Picking Out Logic Operations in a Naphthalene β-Diketone Derivative by Using Molecular Encapsulation, Controlled Protonation, and DNA Binding

Sameena Yousuf,[a] Ritty Alex,[a] Paulraj Mosae Selvakumar,*[a] Israel V. M. V. Enoch,*[a] Palani Sivagnana Subramanian,[b] and Yu Sun[c]

On–off switching and molecular logic in fluorescent molecules are associated with what chemical inputs can do to the structure and dynamics of these molecules. Herein, we report the structure of a naphthalene derivative, the fashion of its binding to β-cyclodextrin and DNA, and the operation of logic possible using protons, cyclodextrin, and DNA as chemical inputs. The compound crystallizes out in a keto-amine form, with intramolecular N–H–O bonding. It shows stepwise formation of 1:1 and 1:2 inclusion complexes with β-cyclodextrin. The amino-pentenone substituents are encapsulated by β-cyclodextrin, leaving out the naphthalene rings free. The binding constant of the β-cyclodextrin complex is 512 M\(^{-1}\). The \(pK_a\) value of the guest molecule is not greatly affected by the complexation. Dual input logic operations, based on various chemical inputs, lead to the possibility of several molecular logic gates, namely NOR, XOR, NAND, and Buffer. Such chemical inputs on the naphthalene derivative are examples of how variable signal outputs based on binding can be derived, which, in turn, are dependent on the size and shape of the molecule.

Introduction

Since the presentation of the first information-processing molecules,[1] examination of chemical compounds in light of their Boolean logic functions and their possible applications has shown remarkable progress.[2,3] It is likely that such logic devices will find applications in medicine,[4] biotechnology,[5] and physiology.[6] These logic devices have shown substantial and intriguing results and have been envisaged to have some notable applications: 1) A reliable detection of pathophysiological conditions is possible with molecular logic systems,[7] 2) secure, economic, and simple methods can be developed for designing complex DNA-based logic devices,[8] 3) the molecular Boolean logic language can function as a medical diagnostic protocol.[9] 4) encapsulated drugs can be trigger-released through logic response.[10] 5) dually controlled photodynamic therapy is practicable using the AND logic gate as the template; 6) the DNA-binding of drugs can be controlled using external or internal stimuli based on a molecular switching principle,[11] and 7) molecular logic gates can be used in fluorescent papers and applied to cryptography.[12]

In one example, an AND logic gate was used as the template in photodynamic therapy involving a boron-dipyrromethene (BODIPY) derivative, and H\(^{+}\) and Na\(^{+}\) served as the inputs.[13] H\(^{+}\) is a sensory and internal input, and could be important because the pH of cancer cells is lower than for healthy cells.[14] By measuring the drop in pH value, a biochemical filtering process can be realized in a signal transduction logic system.[15] The absorption and fluorescence characteristics of luminescent molecules in solution are altered by the change of pH.[16,17] and encapsulation of acidic small molecules by cyclodextrin can alter their \(pK_a\) values.[18] Cyclodextrins (β-CD) are tapered-cone-like cyclic oligosaccharides with a hydrophobic cavity capable of accommodating appropriately sized organic guest molecules.[20] The guest molecule’s fluorescence is generally switched on by the host–guest complex formation with cyclodextrin. Hence, studies on chemical sensing, molecular logic design, and small object recognition could take advantage of cyclodextrin as a host structure.[21] Molecules having multiple light-emissive electronic excited states are pertinent switching structures,[22] and naphthalene derivatives are examples.[23] Their acidities are quite varied, and in their cyclodextrin inclusion complexes, they can show varying acidity responses depending on the part of the derivative molecule en-
gulbed by the cyclodextrin cavity.\cite{24} Hence, it is of paramount importance to study the detailed mode of binding of guest molecules to cyclodextrin.

Besides being sensitive to protonation and cyclodextrin complexation, the fluorescence signals of β-diketone naphthalene derivatives can be switched on and off by the addition of DNA, as these compounds are switchable DNA intercalators.\cite{25} Either DNA itself\cite{26} or the DNA-binding of small molecules could function as biologically relevant logic gates, with possible applications in anticancer drug action.\cite{27} Despite the importance of cyclodextrin and DNA as molecules having size- and shape-specificity in binding events and the on/off switch response of fluorescent molecules upon binding, the combined inputs of these molecules has not yet been reported. Herein, we report the synthesis, mode of binding to cyclodextrin and DNA, and molecular logic operations of (Z)-5-(S-((Z)-4-oxopent-2-en-2-ylamino)naphthalen-1-ylamino)pent-3-en-2-one (abbreviated as H$_2$acacnn).

### Results and Discussion

#### Crystal and molecular structure of H$_2$acacnn

H$_2$acacnn crystallizes in the monoclinic space group P2$_1$/n, and the crystallographic data, selected H-bond lengths, and bond angles are given in Table 1. The Oak Ridge Thermal Ellipsoid Plot (ORTEP) drawing of the molecule is shown in Figure 1.

![Figure 1. ORTEP diagram of the ligand with 50 % probability for the thermal ellipsoid.](image-url)

The single crystal X-ray structure of H$_2$acacnn confirms that the keto amine form is available in the solid state. The aliphatic part of the compound shows an intramolecular N–H–O interaction within the unit (Figure 1). The hydrogen atom H$_1$ from the amine nitrogen N$^1$ is involved in the intramolecular N–H...O interaction with carbonyl group which is in the same plane. In order to make an effective N–H–O interaction, the terminal aliphatic moieties are tilted 43.3° with the NH-substituted naphthyl ring within the hydrogen-bonding distance as depicted in Figure 1. The amino hydrogen NH$_2$ acts as a donor and is involved in the N–H–O bonding, with the O$^1$ of the keto group acting as an acceptor. The hydrogen-bonding distances are as follows: N$^1$–H$^1$–O$^1$: N$^1$–O$^1$ = 2.634 Å, H$^1$–O$^1$ = 1.843 Å, and N$^1$–CH$_2$–O$^1$ = 144.3°. The crystal structure confirms that the $Z$-configuration of the olefinic bond is stabilized by intramolecular N–H–O interactions between the amine and keto groups. The possible tautomers, ionic and hydrogen-bonding interactions, and $E$, $Z$ isomers are shown in Scheme 1.

### Table 1. Crystal data and structure refinement for H$_2$acacnn.

| Crystal Data |  |
|-------------|---|
| CCDC Number$^{[a]}$ | CCDC 927799 |
| Empirical formula | C$_{20}$H$_{20}$N$_2$O$_2$ |
| Formula weight | 322.40 |
| Crystal color and habit | colorless prism |
| Crystal size (mm) | 0.24 x 0.12 x 0.07 |
| Temperature (K) | 150(2) |
| Wavelength (Å) | 1.54184 |
| Crystal system | monoclinic |
| Space group | P2$_1$/n |
| Unit cell dimensions | a = 8.0857(1) Å, α = 90° |
| | b = 9.1244(1) Å, β = 108.278(2)° |
| | c = 12.0067(2) Å, γ = 90° |
| Volume (Å$^3$) | 841.13(2) |
| Z | 2 |
| Calculated density (g cm$^{-3}$) | 1.273 |
| Absorption coefficient (mm$^{-1}$) | 0.0399, wR$_{20}$ = 0.1033 |
| Max. and min. transmission | 1.00000 and 0.91065 |
| Refinement method | Full-matrix least-squares on F$^2$ |
| Data/restraints/parameters | 1336/1/115 |
| Goodness-of-fit on F$^2$ | 1.144 |
| Final R indices | R$_1$ = 0.0341, wR$_2$ = 0.1004 |
| R indices (all data) | R$_1$ = 0.0399, wR$_2$ = 0.1033 |
| Extinction coefficient | 0.011(2) |
| Largest diff. Peak and hole (e Å$^{-3}$) | 0.193/–0.190 |

$^{[a]}$ CCDC 927799 contains the supplementary crystallographic data for the complexes in this study. These data can be obtained free of charge from the Cambridge Crystallographic Data Center via www.ccdc.cam.ac.uk/data_request/cif

### Formation of the β-CD–H$_2$acacnn host–guest complex

The absorption spectra of H$_2$acacnn (c = 1.0 μM) at pH 7 (Figure 2a) is characterized by a structureless band ($\lambda_{max}$ = 326 nm). The presence of β-CD in increasing concentrations leads to a continuous blue shift (hypochromic shift) of the band envelope centered at 326 nm to 324 nm (in 4.0 mM β-CD). This band is due to the n→π∗ transition. Also, the increase in β-CD leads to a regular hyperchromic shift. When the addition of β-CD is continued, above 5 mM of β-CD, an abrupt increase in the absorbance (hyperchromic shift) could be observed. Above this concentration, there is another 2 nm blue shift of the band and a continuous but small hyperchromic shift.
in the preceding lower concentrations [β-CD. Another important observation is that at higher concentrations of β-CD, the absorption spectra of H2acacnn show a vivid isosbestic point at 352 nm which is due to the equilibrium between the free and the β-CD-bound H2acacnn molecules in solution.

Aside from the absorption band at 326 nm, H2acacnn shows another band at 211 nm, corresponding to the π→π* transition. This band is structureless, probably due to the substitutions on the aromatic naphthalene ring. This band shows a bathochromic shift when β-CD is added in aliquots. At 5 mM β-CD, this shifted band is centered at 218 nm, while at 10 mM β-CD, it appears at 221 nm. Moreover, there is a continuous hyperchromic shift with increasing β-CD concentrations. The overall 10 nm bathochromic shift and the regular hyperchromic shift are attributed to the complex formation of H2acacnn with β-CD. The red shift and the increase in intensity of this band can be related to the conformational stability of the substituent aminopenitene chains of H2acacnn, attained due to the restriction offered by the encapsulating β-CD molecule. Presumably, the addition of β-CD results in the decrease in energy level of the excited state accompanying dipole-dipole interactions. Also, in the band corresponding to the π→π* transition, the addition of β-CD shows two distinct sets of absorption trends, namely a regular hyperchromic shift up to 5 mM β-CD and an abrupt increase of absorbance followed by a regular hyperchromic shift above this concentration. Evidently, β-CD forms an inclusion complex with H2acacnn, and there may be two different types of complexes formed. In order to evaluate the stoichiometry of the complex(es) and their binding strength(s), plots were made for the following binding events, assuming 1:1 stoichiometry:[29]

\[
CD + H_2acacnn \leftrightarrow CD : H_2acacnn \quad (1)
\]

\[
K = \frac{[CD : H_2acacnn]}{[CD][H_2acacnn]} \quad (2)
\]

where \([CD]_t = [CD] + [CD : H_2acacnn],\)

\[
[CD]_{t1} = [CD] + K[CD][H_2acacnn], \quad (3)
\]

\[
[CD]_{t1} = [CD](1 + K[H_2acacnn]), \quad (4)
\]

\[
[H_2acacnn]_{t1} = [H_2acacnn] + [CD : H_2acacnn] \quad (6)
\]

Substituting Equation 2 in Equation 6, we get

\[
[H_2acacnn]_{t1} = [H_2acacnn] + K[CD][H_2acacnn] \quad (7)
\]

Substituting Equation 5 in Equation 7, we get

\[
[H_2acacnn]_{t2} = [H_2acacnn] + K[H_2acacnn][CD] \frac{1}{1 + K[H_2acacnn]} \quad (8)
\]

where CD, H2acacnn, CD: H2acacnn, H2acacnn, and K are the free CD, the free H2acacnn, the host-guest complex, the total H2acacnn concentration, and the binding constant, respective-

The blue shift in absorbance is due to the change in the microenvironment of H2acacnn in aqueous solution upon addition of β-CD. The host-guest association between β-CD and H2acacnn leads to the shift since the hydrophobic cavity of β-CD offers an environment different from polar water molecules. The overall small shift of the absorption maximum suggests that there is no strong bonding between the H2acacnn molecule and β-CD. The hyperchromic shift observed upon addition of β-CD is due to the free movement of electrons into various energy levels and the resulting enhanced absorption of UV light. This is a consequence of the molecular interaction between the β-CD and the H2acacnn molecules, which induces structural modifications in the system. The electronic transition probability of the H2acacnn electrons increases in the presence of β-CD; hence, the magnitude of absorbance and the molar absorption coefficient increase. The sheer change of absorbance above 5 mM β-CD implies that the electronic charge distribution and the associated transition moment of H2acacnn are different on β-CD binding, compared with those observed
Equation 8 predicts a linear correlation between $H_{2}acacnn$ and $CD$ which is actually obtained within the concentration range of 0 to 5.0 mM $\beta$-CD (this is the highest concentration used in the plot). The linearity obtained in this plot (Figure 2b) suggests that at these lower concentration limits, there is formation of a 1:1 complex of $\beta$-CD:$H_{2}acacnn$. However, when the concentration higher than this range is included in the plot (the maximum concentration of $\beta$-CD is 10 mM), there is a deviation from linearity. Hence, a nonlinear plot is made (Figure 2c), for the higher order complex formed, considering two cases, namely 1:1 and 1:2 mixed complexes coexist [Eq. (9)] and only the 1:2 complex $(H_{2}acacnn;\beta$-CD) is formed [Eq. (10)].

\[
A = \frac{A_{0}A_{1}K_{1}[CD] + A_{2}K_{1}K_{2}[CD]_{2}}{1 + K_{1}[CD] + K_{1}K_{2}[CD]_{2}}
\]  

\[
A = \frac{A_{0} + A_{2}K_{2}[CD]_{2}}{1 + K_{1}K_{2}[CD]_{2}}
\]

where $A_{0}$, $A_{1}$, and $A_{2}$ are the absorbances of $H_{2}acacnn$ in water, 1:1 $H_{2}acacnn;\beta$-CD complex, and 1:2 complex respectively. $K_{1}$ and $K_{2}$ indicate the binding constants of the 1:1 and 1:2 complexes. [CD]$_{0}$, the initial concentration, can be used to replace [CD], the equilibrium concentration, since it is much smaller than [CD]. Reasonable values of standard errors, correlation coefficient, and confidence intervals could be obtained only when a nonlinear regression analysis using Equation 10 was done.

The binding constants determined for the 1:1 and the 1:2 $H_{2}acacnn;\beta$-CD complexes are $606 \text{ M}^{-1}$ and $2.32 \times 10^{3} \text{ M}^{-2}$ respectively. The fitting procedure was done using the above equations in the region of maximum variation of absorbance changes (325 nm). These binding constant values are independent estimates of the ground state binding affinities of $H_{2}acacnn$ with $\beta$-CD. The stepwise binding constants cannot be determined for the 1:2 complex from curve fitting done with just a single equation.

The fluorescence emission of $H_{2}acacnn$ in water (at pH 7) is a structureless band with a maximum at 407 nm (Figure 2d). When $\beta$-CD is added in stepwise increasing concentrations, $H_{2}acacnn$ is transferred from the aqueous bulk to the hydrophobic $\beta$-CD cavity, and the relative intensity of fluorescence reflects a change in the microenvironment of the guest mole-
cule. The fluorescence behavior of \( H_2\text{acacnn} \) is distinctly different between the two concentration ranges, namely 0 to 5 and 5 to 10 mM \( \beta\text{-CD} \). The usual \( \beta\text{-CD} \) complex formation results in an enhancement of the guest fluorescence intensity owing to the steric hindrance, offered by the \( \beta\text{-CD} \) cavity, to the molecular degrees of freedom leading to a nonradiative deactivation of the singlet excited state. We observed an immediate enhancement of fluorescence of \( H_2\text{acacnn} \) at the first addition of \( \beta\text{-CD} \) (5.0 mM), followed by a quenching of fluorescence of the originally enhanced band continuously up to 4.0 mM of \( \beta\text{-CD} \). This cannot be fitted by a quenching plot as the fluorescence intensity of \( H_2\text{acacnn} \) without \( \beta\text{-CD} \) lies below the fluorescence intensities of \( \beta\text{-CD} \)-bound \( H_2\text{acacnn} \) in the above-mentioned concentration range. Although the abrupt enhancement of fluorescence at the first addition can be easily understood, the subsequent quenching of fluorescence is unusual. The size match between the host cavity and the guest and the stoichiometry of the complex, to a specific microenvironment inside the \( \beta\text{-CD} \) cavity, can control such an effect. Generally, lower- or higher-order complexes govern respectively the less or more tightly binding behavior of the guest molecule with the \( \beta\text{-CD} \) environment. Moreover, in the higher order complexes, the guest is more protected from contact with water. The quenching of fluorescence may occur due to the low concentration of the host cavity by one side chain of the \( H_2\text{acacnn} \) molecule, leaving out the central naphthyl ring outside the host. Now, the aminopentenone chain, having sufficient conformational flexibility and a less restricted degree of freedom, cannot get induced in much variation of its emission. It is usual for a linear carbon chain to rattle around inside the \( \beta\text{-CD} \) cavity.[31] In general, the naphthalene ring can be included deeply into the \( \beta\text{-CD} \) cavity only with the long axis parallel to the CD axis.[31] In \( H_2\text{acacnn} \), since the naphthyl ring is not parallel to the aminopentenone chain, it cannot be accommodated by \( \beta\text{-CD} \).

With 5 to 10 mM of added \( \beta\text{-CD} \), the \( H_2\text{acacnn} \) displays an enhancement of fluorescence (Figure 2e). Again, this enhancement of fluorescence is ascribed to the accommodation of the second aminopentenone chain into the next \( \beta\text{-CD} \) molecule. The fluorescence maximum is not significantly shifted. Since only the side chains of \( H_2\text{acacnn} \) are entering the \( \beta\text{-CD} \) cavity, water molecules can freely move through the cavity along the sides of the encapsulated part of \( H_2\text{acacnn} \), resulting in the effective dielectric constant of the \( \beta\text{-CD} \) cavity to remain similar to that of water. This explains the insignificant shift of fluorescence maximum. Similar to the plot made using Equation 10 which utilizes absorbance, usually, we should be able to use the fluorescence intensity to do a nonlinear plot. However, in the case of \( H_2\text{acacnn} \), the plot could not be made because initially, there is fluorescence quenching at the low concentration range of \( \beta\text{-CD} \). Hence, mainly based on the results of UV/Vis absorption spectral data, it is inferred that \( H_2\text{acacnn} \) forms a higher order (1:2) complex with \( \beta\text{-CD} \) at higher concentrations of \( \beta\text{-CD} \), with the \( \beta\text{-CD} \) molecules encapsulating the aminopentenone chains.

NMR spectral analysis

In order to obtain direct evidence for the formation of the host–guest complex and to provide additional and concrete evidence for the mode of inclusion discussed in the previous section, \(^1\text{H} \) NMR spectra were recorded for the free \( H_2\text{acacnn} \), the free \( \beta\text{-CD} \), and the \( \beta\text{-CD}–H_2\text{acacnn} \) complex. The significant differences in the chemical shift values between these spectra confirm the formation of the inclusion complex. Larger and pronounced chemical shifts are observed for the \( H_1^2 \) and \( H_4^2 \) protons of \( \beta\text{-CD} \), compared to \( H_4^1 \) and \( H_4^4 \) protons. The \( H_4^1 \) and \( H_4^4 \) protons are located inside the hydrophobic cavity of \( \beta\text{-CD} \). The first inference is that the complex is formed between \( \beta\text{-CD} \) and \( H_2\text{acacnn} \) through the host–guest association. The \( H_2^5 \) and \( H_4^5 \) protons are on the outer surface of \( \beta\text{-CD} \), hence their signals are less affected by the guest molecule. The chemical shifts of the protons of the free \( H_2\text{acacnn} \) and the \( \beta\text{-CD}–H_2\text{acacnn} \) host–guest complex are listed in SI 1 in the Supporting Information.

The geometry of the inclusion complex and the structural information was provided by rotating-frame Overhauser effect spectroscopy (ROESY). The distance between the nuclei that are interacting with each other is directly proportional to the intensities of the cross peaks in the ROESY spectrum. When the internuclear distance between protons decreases, the cross peak intensity decreases. The ROESY spectrum of the \( \beta\text{-CD}–H_2\text{acacnn} \) complex displays cross peaks between the protons lined inside the cavity of \( \beta\text{-CD} \), namely \( H_1^2 \) and \( H_4^2 \), and the protons of the aminopentenone substituent of \( H_2\text{acacnn} \) shown as \( H_2^{15} \) and \( H_2^{26} \) in Figure 3a. There are two observations from these results: 1) the two aminopentenone chains are encapsulated by \( \beta\text{-CD} \), forming a 1:2 complex, and 2) the cross peaks are intense and, hence, the alkenyl protons are well placed inside the \( \beta\text{-CD} \) cavity, close to its \( H_1^2 \) and \( H_4^2 \) protons. The other weak cross peaks between the secondary hydroxy proton signals of \( \beta\text{-CD} \) and the aromatic protons (\( H_1^2 \) and \( H_4^2 \)) signals of \( H_2\text{acacnn} \) imply that the two \( \beta\text{-CD} \) molecules slide through the aminopentenone chains and get closer to the aromatic naphthyl ring although not encapsulating it. Very weak cross peaks are observed for the interaction between the end methyl protons of \( H_2\text{acacnn} \) (\( H_1^{17} \) and \( H_1^{29} \)) and the cavity protons of \( \beta\text{-CD} \), suggesting that the methyl end group penetrates deeply and comes outside the \( \beta\text{-CD} \) cavity through the other rim, as shown in the schematic diagram of the 1:2 inclusion complex (Figure 3b).

We performed diffusion-ordered NMR spectroscopy (DOSY) for \( H_2\text{acacnn} \) and the \( \beta\text{-CD}–H_2\text{acacnn} \) complex (SI 2 in the Supporting Information) in order to demonstrate that the complex is stable and to calculate the binding constant. The diffusion coefficient of free \( H_2\text{acacnn} \) is \( 4.515 \times 10^{-10} \text{ m}^2\text{s}^{-1} \) and that of \( \beta\text{-CD} \) is \( 3.701 \times 10^{-10} \text{ m}^2\text{s}^{-1} \). In the DOSY spectrum of \( \beta\text{-CD}–H_2\text{acacnn} \) complex, the signals due to the unbound \( H_2\text{acacnn} \) are not found suggesting that it is fully complexed to \( \beta\text{-CD} \). The diffusion coefficient of \( \beta\text{-CD} \) is \( 3.254 \times 10^{-10} \text{ m}^2\text{s}^{-1} \). The application of DOSY data to the binding of \( H_2\text{acacnn} \) to \( \beta\text{-CD} \) gives a binding constant value of \( 512 \text{ m}^{-1} \), which indicates strong binding and good stability of the complex. It should be
noted that this is close to the calculated value using the steady-state fluorescence spectral data.

Effect of pH on the absorption and fluorescence characteristics of H$_2$acacnn

The absorption and the fluorescence spectra of H$_2$acacnn are studied in the pH range of 1–7. With the decrease of pH from 7, the absorption spectrum of H$_2$acacnn (\(\lambda_{\text{max}} = 326\) nm) is considerably blue shifted (SI 3a in the Supporting Information). Below pH 4.5, the 326 nm band disappears completely. Parallel to this observation, there is a corresponding increase of absorbance at 275 nm, with the formation of a new band at around 291–292 nm. A large blue shift of absorbance is characteristic of the formation of a cation due to the H$^+$ ions protonating the nitrogen attached to the aromatic ring of the naphthalene through its unshared pair of electrons. This leads to a shortening of the conjugation of electrons in the chromophore and, hence, to a blue shift of the absorbance band. The presence of an isosbestic point in the spectra indicates that there is an equilibrium existing between two forms of the H$_2$acacnn molecule, namely the neutral and the cationic forms.

There is also another isosbestic point observed at 220–221 nm. The absorption spectrum of the protonated H$_2$acacnn resembles the spectrum of the unsubstituted naphthalene. This result suggests that the amino nitrogens attached to the naphthalene are protonated at a low pH, decreasing the conjugation of electrons and the length of the chromophore. The ground state p$K_a$ of this equilibrium in solution is calculated as follows:

\[
P_{K_a} = \text{pH} + \log \frac{C_1}{C_2}
\]

In the above equations, \(C_T\) represents the total concentration of H$_2$acacnn in neutral and cationic forms and \(e_{\lambda_1} (\lambda_1), e_{\lambda_2} (\lambda_2), e_{\lambda_3} (\lambda_3), e_{\lambda_4} (\lambda_4)\) are the molar extinction coefficients of the protonated and neutral forms at the two different wavelengths \(\lambda_1\) and \(\lambda_2\) respectively (275 nm and 325 nm in this case). The calculated p$K_a$ for the neutral-cation equilibrium of H$_2$acacnn in water, calculated using the above equations, is 3.7 ± 0.2.

The effect of acid strength on the absorption spectrum of H$_2$acacnn is shown in SI 3b in the Supporting Information. The longer wavelength absorption band (325 nm) decreases in absorbance with a corresponding increase at a shorter wavelength (230 nm). There is an isosbestic point observed in the range of 268 to 272 nm, which is not as clear as obtained for the prototropic equilibrium of H$_2$acacnn in water. This may be due to the presence of \(\beta\)-CD. The ground state p$K_a$ value of H$_2$acacnn in the presence of \(\beta\)-CD was determined in the spectrophotometric method as 3.5 ± 0.2. This value is slightly lower than the p$K_a$ obtained for the same equilibrium in water (3.7 ± 0.2), which is due to the hindrance offered by the \(\beta\)-CD molecule as it covers up the aminopentenone chains of H$_2$acacnn.

The fluorescence spectra of H$_2$acacnn in water, at various pH levels, are shown in SI 3c in the Supporting Information. The fluorescence spectra were recorded with excitation at the isosbestic points. When the pH is reduced from 7, the fluorescence emission band at 411 nm starts to quench with a red shift.
shift of fluorescence. The decrease of intensity at this wavelength is accompanied by a corresponding increase in intensity at a longer wavelength. The band at 411 nm completely disappears in the pH range below 5. The new band formed is centered at 454 nm. This band is formed due to the protonation of the H$_2$acacnn molecule. There is an isoemissive point around 420 nm, indicating that there exists a prototropic equilibrium between the neutral and the protonated forms of H$_2$acacnn. The fluorometric titration curves plotted as $I/I_0$ versus pH at the wavelengths 411 nm and 454 nm are sigmoidal, and these intersect at the middle of inflection. This is due to the occurrence of a single phenomenon, that is, the protonation of H$_2$acacnn. The pK$_a^*$ value (the excited-state pK$_a$) of the neutralization equilibrium of H$_2$acacnn could be directly obtained from the intersection of the fluorimetric titration curves, and the observed value is 3.4 ± 0.2.

A similar pH-based fluorimetric titration was done with H$_2$acacnn in the presence of β-CD (SI 3d in the Supporting Information). A red shift of the fluorescence band of the neutral form is observed on decreasing the pH below 7, which is similar to the observation of the fluorescence shift of H$_2$acacnn in water. However, the newly formed red-shifted band, which corresponds to the protonated form of H$_2$acacnn, is more intense than the fluorescence band of the neutral form. There is no single isoemissive point albeit most of the spectra, in the studied pH range, passes through a meeting point at around 395 nm. The absence of a clear isoemissive point may be due to two reasons: 1) the proton that tends to add to the nitrogen lone pair of the substituent on H$_2$acacnn might be in equilibrium between its different tautomeric forms and 2) the β-CD covers one of the aminopen-tenones at this concentration (4.0 mm); and so there may be two nonuniform sites of protonation, in the sense that the nitrogen of one of the aminopen-tenones is open and the other is β-CD-encapsulated. The excited-state pK$_a^*$ calculated for H$_2$acacnn in the presence of β-CD was calculated from the fluorimetric titration curves in the plot of $I/I_0$ versus pH as 3.2 ± 0.2.

**Binding of H$_2$acacnn to DNA**

The interaction of H$_2$acacnn with calf thymus DNA was studied by using absorption and fluorescence spectroscopy. The binding titration was carried out by keeping the concentration of H$_2$acacnn constant at 1.0 μM and adding aliquots of DNA. The absorption spectra of H$_2$acacnn with the various added amounts of DNA are shown in Figure 4a. The range of concentration of the added DNA is 0–1.4 μM. With an increase in the concentration of DNA, the absorbance of the 322 nm band decreases along with a hypsochromic shift of about 2 nm. The lesser availability of the H$_2$acacnn molecule to the surrounding solvent molecules, due to binding to DNA, results in the decrease in absorbance. The hypsochromic shift may be due to the intercalation of the naphthalene moiety of H$_2$acacnn into the DNA helix. A decrease in absorbance is an indication of the close proximity of the small molecule (H$_2$acacnn) to the DNA bases. Moreover, hypsochromism is generally a result of the contraction of the helical axis of the DNA and a change of conformation of DNA. On the other hand, damage to the DNA double helix would result in a hyperchromic shift of absorbance. We do not observe such a hyperchromism; hence, it can be inferred that there is no damage to the DNA upon H$_2$acacnn binding. Further evidence for intercalative binding is discussed in the section on molecular docking. A double recip-
rocal plot was used to determine the binding constant of the H$_2$acacnn–DNA binding using the following equation$^{[36]}$

$$
\frac{A_b}{A - A_b} = \frac{E_{0}}{E_{1} - E_{0}} + \frac{E_{1} - E_{0}}{(E_{1} - E_{0})K_b[\text{DNA}]}
$$

(14)

where $A_b$ and $A$ are the absorbances of the free H$_2$acacnn in water and at various concentrations of the added DNA and $E_{0,1}$ are the corresponding absorption coefficients. The linear plot of $A_b/A - A_b$ versus 1/[DNA] (correlation coefficient, 0.99) is shown in SI 4a in the Supporting Information. The calculated binding constant $K_b$ is 2.09 × 10$^5$ M$^{-1}$.

The H$_2$acacnn–DNA binding was also studied using fluorescence spectroscopy. The fluorescence titration of H$_2$acacnn with DNA shows a quenching of fluorescence (Figure 4b), along with a significant blue shift of the band at 407 nm. The Stern–Volmer quenching constant ($K_{SV}$) is calculated from the plot (SI 4b in the Supporting Information) made using the fluorescence titration data following the equation$^{[37]}$

$$
\frac{F_0}{F} = 1 + K_{SV}[Q] = k_q\tau_0[Q]
$$

(15)

where $F_0$ and $F$ are the intensities of fluorescence of H$_2$acacnn in the absence and the presence of DNA, $k_q$ is the bimolecular quenching constant, and $\tau_0$ is the fluorescence lifetime of H$_2$acacnn, and [Q] is the concentration of the quencher (DNA). The $K_{SV}$ which is proportional to the quenching efficiency of DNA, is 3.75 × 10$^4$ M$^{-1}$.

The binding data were cast into a plot of $\log(F_0/F - F)$ versus log [Q], where $F_0$ is the fluorescence intensity of H$_2$acacnn with no added quencher and $F$ is the fluorescence intensity at each concentration of the quencher. They were straight-line-fitted (SI 4c in the Supporting Information) to the model$^{[38]}$

$$
\log\left(\frac{F_0}{F} - F\right) = \log K_b + n \log[Q]
$$

(16)

where $n$ is the number of binding sites, [Q] is the concentration of the quencher (DNA), and $K_b$ is the binding constant.

The influence of $\beta$-CD encapsulation on the binding of H$_2$acacnn with DNA was studied using the titration of H$_2$acacnn–$\beta$-CD complex against DNA. H$_2$acacnn–$\beta$-CD exhibits absorption bands with $\lambda_{\text{max}}$ at 209, 228, and 321 nm (Figure 4c). Stepwise addition of DNA up to 1.4 μM results in the hypsochromic shift of absorbance along with a small blue shift. In the presence of $\beta$-CD, H$_2$acacnn is less available for energy transfer to the surrounding solvent molecules than in pure water. This leads to a decrease in the absorbance of H$_2$acacnn. The naphthalene moiety of H$_2$acacnn may have intercalated into the DNA helix as the competitive binding between cyclodextrin–H$_2$acacnn and DNA–H$_2$acacnn takes place, where the fluorophore sheds the two cyclodextrin units in favor of the DNA. The calculated binding constant is 5.67 × 10$^5$ M$^{-1}$. The binding constant reported for the 1:2 complex with DNA is nearly the same value as the binding constant for the DNA–H$_2$acacnn complex divided by the 1:1 $\beta$-CD–H$_2$acacnn binding constant. The reciprocal plot showing the relative changes of absorbance of H$_2$acacnn–$\beta$-CD complex with variation in the DNA concentration is given in SI 4d in the Supporting Information. A decrease of binding constant of the binding interaction between H$_2$acacnn and DNA in the presence of $\beta$-CD was observed. A similar binding titration between H$_2$acacnn–$\beta$-CD and DNA was also done using fluorescence spectroscopy (Figure 4d). The addition of DNA to the $\beta$-CD-bound H$_2$acacnn leads to a quenching of fluorescence, and the Stern–Volmer plot of the quenching is shown in SI 4e in the Supporting Information. The $K_{SV}$ is 1.74 × 10$^4$ mol$^{-1}$ L$^{-1}$. The binding constant determined for the binding of H$_2$acacnn–$\beta$-CD to DNA is 2.07 × 10$^5$ M$^{-1}$, using the plot as in SI 4f in the Supporting Information. The mode of binding of H$_2$acacnn to DNA can be visualized as given in Scheme 2.

Scheme 2. Pictorial representation of the H$_2$acacnn–DNA association.

**Molecular docking**

The interaction profile of $\beta$-CD with H$_2$acacnn shows the existence of hydrogen, electrostatic, and hydrophobic interactions SI 5a in the Supporting Information. The interaction provides a the glide score of −1.99 kcal mol$^{-1}$ at the binding site, aminopentenone moiety of H$_2$acacnn, and the secondary hydroxy groups of $\beta$-CD through the hydrogen bonds with the bond distances of 2.42 Å and 1.96 Å. The entry of the aminopentenone moiety of H$_2$acacnn into the hydrophobic cavity of $\beta$-CD is shown in Figure 3b. This observation is in accordance with the experimental binding studies. The types of interactions of H$_2$acacnn with DNA and the hydrogen bond length are given in SI 5 in the Supporting Information. H$_2$acacnn provides the best glide score with a G-score of −1.99 kcal mol$^{-1}$ for binding to $\beta$-CD and −3.67 kcal mol$^{-1}$ for binding to DNA. H$_2$acacnn is found to interact with DNA in two different sites by means of hydrogen bonding with a bond length of 2.113 Å and 9.068 Å in both the A and B-chains of DNA. The docked structure is shown in SI 5b in the Supporting Information. The mode of binding is pictorially represented in Scheme 2.

**Logic gates**

In Figure 5a, the change of absorbance of H$_2$acacnn with the concentration of protons and $\beta$-CD is represented. The absorbances at 235 nm and 215 nm are considered as output 1 and 2 respectively. When the pH is 6, that is, when the proton con-
In such a condition, the operation of AND and NOR gate operations are observed regardless alone, whereas in all the other conditions as shown in Figure 5b, the output 2 is a digital 1 when the input is smaller at 235 nm. The protonated H-acacnn in the presence of β-CD shows OFF and ON signals at λ_{325} and λ_{315}, respectively. Hence, the system functions as a two-input XOR logic gate at λ_{325} and as a Buffer at λ_{315}. The truth table listing the inputs and outputs is represented in Figure 6a.

Figure 5b shows the fluorescence spectra of H-acacnn in its various forms viewed as operating logic gates at λ_{400} and λ_{450}. With a threshold intensity set-up at 100, and considering the inputs as H⁻ and β-CD, the output 1 (λ_{400}) results in a digital 1 (or ON) initially when there is no input, and in a digital 0 (or OFF) when the inputs are either H⁻, β-CD, or both. This suggests that H-acacnn can function as a NOR logic gate at λ_{400}. Similarly, at λ_{450}, the output 2 is a digital 1 when the input is H⁻ alone, whereas in all the other conditions as shown in Figure 3b, it is a digital 0. The truth table and the logic gates are shown in Figure 6b. Hence at λ_{450} the system operates as an INHIBIT logic gate. Considering the same fluorescence spectra of H-acacnn with the same inputs, but having the threshold set at the intensity of 50 units, the digital output at λ_{400} shows the function as a Buffer as opposed to the INHIBIT logic gate observed when the threshold value is 100 (discussed above). However, at λ_{400}, NOR gate operations are observed regardless of the threshold intensity of fluorescence. Figure 6c shows the truth table and the logic gate symbols corresponding to the above logic operations using fluorescence with threshold intensity value 50.

Figure 6a.

Considering the absorbances of the free- and the β-CD-bound H-acacnn interacting with DNA (Figure 5c), the changes are monitored in the view of digital outputs. The inputs 1 and 2 are β-CD and calf thymus DNA respectively. The outputs are read out at λ_{265} and λ_{260}. With no inputs or with single input of β-CD, the output 1 (at λ_{260}) is a digital 0. With a single input of DNA alone, or β-CD/DNA, the output 1 is a digital 1. Hence at λ_{265} the operation of a Buffer gate is observed. At λ_{260}, the output 2 is a digital 1 when there is no input or there are the inputs of β-CD or DNA alone. When both β-CD and DNA are the inputs, the output 2 is a digital 0. Hence, it is an operation of a NAND logic gate at λ_{325}. The truth table and the symbols of the two-input logic gates are given in Figure 6d.

Cancer cells show an aberrant regulation of hydrogen ion dynamics. This results in a reversal of the intracellular to extracellular pH gradient in cancer cells as compared to normal cells. This perturbation in pH dynamics rises very early in carcinogenesis and is one of the most common pathophysiological hallmarks of tumors. In such a condition, the operation of the logic gates discussed above is relevant in the sense that H⁺ ions are involved in the pH-dependent signal on/off switching of the H-acacnn absorption and fluorescence, and β-CD can mimic the movement-restricted microenvironment of tumors. Moreover, a frequently used method for anticancer treatment is the application of a drug that binds to DNA, which affects cell proliferation. β-CD can tune the drug binding to DNA, and the small molecule–DNA–β-CD chemical equilibrium can be mimicked by the logic gates discussed in the preceding paragraphs involving DNA and β-CD.

Conclusions

In the (Z)-5-{5-[(Z)-4-oxopent-2-en-2-ylamino)naphthalen-1-ylamino]pent-3-en-2-one (H-acacnn) crystal, the terminal moieties are oriented along the NH-substituted naphthyl rings. β-Cyclo-dextrin forms a 1:2 inclusion complex with H-acacnn. Fluorescence enhancement on the complex formation occurs at a higher concentration range of β-CD. The aminopentenone chains of H-acacnn are encapsulated by β-CD. In the lower concentration range of β-CD, since only one aminopentenone chain (which does not match the cavity size) gets into the host cavity, the complex gets greatly stabilized only at the formation of the 1:2 complex. The naphthyl rings are not encapsulated by β-CD as it can approach the rings only by sliding over the aminopentenone chains. The methyl end groups of the chains come outside the narrower rim of β-CD after entering through the larger rim. The pHₐ of H-acacnn is very slightly different between the measurements in water and in...
aqueous β-CD media. The case is similar for the excited-state pKa, and this means that the prototropic equilibrium is attained in the ground state and does not greatly vary in the excited state.

The system is suitable for functioning as a reliable logic gate platform as the binding to either β-CD or DNA is sufficiently strong. Binding to β-CD and to DNA results in the possible operation of different logics, and this can be related to the different modes of binding in these events and to the types of bonding and hydrophobic interactions involved. These signaling differences are relevant in human pathophysiological conditions. Both universal gates (NAND, NOR) and basic logical operators (Buffer, INHIBIT) are possible with this simple naphthalene-derivative-based molecular logic substrate. In mimicking the molecular scale biodynamics, this approach based on binding events is advantageous over photochemical switching due to its photostationary state dependence and the laggardness of the latter.

**Chemicals and preparation of solutions**

Analytical-reagent-grade β-CD (Sigma–Aldrich) and calf thymus DNA (Genei, India) were used as received. The purity of DNA was tested before the spectral studies. The absorbance ratio (A_{260}/A_{280}) of DNA was greater than 1.8. The pH of the solutions was adjusted using a phosphate buffer, and the lower pH solutions (< pH 2) were prepared according to the modified Hammett’s acidity scale using sulfuric acid. Spectral grade solvents were used as received.

Stock solutions of H$_2$acacnn and β-CD were prepared in MeOH and triple-distilled H$_2$O, respectively. The stock solution of DNA was made in 50 mM of NaCl. Test solutions of H$_2$acacnn in H$_2$O/β-CD were prepared at pH 7 for the study of the inclusion complex formation. All the test solutions contained H$_2$acacnn in 1% MeOH (v/v) according to its appropriate dilution. Aliquots of DNA used for the DNA–H$_2$acacnn binding studies were made at pH 7 by diluting the stock solution. The experiments were done at 25 ± 2 °C. All the test solutions were homogeneous after the addition of the respective additives.

### Table 1: Truth tables showing various logic gates.

| INPUT (H$^+$) | OUTPUT (H$^+$) |
|---------------|----------------|
| A (H$^+$)     | B (β-CD)       |
| A (H$^+$)     | B (β-CD)       |
| A (H$^+$)     | B (β-CD)       |

### Table 2: Truth tables showing various logic gates.

| INPUT (Threshold: 100) | OUTPUT (Threshold: 100) |
|------------------------|------------------------|
| A (H$^+$)              | B (β-CD)               |
| A (H$^+$)              | B (β-CD)               |
| A (H$^+$)              | B (β-CD)               |

### Table 3: Truth tables showing various logic gates.

| INPUT (Threshold: 50) | OUTPUT (Threshold: 50) |
|-----------------------|------------------------|
| A (H$^+$)             | B (β-CD)               |
| A (H$^+$)             | B (β-CD)               |
| A (H$^+$)             | B (β-CD)               |

### Table 4: Truth tables showing various logic gates.

| INPUT (Threshold: 50) | OUTPUT (Threshold: 50) |
|-----------------------|------------------------|
| A (DNA)               | B (β-CD)               |
| A (DNA)               | B (β-CD)               |
| A (DNA)               | B (β-CD)               |

Figure 6. Truth tables showing various logic gates. a) Results of the absorption spectra (threshold absorbance: 0.8). b) Results of the fluorescence spectra (threshold intensity: 100). c) Results of the fluorescence spectra (threshold intensity: 50). d) Results of the absorption spectra (threshold absorbance: 0.15).
Instruments and recording of spectra
UV/Vis and fluorescence spectra were recorded using a double-beam Jasco V630 spectrophotometer (Mary's Court, USA) and a PerkinElmer LS55 spectrophotometer (Waltham, USA), respectively. To record the spectra, cuvettes of path length 1 cm were used. The spectrophotometer used a 120 W xenon lamp as the excitation source, and the excitation and emission bandwidths were fixed at 4 nm. pH was measured using an Elco Li 121 pH meter (Hyderabad, India). The absorption spectra were recorded against appropriate reference solutions which did not contain H₂acacn.

Microanalysis of the complexes was done using a PerkinElmer PE 2400 series II CHNS/O elemental analyzer. Infrared (IR) spectra were recorded using KBr pellets (1% w/w) on a PerkinElmer Spectrum GX FT-IR spectrophotometer. Electronic spectra were recorded on a Shimadzu UV 3101PC spectrophotometer (Kyoto, Japan). Mass spectrometric analysis was performed using an electron spray ionization (ESI) technique on an LC Waters Q-TOF-micro mass spectrometer (Milford, USA) (spectrum shown in SI 6 in the Supporting Information).

Molecular docking
The structure of calf thymus DNA was retrieved from the RCSB Protein Data Bank (PDB) based on resolution of the structure, experimental feasibility by matrix data, comparative value of crystallographic model, and X-ray diffraction data. A 10% removal of data for the structure observed and a comparison of 90% of the structure with the crystallographic model yielded the module structure. Docking was performed using Maestro v. 9.6 software (Schrodinger, New York, USA) to optimize the interaction of H₂acacn with β-CD and calf thymus DNA. The structure of β-CD was built and optimized by molecular mechanics. The structure of calf thymus DNA was downloaded from the PDB (ID No. 3GJH) and preprocessed prior to docking. The DNA grid was set up and generated from the receptor grid generation panel.

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Keywords: β-cyclodextrin • fluorescence spectroscopy • host–guest interaction • molecular logic gates • naphthalene • supramolecular chemistry

Preparation of H₂acacn
Naphthalene 1,5-diamine (0.002 mol), acetylacetone (0.002 mol), and few drops of 6.2 M HOAc were mixed in EtOH (50 mL) and stirred constantly at reflux for 3 h and cooled to rt. The mixture was slowly evaporated in vacuo at rt for 48 h to afford yellowish-brown crystals (309 mg, 96%). 1H NMR (CDCl₃, 500 MHz): δ = 12.78 (broad s, 2H), 7.97, 7.95 (d, J = 10 Hz, Ar–CH), 7.52, 7.50, 7.49 (t, J = 10 Hz, 5H z, Ar), 7.32, 7.31 (d, J = 5 Hz, Ar–CH, 2H), 5.32 (s, =CH₂, 2H), 2.17 (s, CH₃, 6H); IR (KBr): ν = 4000—3600 (br), 3500 (br), 2995, 1604, 1573, 1516, 1433, 1355, 1274, 1020, 956, 788 cm⁻¹; ESI-MS m/z [M + 1]⁺ calcld for C₇H₁₄N₂O₃: 222.40, found 223.56; Anal. calcld for C₇H₁₄N₂O₃: C 74.51, H 6.88, N 8.69, found: C 74.72, H 6.59, N 8.78.

Single-crystal X-ray analysis
A crystal of suitable size was selected and mounted on the tip of a glass fiber and cemented using epoxy resin. Intensity data for the crystal was collected using Mo-Kα (λ = 0.71073 Å) radiation on a Bruker SMART APEX diffractometer equipped with CCD area detector. The hydrogen atom H1N, which is bound to the nitrogen atom N1, was located in the difference Fourier synthesis, and was refined semifreely with the help of a distance restraint, while constraining its U-value to 1.2 times the U(eq) value of N1. All other hydrogen atoms were placed in calculated positions and refined by using a riding model.

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