,⁹³ has found significant use in synthetic transformations, in the design of chelating pockets for coordination, in molecular recognition, in the design of

molecular switches, and in crystal engineering, among many other applications in organic chemistry.⁹⁴

5.2 | Aromaticity and antiaromaticity

Dewar first recognized a relationship between hydrogen bonding and aromaticity, when he proposed the structure of stipitatic acid, calling it a nonbenzenoid aromatic. He suggested that stipitatic acid and many tropolone derivatives might be considered to be "aromatic," since intramolecular hydrogen bonding between C=O and OH groups at ortho positions could polarize the ring π -electrons to increase $[4n+2]\pi$ -aromaticity in the seven membered ring (Figure 12a).⁹⁵ More examples relating the effects of aromaticity gain and hydrogen bonding appeared later on. Aromaticity gain was found to affect the tautomeric equilibria of hydrogen bonding compounds,^{96,97} to increase the basicity of organic superbases,^{98,99} and to enhance the hydrogen bonding ability of heterocycles.^{100–103} It was suggested that hydrogen bonding of squaramide, at the two carbonyl and two amine groups, increased cyclic two π -electron aromaticity in the four membered ring (Figure 12b).^{102,103} In essence, the effect of aromaticity gain and loss on hydrogen bonding can be considered as a manifestation of the RAHB concept. While π -conjugated hydrogen bonding compounds can all benefit from "resonance-assistance," the energetic consequences of RAHB are especially pronounced when aromaticity gain happens. Interestingly, Gilli's original depiction of the supposed effects of RAHB in the guanine-cytosine and adenine-thymine base pairs, considered only the six membered ring of guanine and not the ring moieties of other nucleobases (see Figure 11b, red dotted lines) to have special importance for resonanceassistance (note resonance form showing aromaticity gain in guanine, see Figure 12c).

In 2014, a proof-of-concept paper by Wu et al. generalized the relationship between aromaticitiy gain and loss and hydrogen bonding, delineating its possible implications for hydrogen bond design.¹⁰⁴ They reported that hydrogen bonds are stronger than expected when they increase [4n + 2] cyclic π -electron delocalization (aromaticity gain) in the hydrogen bonding compounds, and are weaker than expected when they decrease [4n + 2]cyclic π -electron delocalization (aromaticity loss) in compounds (Figure 13a). Later works from Wu and Jackson et al. ^{105,106} extended the original idea to show that the opposite happens for [4n] "antiaromatic" rings, and these reciprocal relationships were later applied to rationalize the trends of hydrogen bonding in self-assembling systems,^{80,107–109} in multipoint arrays,^{78,79} and may rationalize short, strong hydrogen bonds in enzymes.¹¹⁰

Whether light irradiation strengthens or weakens a hydrogen bond also can be related to changes in (anti)aromaticity of hydrogen bonding compounds in the excited state.¹¹¹ Just as the rules for orbital interactions in organic reactions reverse in the excited state, the electron-counting rules for aromaticity and antiaromaticity also reverse at the first singlet and triplet $\pi\pi^*$ states. According to Baird's rule: $[4n] \pi$ -ring compounds are excited-state aromatic, and $[4n + 2] \pi$ -ring compounds are excited-state antiaromatic.^{112,113} By connecting Baird's rules to the effects of hydrogen bonding, Wu et al. deduced and demonstrated that,¹¹¹ upon photoexcitation, hydrogen bonding interactions that polarize ring π -electrons to increase excited-state antiaromaticity in compounds are weakened (Figure 13b). Conversely, hydrogen bonds that decrease excited-state antiaromaticity in compounds are strengthen

(Figure 13b), and in the extreme, relief of excited-state antiaromaticity can drive excitedstate proton transfer reactions.¹¹¹ Although not properly recognized in a large body of supporting examples,^{114–118} this relationship—between excited-state (anti) aromaticity and excited-state hydrogen bonds—explains why photoexcitation strengthens some hydrogen bonds but weakens others.

6 | COOPERATIVE EFFECTS

Networks of hydrogen bonds can give rise to strong hydrogen bonding interactions. Water clusters can form through networks of hydrogen bonds. Enzymes can engage multiple hydrogen bonding interactions to stabilize charges and facilitate catalysis (Figure 14a). 119,120

In an elegant experiment, Shan and Herschlag demonstrated that networks of intramolecular hydrogen bonds could significantly increase the acidity of benzoic acid in dimethylsulfoxide; the pK_a of benzoic acid decreases by 4 units, when one hydrogen bonding OH group is placed ortho to the carboxylic acid group, and by an enormous 8 units, when two OH groups are placed ortho to the carboxylic acid group (Figure 14b).¹²¹ Based on a series of covalent polyol models (Figure 14c), Wang, Kass, and coworkers showed that hydrogen bonding networks can stabilize charged centers, and that the compounding effects of having multiple hydrogen bonds may explain how charges in enzyme active sites affect catalysis and conformational changes.^{122,123} Rather than attributing the stabilization of charges in enzymes to the possible existence of a single short, strong "low-barrier" hydrogen bond, these authors proposed that multiple hydrogen bonds stabilize charged centers in enzymes. Neutral systems also can display strong, short-range cooperativity. Based on a series of synthetic molecular balances, Cockroft et al. demonstrated that neutral hydrogen bonds (see OH in bold, Figure 14d) could be strengthened by increasing numbers of cooperative, intramolecular hydrogen bonding interactions.¹²⁴ Networks of hydrogen bonds have been applied to the design of inhibitors, catalysts, and molecular receptors.^{125–128}

7 | CONCLUSION

Chemistry is largely a science of molecular design, and hydrogen bonding interactions are the workhorse for linking molecules and molecular fragments in a chemically intuitive way. Since the 1920 paper of Latimer and Rodebush, a hundred years has passed, and discussions surrounding "The Hydrogen Bond" has evolved from debates concerning the nature of the interaction, to milestones in crystallography, to explosive developments in the concept and application of hydrogen bond design principles (Figure 1). A potential area of growth is to rationalize the effects of external stimuli (e.g., light, pressure) on hydrogen bonding. We close our overview of the topic by emphasizing the value of simple hydrogen bonding principles, like the secondary electrostatic interaction model and the resonance-assisted hydrogen bonding concept, and their imperative roles in pushing the realms of molecular design in many areas of chemistry (e.g., in supramolecular catalysis, recognition, and assembly). Concepts like these are powerful, not because of theoretical rigor but because of conceptual simplicity—any chemist can pick up a pen and a piece of paper and sketch out the next ideas for an experiment. Whether or not the next experiments "work" is a separate

issue, what matters more is that these principles influence the evolution of molecular design in chemistry.

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BOX 1

HYDROGEN BONDING—A HARD-TO-SWALLOW IDEA

As contemporaries with Latimer and Rodebush in the lab of G. N. Lewis at Berkeley, Huggins first suggested a crude idea of hydrogen bonding in a class term paper in 1919. His idea was met with dismissal from Bray, who taught the course at the time and who commented that, "*Huggins, there are several interesting ideas in the paper, but there is one you'll never get chemists to believe: the idea that a hydrogen atom can be bonded to two other atoms at the same time.*" Even though Latimer and Rodebush described the idea of hydrogen bonding in their 1920 publication, the phrase "hydrogen bond" only appeared for the first time in Lewis' Valence and the Structure of Atoms and Molecules, in 1923, and the idea remained largely ignored until the mid-1930's.



FIGURE 1.

Timeline for the development of hydrogen bond design principles



FIGURE 2.

Depictions of hydrogen bonding, in (a) H_2OH_2O and (b) ammonium hydroxide, based on the early works of Latimer and Rodebush¹



FIGURE 3.

Effects of electronegativity on O—H...N vs. N—H...N hydrogen bond strength. Explanations based on: (a) donor–acceptor orbital interactions (using water...ammonia and ammonia...ammonia as examples), and (b) dipole–dipole interactions



FIGURE 4.

(a) Schematic illustration of CH...Y interactions (Y = electronegative atom or anion). (b) F_2C —H...O hydrogen bonding. (c) Examples of anion receptors based on C—H...anion interactions



FIGURE 5.

(a) Schematic illustration of XH... π interactions. (b) and (c) Examples of OH... π hydrogen bonding. (d) Example of B—H... π hydrogen bonding



FIGURE 6.

(a) Schematic illustration of dihydrogen bonding. Dihydrogen bonds were shown to direct the (b) preassembly of covalent materials, and (c) the diastereoselectivity of borohydride reduction



FIGURE 7.

(a) The original "proton sponge," DMAN. (b) Protonation of the diamine relieves lone pair repulsion, resulting in low-barrier [N...H...N]⁺ hydrogen bonding



FIGURE 8.

Examples of other diamine-based proton sponges: TMGN, P₂-TPPN, *t*-Bu–P₂, vinamidine, and a fluorene-based proton sponge



FIGURE 9.

Secondary electrostatic interactions (SEIs) between proton donors ("D," in orange) and acceptors ("A," in blue) in: (a) doubly, (b) triply, and (c) quadruply hydrogen bonded arrays. Solid lines indicate attractive interactions and dashed lines indicate repulsive interactions



FIGURE 10.

Examples of quadruply hydrogen bonded arrays prepared based on the secondary electrostatic interaction (SEI) model. (a) 2-Ureido-4-pyrimidone,⁶⁷ and (b) Blight's AAAA-DDDD array⁶⁸



FIGURE 11.

(a) Intramolecular and intermolecular resonance-assisted hydrogen bonding (RAHB) in β -diketone. (b) Possible RAHB effects in hydrogen-bonded dimers and base pairs



FIGURE 12.

Resonance structures showing the effects of aromaticity gain in (a) tropolone, and in hydrogen-bonded (b) squaramide and C) guanine



FIGURE 13.

(a) Aromaticity gain and loss in hydrogen-bonded heterocycles, and (b) a reversed effect in photoexcited states



FIGURE 14.

(a) Hydrogen bonding networks in the active site of triosephosphate isomerase. (b) Cooperative hydrogen bonds can stabilize the conjugate base of OH substituted benzoic acid (note pK_a values for the acid). (c) Example of a covalent polyol system. (d) Enhanced hydrogen bonding resulting from neutral, short-range networks (see OH group in bold)