

Combining Excellent Selectivity with Broad Target Scope: Biosensing with Arrayed Deep Cavitand Hosts

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Cite This: *Acc. Chem. Res.* 2022, 55, 1035–1046



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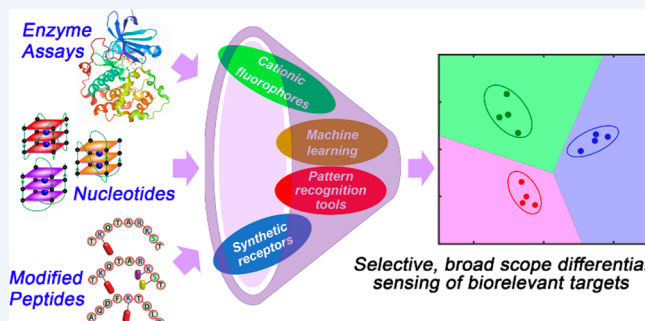
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CONSPECTUS: Simple macrocyclic water-soluble hosts such as cucurbiturils, cyclophanes, and calixarenes have long been used for biosensing via indicator displacement assays. Using multiple hosts and dyes in an arrayed format allows pattern recognition-based “chemical nose” sensing, which confers exquisite selectivity, even rivaling the abilities of biological recognition tools such as antibodies. However, a challenge in indicator displacement-based biosensing with macrocyclic hosts is that selectivity and scope are often inversely correlated: strong selectivity for a specific target can limit wide application, and broad scope sensing can suffer from a lack of selectivity between similar targets. This problem can be addressed by using water-soluble, self-folding deep cavitands as hosts. These flexible bowl-shaped receptors can be easily functionalized with different motifs at the upper and lower rim, and the large cavities can bind many different fluorescent dyes, causing either fluorescence enhancement or quenching upon binding.

Cavity-based affinity is strongest for NMe_3^+ groups such as trimethyl-lysine, and we have exploited this for the site-selective recognition of post-translational lysine methylations in oligopeptides. The host recognizes the NMe_3^+ group, and by applying differently functionalized hosts in an arrayed format, discrimination between identical modifications at different positions on the oligopeptide is possible. Multiple recognition elements can be exploited for selectivity, including a defined, yet “breathable” cavity, and variable upper rim functions oriented toward the target.

While the performance of the host/guest sensing system is impressive for lysine methylations, the most important advance is the use of multiple different sensing mechanisms that can target a broad range of different biorelevant species. The amphiphilic deep cavitands can both bind fluorescent dyes and interact with charged biomolecules. These non-cavity-based interactions, when paired with additives such as heavy metal ions, modulate fluorescence response in an indirect manner, and these different mechanisms allow selective recognition of serine phosphorylation, lysine acetylation, and arginine citrullination. Other targets include heavy metals, drugs of abuse, and protein isoforms. Furthermore, the hosts can be applied in supramolecular tandem assays of enzyme function: the broad scope allows analysis of such different enzymes as chromatin writers/erasers, kinases, and phosphatases, all from a single host scaffold. Finally, the indirect sensing concept allows application in sensing different oligonucleotide secondary structures, including G-quadruplexes, hairpins, triplets, and i-motifs. Discrimination between DNA strands with highly similar structures such as G-quadruplex strands with bulges and vacancies can be achieved. Instead of relying on a single highly specific fluorescent probe, the synthetic hosts tune the fluorophore–DNA interaction, introducing multiple recognition equilibria that modulate the fluorescence signal. By applying machine learning algorithms, a classification model can be established that can accurately predict the folding state of unknown sequences. Overall, the unique recognition profile of self-folded deep cavitands provides a powerful, yet simple sensing platform, one that can be easily tuned for a wide scope of biorelevant targets, in complex biological media, without sacrificing selectivity in the recognition.

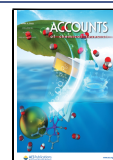


KEY REFERENCES

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Received: January 15, 2022

Published: March 18, 2022



- oligopeptides and an analysis of the dual-mode recognition properties of the hosts.*
- Liu, Y.; Lee, J.; Perez, L.; Gill, A. D.; Hooley, R. J.; Zhong, W. Selective Sensing of Phosphorylated Peptides and Monitoring Kinase and Phosphatase Activity with a Supramolecular Tandem Assay. *J. Am. Chem. Soc.* **2018**, *140*, 13869–13877.² By changing the nature of the dye, we could alter the mechanism of sensing of the arrayed host/dye complexes, selectively targeting serine phosphorylation rather than lysine methylation. This also includes applications in monitoring kinase and phosphatase activity.
 - Liu, Y.; Perez, L.; Gill, A. D.; Mettry, M.; Li, L.; Wang, Y.; Hooley, R. J.; Zhong, W. Site-Selective Sensing of Histone Methylation Enzyme Activity via an Arrayed Supramolecular Tandem Assay. *J. Am. Chem. Soc.* **2017**, *139*, 10964–10967.³ In this work, the sensor arrays were applied to monitor enzymatic activity of methyltransferases and demethylases. The arrayed host/guest complexes were functional in the presence of the enzymes and their cofactors, and site-selective enzyme activity monitoring was possible.
 - Chen, J.; Hickey, B. L.; Wang, L.; Lee, J.; Gill, A. D.; Favero, A.; Pinalli, R.; Dalcanale, E.; Hooley, R. J.; Zhong, W. Selective Discrimination and Classification of G-Quadruplex Structures with a Host:Guest Sensing Array. *Nat. Chem.* **2021**, *13*, 488–495.⁴ The “indirect” sensing mechanism of arrayed host/dye complexes was exploited to detect structural changes in DNA G-quadruplexes, including discrimination and classification of structures that varied only in backbone topology.

INTRODUCTION

The field of biosensing has been greatly expanded in recent years by the application of macrocyclic synthetic receptors.⁵ To function as an effective host sensor for biological species, the synthetic receptor must contain a defined cavity into which the target species can bind and also allow pairing with a suitable optical reporter molecule. This dye can be displaced upon recognition of the target, providing an optical output that converts simple recognition into “sensing”, using indicator displacement assays (Figure 1a).⁶ Small molecule hosts (as opposed to surface-attached sensors⁵) are also greatly improved if they are water-soluble. Achieving all these requirements is not trivial, and the “best” (i.e., most popular) receptors for this process are macrocycles such as cucurbit[*n*]urils (CBs),⁷ cyclophanes,⁸ sulfonato-calixarenes (CXs),⁹ and shallow phosphonate cavitands¹⁰ (Figure 1c), as well as other aromatic macrocycles such as pillararenes.¹¹ Each receptor type has its advantages and disadvantages, and some exquisite applications have been shown that highlight the creativity and technical prowess of the scientists involved.¹² Substrate selectivity can be enhanced by applying the hosts as differential sensors.¹³ By use of multiple hosts and fluorophores in an array, a fluorescence fingerprint can be generated that, after statistical data processing,¹⁴ provides a simple output that allows differentiation and classification of highly similar substrates (illustrated in Figure 1b). The differential sensing strategy allows synthetic receptor-based sensors to approach levels of selectivity in biosensing that are usually only associated with antibodies or other platforms that exploit biomacromolecules, with none of the synthetic complexity or expense associated with biological sensors.¹⁵

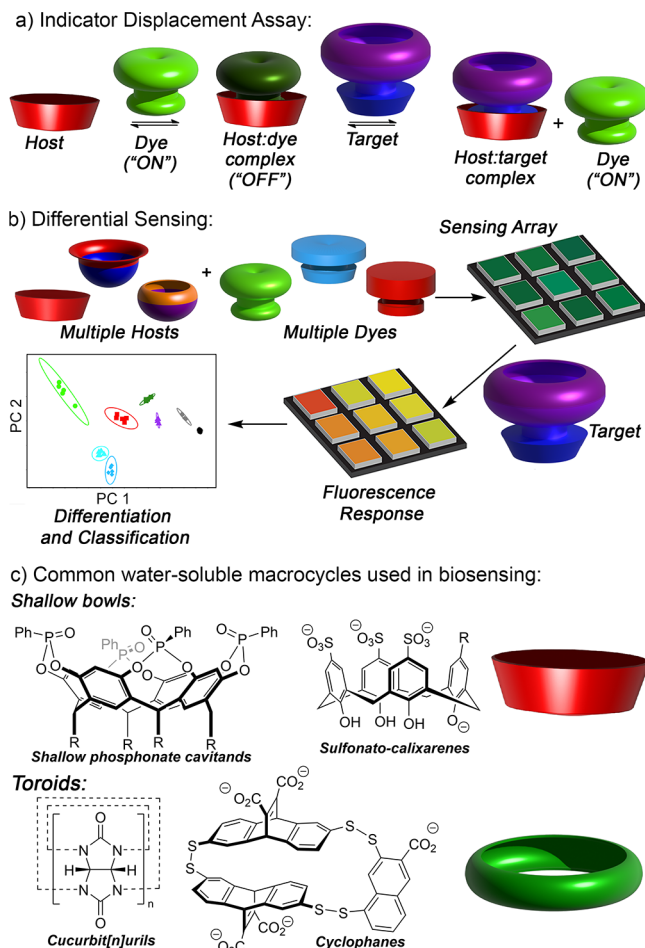


Figure 1. Synthetic receptors as biosensors. (a) Illustration of the indicator displacement assay concept; (b) extension to differential sensing; (c) examples of other well-known macrocycles commonly used for molecular recognition and sensing in water.

Challenges still remain, however. The major limitation with using these privileged hosts is that they tend to favor specific targets: each host binds one specific type of molecular structure very strongly. For example (and with apologies for this simplistic generalization), the “best substrates” for CBs are *N,N'*-alkyldiammonium ions,¹⁶ calixarenes and phosphonate cavitands favor methylammonium ions (R-NMe_3^+ ¹⁷ and $\text{R-NH}_2\text{Me}^+$ ^{18,19} respectively), and cyclophanes favor soft cations such as methylated ammonium²⁰ or guanidinium ions.²¹ Molecular tweezers are also effective and target unmodified lysines in proteins.^{22,23} All these species are essential molecular markers and can be found as post-translational modifications (PTMs) in proteins,²⁴ drugs of abuse,²⁵ and neurotransmitters,²⁶ among others. They are quite similar in structure to each other, however, mostly falling into the category of “soft nitrogen-containing cations”, and as such, many of the most impressive applications involve a narrow subset of targets. Other vital targets such as small anions,²⁷ lipids,²⁸ sugars,²⁹ phosphates,³⁰ and nucleotides³¹ require other types of sensors for detection, with varying degrees of success.

Obviously, these challenges are endemic for biosensing with small molecule receptors: small, inflexible cavities cannot rival antibodies or RNA for both substrate binding affinity and scope, as their defined binding site is limited in the size, shape, and charge of groups that can be recognized. As a result, there is an

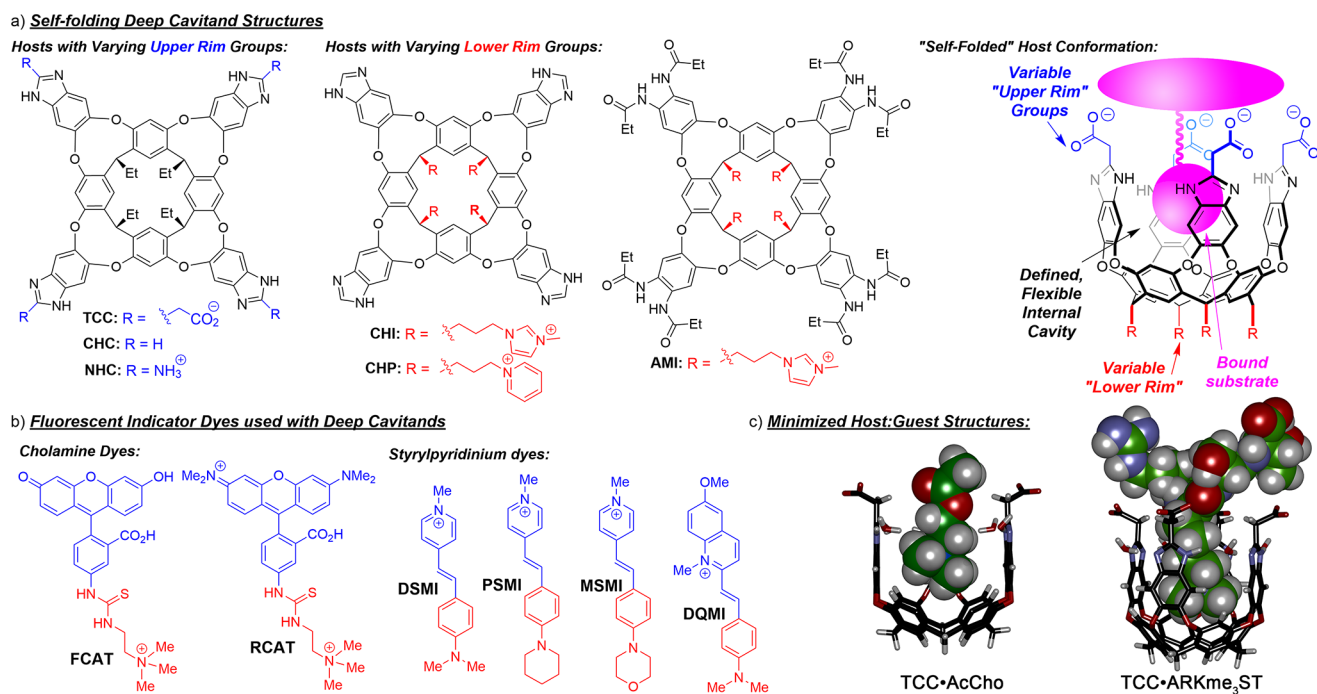


Figure 2. Structures of sensor components. (a) Deep cavitand structures used for sensing biological targets, and the folded conformation in solution; (b) suitable fluorophores; (c) minimized structures of host/guest complexes between TCC and R-NMe₃⁺-containing guests.

unfilled niche in this area: can more flexible, complex synthetic receptors be used that exploit multiple different recognition and sensing mechanisms for different types of targets and are not restricted to sensing via 1:1 host/guest binding and indicator displacement assays? In this Account, we describe our work on biosensing with water-soluble, flexible deep cavitands and hopefully illustrate the power of these hosts to combine both strong selectivity and broad scope in sensing a wide array of complex biological targets.

MOLECULAR RECOGNITION WITH WATER-SOLUBLE DEEP CAVITANDS

Self-folding deep cavitands, pioneered by Rebek,^{32–34} can act as host molecules for a variety of targets by presenting a large, defined cavity with a single entry portal to suitably sized guests. These larger cavities introduce a challenge for solubility (especially in water) and conformational stability: species with large open cavities tend to fill those spaces, by either aggregation,³⁵ dimerization,³⁶ or simply adopting an “undesired” conformation.³⁷ One solution is to synthesize large, rigid deep cavitands, which has been explored to great success by Gibb.³⁸ Another technique is to use flexible walls and rely on “self-folding” for conformational stability. This can be achieved by introducing self-complementary hydrogen bonding groups at the upper rim of the hosts. Depending on the groups used, the self-folding technique can even confer conformational stability on the hosts in water. This, of course, is essential for biorecognition, which at minimum must be performed in aqueous solution and hopefully in complex biological media. Water solubility is usually conferred on the cavitands by incorporating charged functional groups, at either the upper or lower rim. We have generally focused on the benzimidazole deep cavitand scaffold (Figure 2a), which uses water molecules as the hydrogen-bonding agents to ensure proper folding. It should be noted here that Rebek and Yu have investigated similar types of cavitand scaffolds as small molecule hosts and reactors.³⁹

Our most successful cavitand for biorecognition purposes is the anionic cavitand TCC (Figure 2a).⁴⁰ When folded, the cavity is highly electron rich, with 8 aromatic panels surrounding an internal space of ~180 Å³. This is inspired by the acetylcholinesterase active site, which consists of an “aromatic box” of tryptophan and tyrosine side chains.⁴¹ As such, TCC is an excellent host for R-NMe₃⁺ ions (Figure 2c), notably choline and acetylcholine^{40,42} and can also bind hydrocarbons of the correct size and shape.⁴³ Recognition of R-NMe₃⁺ ions is generally both enthalpically and entropically favorable. Cation-π and CH-π interactions between host and guest provide an enthalpic driving force, and as R-NMe₃⁺ ions are relatively “greasy”, entropically favorable desolvation of the surrounding water molecules also occurs upon binding. Importantly, as the cavitand is open-ended, the selectivity extends to a wide range of “R” groups. Examples include surfactants,⁴⁴ proteins,⁴⁵ polymers,⁴⁶ and modified fluorophores.⁴⁷

There is one more important trait shown by TCC: as the charged carboxylates reside only on the upper rim of the host, it is lipophilic and quite prone to self-aggregation. This has allowed application as a host and sensor in membrane bilayers^{44–46,48} but also has a large (and complex) effect on the recognition and sensing of charged and hydrophobic species in water, which is one of the keys to its broad target scope.

SELECTIVE RECOGNITION AND SENSING WITH ARRAYED CAVITAND·FLUOROPHORE COMPLEXES

While the molecular recognition aspects of deep cavitands have been extensively analyzed by NMR, coupling this recognition to an optical output is more challenging. Indicator/fluorescence displacement assays⁶ have been widely used with calixarenes (especially tetrasulfonatocalix[4]arene, CX4,⁴⁹ or tailored CX4 derivatives⁵⁰), cucurbit[n]urils,⁷ cyclophanes,²⁰ and pillararenes.¹¹ Unfortunately, common fluorophores such as lucigenin (LCG) do not bind in deep cavitands, so we have focused on two

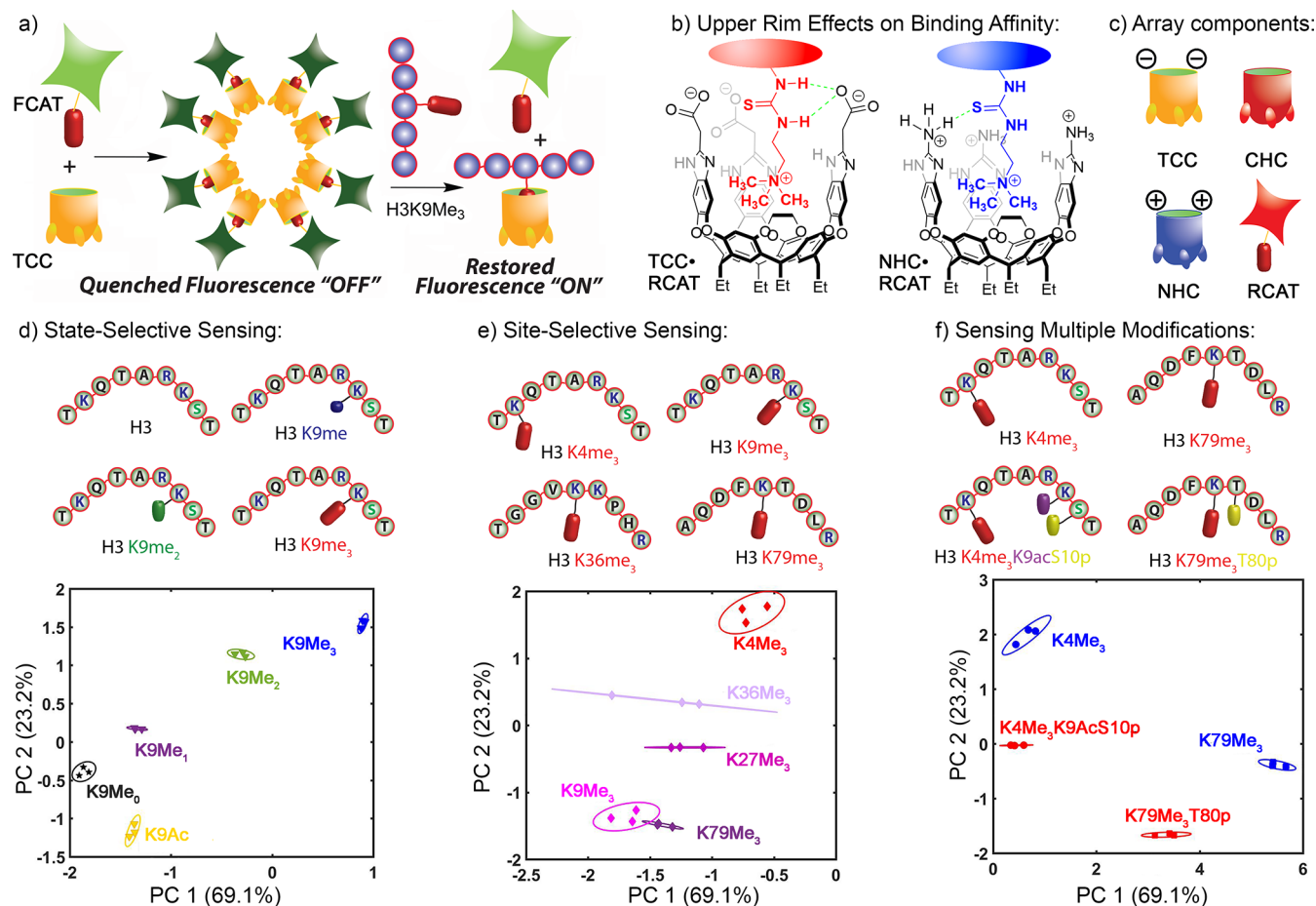


Figure 3. Differential sensing of lysine trimethylations in oligopeptides.¹ (a) Illustration of the aggregation-dependent mechanism of indicator displacement shown by the TCC cavitand; (b) upper rim groups provide a second layer of binding selectivity; (c) array components used for differential sensing. Subsets of the PCA scores plot from the observed fluorescence data indicating discrimination of (d) methylation state, (e) methylation site, and (f) multiple methylations. Adapted with permission from ref 1, copyright 2017, Royal Society of Chemistry.

different types of fluorophores: combinations of fluorescein or rhodamine isothiocyanates with cholamine (FCAT and RCAT, respectively), and styrylpuridinium dyes such as DSMI, PSMI, MSMI, and DQMI (Figure 2b). Each of these dyes shows differing affinity and fluorescence response with different cavitands, which is ideal for broad target scope and differential sensing. Our initial focus was to establish the scope of the recognition/sensing system for peptides modified with methylated lysines: as cavitands such as TCC, CHC, and NHC bind R-NMe₃⁺ ions, they are perfectly suited for selective recognition of trimethyl-lysine groups. Using arrayed deep cavitands has a key advantage over simple hosts: while the cavities in different deep cavitands are essentially identical, the upper rim functional groups are oriented directly at the peptide backbone. This introduces secondary interactions with neighboring side chains, which, in addition to recognizing the Kme₃ group, can be used to further increase target selectivity.

The simple TCC-FCAT host/dye pair is a powerful sensor for Kme₃ groups in oligopeptides.⁵¹ The mechanism is slightly more complex than simple displacement: upon binding, TCC causes 80% quenching of FCAT via a triggered aggregation mechanism (Figure 3a). The host/guest binding interaction between TCC and FCAT triggers the formation of a micellar aggregate of the amphiphilic cavitand, and FCAT is self-quenched upon aggregation. Upon addition of the modified histone H3 peptide H3K9me₃(1–21), the Kme₃ residue outcompetes the dye,

displacing it and turning on the sensor. Neither the parent unmethylated peptide (H3K9) nor the H3K9me variant caused appreciable recovery of the fluorescence signal: this simple cavitand-dye sensor is *state-selective* for different lysine methylations. It should be noted that this selectivity, while an exciting proof-of-principle for us, is also seen with other known hosts such as calixarenes⁵² and cyclophanes,⁵³ which show stronger affinities for Kme₃ and greater levels of Kme₃/Kme₂/Kme selectivity⁵⁵ than TCC. However, the most important advantage of the deep cavitand system is its adaptability: state selectivity can be achieved with a single host/guest complex, but for more challenging targets and a finer level of selectivity, multiple arrayed host/guest complexes are needed. The constituents of these arrays can be quite varied, including different cavitand hosts, dyes, or other environmental factors.

The initial array focused on varying cavitand structure and pH: three hosts (negative TCC, neutral CHC and cationic NHC, Figure 2, 3c) were combined with RCAT at different pH values.¹ The thioureas in RCAT can interact synergistically with the cavitand upper rims: the C=S group can favorably interact with acids, whereas the urea NH groups can interact with bases. Biasing the pH of the system changes the balance of these interactions, adding a subtle layer of recognition selectivity between the hosts and dyes (Figure 3b), as well as the hosts and the oligopeptide targets. This effect was shown by discriminating a series of cholamine derivatives¹ and then applied to more

complex methylated lysines. Fourteen peptides were tested, based on the histone H3 sequence, with a variety of different modifications. These included varying the methylation state at K9 (K9me_0 – K9me_3), changing the position of the trimethylation on the backbone (K4/9/27/36/79me₃), combining other PTMs such as phosphorylations and acylations with Kme₃ (e.g., K4me₃K9AcS10p, K79me₃T80p), and varying the peptide length (Figure 3).

A six-factor array was used (the three cavitands (TCC, CHC, and NHC), RCAT dye, in either pH 7.4 or 9.0 buffer). Fluorescence changes were monitored upon addition of peptide to each sensor well in the array, and the outputs were analyzed by principal component analysis (PCA, Figure 3d–f). Differentiating methylation state is simple, but the much greater challenge is to differentiate between identical modifications at different positions on the backbone. This can be achieved with the 6-component array: peptide strands of the same length with Kme₃ at different positions (K4/9/27/36/79) could almost all be differentiated with 95% confidence (Figure 3e). The differentiation here is subtle, as the cavitands all bind the Kme₃ group, but the relative response is affected by interactions between the upper rim groups of the cavitand and the adjacent residues on the peptide backbone. The exact interactions that bias the affinity in each state are not obvious, but two more sets of experiments shed some light on the situation. If modifications are added to residues adjacent to the Kme₃ group (e.g., K79me₃T80p, Figure 3f), significant changes in response are observed, presumably due to the large change in charge gained from converting a threonine OH to a phosphate group. The changes need not be adjacent, however; K4me₃K9AcS10p can be distinguished from K4me₃, which indicates that the larger peptide structure is affecting the recognition. In addition, the sensor can discriminate peptides where the Kme₃ group resides at the same position, but the length of the peptide changes.¹ As opposed to smaller, rigid hosts such as CX4 and CB7 that are highly specific for small molecule targets and functional groups on peptides, the more lipophilic cavitands are affected by the superstructure of long peptides. The triggered aggregation process seen by the TCC·FCAT complex is a good illustration of this: not only are the cavitands affected by what binds *inside* the cavity, but also *external* interactions affect the fluorescence response. This has drawbacks: these hosts are notably poor at sensing individual amino acids, and teasing out the mechanisms of sensing becomes challenging, but the benefits in enhanced selectivity and scope are large.

■ INDIRECT SENSING OF NON-METHYLATION MODIFICATIONS

The next step was to utilize these unexpected, non-cavity-based interactions (i.e., *indirect sensing*) and extend the target scope to modifications other than lysine methylations. The anionic TCC can interact with cationic peptides in solution via charge-based interactions rather than cavity-based recognition. This effect is not dominant when binding Kme₃ groups, as the cavity binding outcompetes charge interactions. However, when 1 is combined with styrylpyridinium dyes such as DSMI (Figure 2b), the TCC·DSMI complex can be disrupted by a cationic peptide, which changes the emission of the dye. This allows host/guest arrays to be applied in targeting *non-methylation* PTMs, using the cavity to bind a suitable dye and the upper rim carboxylates to target cationic peptides.

The most obvious application is one that gives a large change in charge to maximize selectivity, namely, phosphorylation of

alcoholic side chains (Ser, Tyr, Thr).² Figure 4b shows the effect of a series of cationic peptides and their phosphorylated

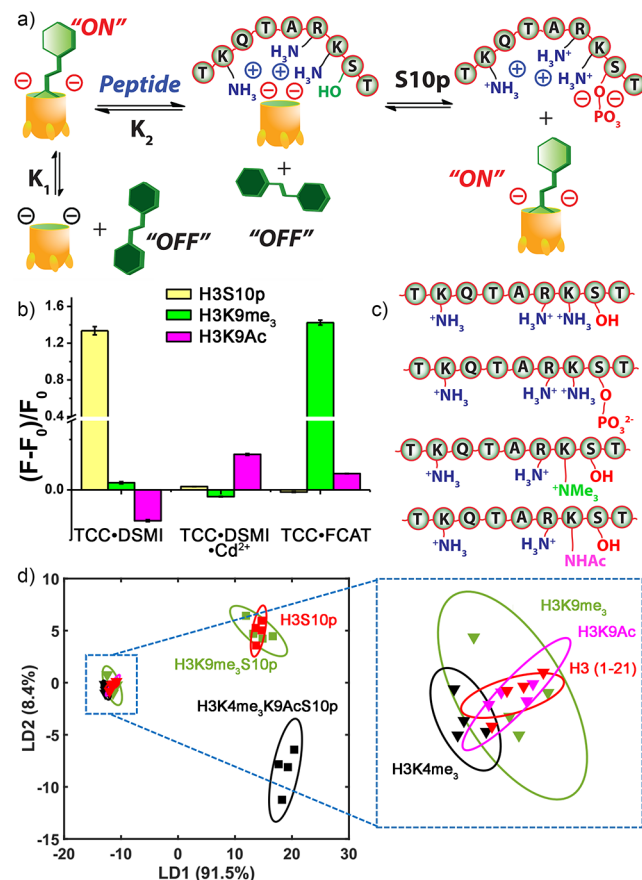


Figure 4. Differential sensing of peptide phosphorylations.² (a) Indirect sensing mechanism; (b) fluorescence responses of modified peptides to different host-dye complexes; (c) representative peptide sequences; (d) PCA score plots from the observed fluorescence data indicating discrimination between phosphorylated peptides and those with other modifications. Adapted from ref 2. Copyright 2018 American Chemical Society.

counterparts on the fluorescence of the host/dye complexes, illustrating the selectivity of TCC·DSMI. Whereas the cationic peptide interacts with the host and competitively displaces DSMI (which is protruding from the top of the host), this interaction is reduced with a phosphorylated peptide. The sensor is not targeting the phosphate group, but the *absence* of cationic peptide interactions. Importantly, DSMI binds more weakly in the TCC than the FCAT or RCAT dyes,^{1,2} and so can be displaced upon addition of the cationic peptide. The more strongly binding dyes can only be displaced by R-NMe₃⁺ groups, so they are minimally affected by unmodified peptides. This allows tailored selectivity for different modifications, depending on the dye used.

To achieve the greatest possible differentiation, arrayed sensing would be ideal, but as only the anionic TCC can interact with the cationic peptides, other non-cavatand elements must be incorporated. Fortunately, TCC shows a surprising affinity for heavy metal ions in solution.⁵⁶ Even though the only functional groups present in the host are carboxylates, micromolar affinity for large metal ions is seen, mainly due to the fact that the carboxylate groups in TCC (when folded) are preoriented to chelate large metals. When TCC is paired with a dye (either

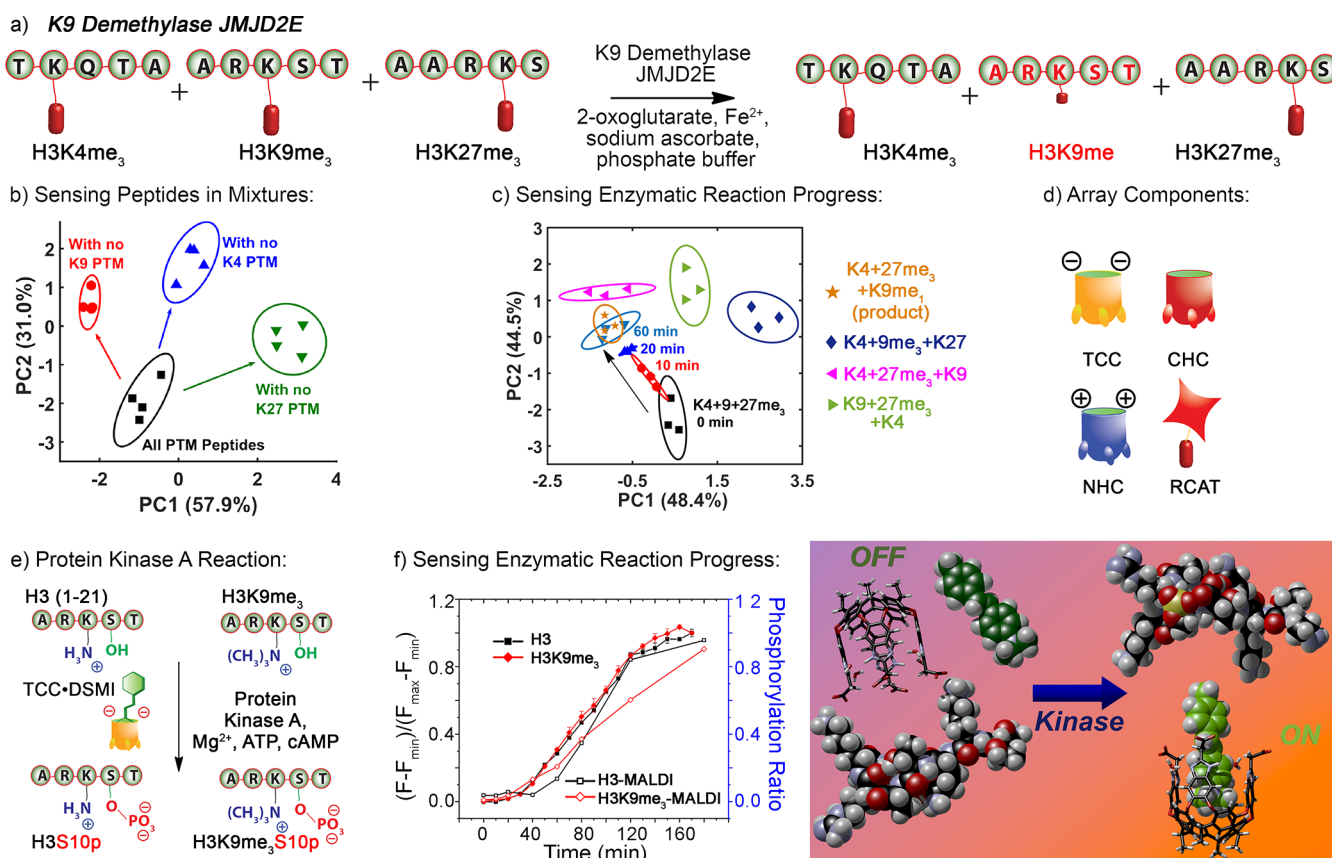


Figure 5. Supramolecular tandem assays. (a) K9 demethylase process;^{3,51} (b) PCA score plots from the observed fluorescence data indicating discrimination between lysine methylations in peptide mixtures;³ (c) PCA monitoring the JMJD2E demethylation reaction; (d) array components used for the JMJD2E assay; (e) protein kinase A dephosphorylation process; (f) cavitand-based monitoring of the protein kinase A reaction.² Reproduced from refs 2, 3, and 51. Copyright 2016–2018 American Chemical Society.

RCAT or DSMI) and metal ions, this introduces multiple different fluorescence responses. The metals can either bind to the TCC-dye complex or displace the dye from the cavitand or both. Also, the fluorescence response of the TCC-dye-M complexes is variable: heavy metals quench the dye, whereas filled shell metals (e.g., Zn²⁺) do not. This has interesting applications in environmental analysis,⁵⁶ but for the purposes of this Account, the most important result is that simply adding 10–50 μM metal ion easily expands the array without needing to synthesize new hosts or dyes.

A 4-component phosphate sensing array, consisting of TCC, DSMI, and Cu²⁺/Co²⁺/Ni²⁺/no metal was exposed to a series of modified peptides as before (some of which are shown in Figure 4c), and the fluorescence responses were analyzed by linear discriminant analysis (LDA). The TCC·DSMI·M²⁺ sensor array can discriminate phosphorylated peptides from their unphosphorylated counterparts, even when those peptides contain Kme₃ modifications. For example, the three phosphorylated peptides H3S10p, H3K4me₃S10p, and H3K4me₃K9AcS10p are fully separated from H3, H3K4me₃, H3K9me₃ and H3K9Ac (Figure 4d). The array is also capable of selectively discriminating phosphorylation at tyrosine or threonine from serine in different peptide strands. Most interestingly, the sensing of peptide phosphorylations can be performed in phosphate buffer! As the target of the sensor is the unmodified peptide, not the phosphate group, the sensing of peptide phosphorylations is tolerant to phosphate in the media. Some degradation of performance is seen in phosphate buffer when

compared to Tris, but the fact that phosphate sensing is functional in phosphate buffer at all is remarkable.

This concept can be extended to other types of non-methylation PTMs, although this becomes significantly more challenging as the charge differences lessen. Lysine acylations can be discriminated from unmodified peptides, as the positive charge in the NH₃⁺ group is removed upon modification. Similarly, arginine citrullinations (Ci) can be targeted:⁵⁷ the change in side chain in this case involves converting the guanidinium group in Arg to a urea, that is, C=NH₂⁺ becomes C=O. After some optimization, a 4-factor array (TCC·FCAT, with no metal, La³⁺, Ni²⁺, or Zn²⁺) was shown to distinguish H3R2_{Ci} and H3R2_{Ci}R8_{Ci} from H3(1–21), so in this case, the sensor shows modification state selectivity. However, positional selectivity is more challenging: discriminating between H3R2_{Ci} and H3R8_{Ci} requires two dyes, two metals, and analysis via 3D PCA for full separation. Still, this is a challenging target for sensing, so even this moderate selectivity is impressive.

SUPRAMOLECULAR TANDEM ASSAYS

Small molecule hosts such as CB[7] and CX4 have been widely used in supramolecular tandem assays of enzyme function,⁵⁸ including cholinesterases,⁵⁹ proteases,⁶⁰ oxidases,⁶¹ lysine methyltransferase,⁶² and kinases,⁶³ among others.^{64,65} These systems usually monitor indicator displacement by either the reactant or product of the enzymatic reaction over time. This is a simple, effective monitoring method that relies on the affinity of small molecules for the host. However, this can be challenging to

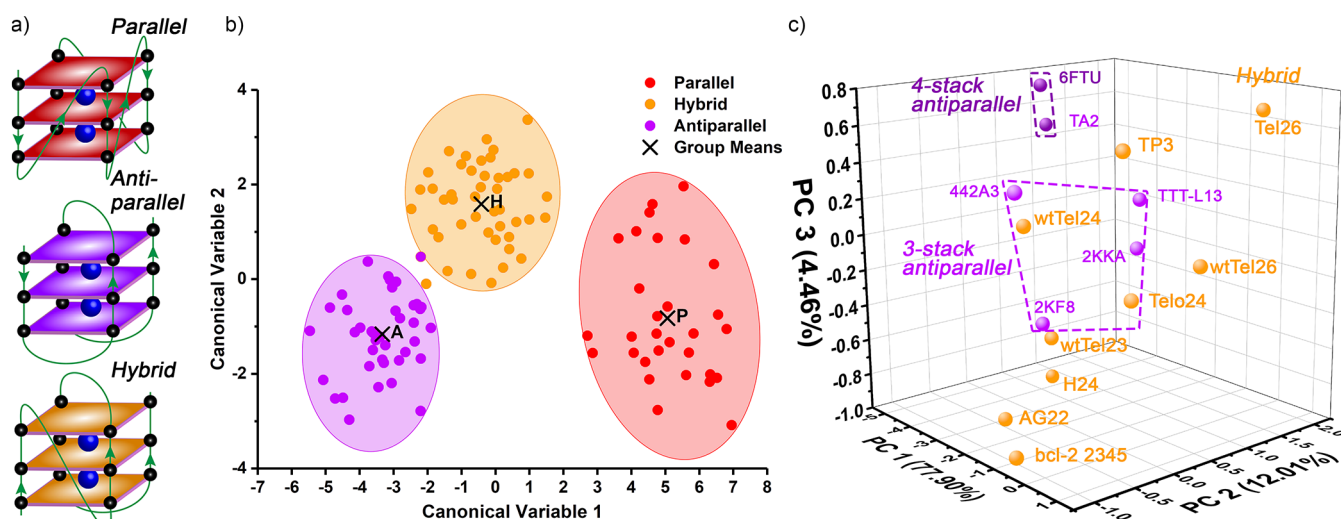


Figure 6. Nucleotide Sensing. (a) Different G4 topologies; (b) CDA score plot of 115 G4 samples, grouped by topology;⁴ (c) 3D PCA score plot of 15 antiparallel and hybrid G4 strands with 10-component cavitand/dye sensing array.⁴

apply for large, complex targets and cannot detect positionally selective enzyme reactions such as those of chromatin writers and erasers. As the deep cavitand sensors can provide positionally selective sensing, the obvious next step was to apply this to enzyme reaction monitoring.^{3,51}

The key to all supramolecular tandem assays is to ensure that the target binding process is not affected by the required conditions for the enzymatic reaction. Whereas simple recognition of Kme₃ peptides can be performed in a controlled aqueous environment, monitoring the function of lysine demethylase JMJD2E requires that the sensor be tolerant to the enzyme itself, as well as cofactors such as Fe²⁺, sodium ascorbate, and 2-oxoglutarate. Fortunately, the TCC·FCAT sensing pair is not affected by these conditions. While Fe²⁺ has some effect on the fluorescence response, it is small enough in this case to not affect peptide detection. Upon addition of TCC·FCAT to a solution of H3K9me₃ in the presence of JMJD2E and required cofactors, the modified peptide displaces FCAT from the host, “turning on” the fluorophore.⁵¹ Over time, as the enzyme removes the methyl groups from H3K9me₃, the fluorophore is reclaimed by the host, and the drop in fluorescence over time allows monitoring of enzyme activity, including determining the K_i of a suitable inhibitor.⁵¹ This shows that the TCC·FCAT sensing pair can monitor lysine demethylase activity, but this is not unique, as Nau has shown this type of assay using CB[7].⁶² However, the cavitand-based arrays are capable of much more than simply monitoring activity: by using multiple cavitands and dyes, site-selective monitoring of enzyme activity is possible in complex mixtures of modified peptides.³ The arrayed tandem assay follows the same principle as the simple one-component FDA process: Kme₃ groups displace fluorophores from the cavitand(s), and enzymatic removal of those methyl groups allows recombination of the cavitand-dye complex, lowering the observed fluorescence. In this case, a three-component methylation-sensitive array was used, with TCC, CHC, and NHC cavitands and RCAT dye at pH 7.4 (to ensure proper enzyme function). For monitoring the site-selective JMJD2E demethylation process, three modified peptides were combined and subjected to enzymatic demethylation (Figure 5a): K4me₃, K9me₃, and K27me₃. Only K9me₃ is demethylated by JMJD2E, and the product is the monomethylated peptide K9me. The reactions

were added to each array well, and the sensor responses at various time points were subjected to PCA, together with a series of controls corresponding to all possible outcomes of the reaction (i.e., demethylation of the other peptides as well as the “correct” target).

As the reaction proceeds (Figure 5a), the reaction mixture moves toward the position where K9me + K4me₃ + K27me₃ is located, and further away from the other two controls, indicating that the enzyme acts on only the K9 methylation site, but not K4 nor K27 (Figure 5b,c). The analysis is capable of distinguishing not only incorrect reaction products but also the state of the product: fully demethylated K9 is distinguishable from K9me. This methylation-sensitive array is also capable of site-selective monitoring of methylation writer enzymes.

As well as sensing the “obvious” targets of lysine methylation enzymes with cavitand-dye arrays, the indirect sensing concept can be used to monitor kinase (Figure 5e) and phosphatase activity. The challenge here is that kinases require the presence of both ATP and cAMP for reaction, which is quite difficult for sensors that detect the phosphate group! Fortunately, as the TCC·DSMI sensor shows no affinity for small phosphates and its fluorescence is mediated by the nature of the cationic peptide, it is capable of monitoring kinase activity. Figure 5f shows one example of the sensing performance: phosphorylation of peptides H3(1–21) and H3Kme₃ by protein kinase A was monitored by both the TCC·DSMI sensor and MALDI-MS analysis of aliquots drawn from solution. The fluorescence changes follow the phosphorylation process well, and this sensor is not affected by the presence of a Kme₃ group, despite using the same cavitand host. Other examples of enzymes that can be analyzed include aurora kinase A and alkaline phosphatase.²

■ INDIRECT SENSING MECHANISMS FOR STRUCTURAL ANALYSIS OF DNA 3D STRUCTURE

The indirect sensing mechanism for phosphate sensing illustrates that cavitands are capable of sensing biorelevant targets that do not bind in the cavity, as there are multiple mechanisms of interaction and fluorescence response that can be exploited. The logical extension of this concept is to use the hosts as fluorescence modulators rather than the recognition element, that is, use dyes as the primary recognition motif, and

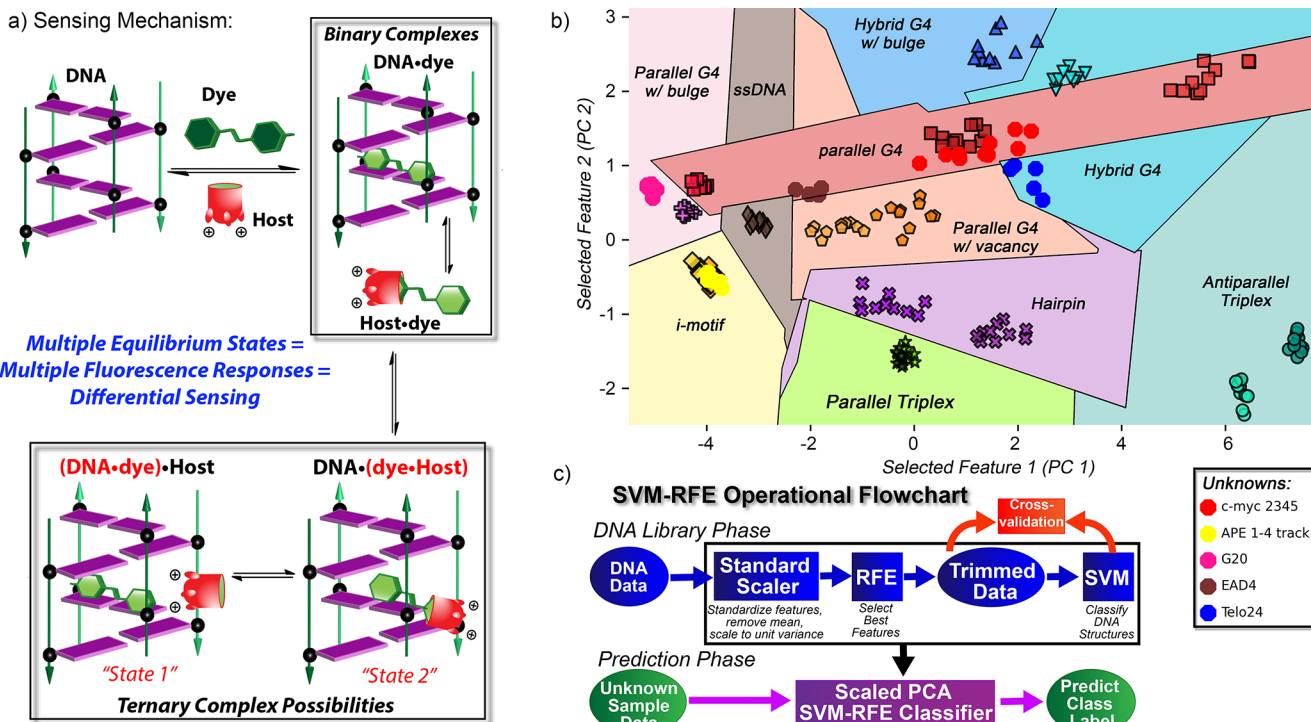


Figure 7. Machine learning optimized sensing. (a) Schematic of the sensing mechanism of the synthetic receptor array using an i-motif as example; (b) SVM-RFE decision region boundary (DRB) plot showing classification of different DNA folding motifs; (c) operational flowchart of the SVM-RFE process.

add the hosts as competitive sensing elements. This allows sensing of unexpected targets, namely, noncanonically folded nucleic acid strands.^{4,66} The anionic cavitand TCC has no affinity for DNA or RNA at all, which is quite obvious. However, the styrylpyridinium dyes are ideally shaped for interaction with DNA: they are rigid, cationic, and slightly kinked, mirroring the general structure of known DNA ligands.⁶⁷ As there are myriad small molecule dyes that can simply “detect” DNA, we focused on a different goal: can arrayed host/guest complexes sense small differences in DNA structure that cannot be simply detected by sequence analysis? Nucleotide strands can form complex 3D structures other than the paradigmatic double helix: examples of noncanonical folds include G-quadruplexes (G4s), Hoogsteen triplets, and hairpins and i-motifs, among others. Despite some notable recent advances,^{68,69} it remains challenging to analyze noncanonical structures from sequences alone and often requires modified nucleotides,^{70–72} X-ray crystallography, or multidimensional NMR spectroscopy for complete structural analysis.

The initial target was a series of DNA G4 strands. G4s are generally classified into three different structural topologies: parallel, antiparallel or hybrid (Figure 6a). The structural differences between the topologies are small, varying in the conformation of the guanine glycosidic angles (*syn* or *anti*) and the relative orientation of the strands. The styrylpyridinium dyes bind G4 DNA strands with micromolar affinity^{4,66} and show enhanced emission while bound. They also show strong affinity for cavitand hosts in water. As such, we combined a series of hosts, TCC, CHI, CHP, and AMI (Figure 2a), and a shallow phosphonate cavitand provided by the Dalcanale group¹⁰ with DSMI and PSMI dyes and monitored the fluorescence changes upon addition of a series of DNA strands. This 10-component array was highly responsive to changes in nucleotide structure.

The potential of the array was shown by focusing on different G4s: 23 different sequences (6 parallel, 8 antiparallel and 9 hybrid, all with lengths of 22–26 bases) could all be fully discriminated based on their folding type. This result was further confirmed by canonical discriminant analysis (Figure 6b), which shows correct classification of 114 out of 115 samples (including repeats) into the three topologies.

While the classification ability of the array is excellent, the discriminatory power is also impressive. The sensor can easily discriminate the parallel G4 structures from the hybrid and antiparallel topologies. Figure 6c focuses on the differentiation between hybrid and antiparallel strands, which both contain one or two of the four G strands in an orientation opposite to the others and display *anti* and *syn* guanines. They are even sometimes considered as a single topology group,⁷³ making their differentiation by simple methods highly challenging. The antiparallel and hybrid G4s that colocalize close to each other are all of similar lengths (22–26 nts) and all display three G-quartet stacks. The antiparallel structures that vary in the number of G-quartet stacks are fully separated from the 3-stack G-quartets (either antiparallel or hybrid). Some impressive discrimination effects can also be seen for the 3-stack G4s (Figure 6c). Although 2KKA has high sequence similarity to 2KF8, AG22, wtTel23, and wtTel24 (varying only by the absence of one T or one A on the 3' or 5' end), it is well separated from the other strands. The limit of detection (LOD) ranges from 3 to 36 nM, and the material usage is low: the minimum total DNA consumption is <12 pmol. The sensor also performs well in the presence of interfering compounds like lysozyme, lactose, and serum and can sense topology switches triggered by substituting K⁺ with Na⁺ in the buffer.

The excellent selectivity shown by the initial array in discriminating G4 structures allowed application to other

folds.⁶⁶ The key to the detection and the differential selectivity is the synergistic combination of *both* host and dye interacting with DNA folded structures. Figure 7a shows a representation of the sensing mechanism, which illustrates the complexity of the system. The dyes all bind to the DNA and show differential emission enhancement when bound. The host-dye complexes can also form ternary complexes with folded DNA. Cationic cavitands can interact with structured DNA while binding the dyes, and this complex interaction adds a level of selectivity that is not possible with the dyes alone. The hosts are essential for selective sensing, even though the dyes are the main reporters.

An improved modified sensor array (hosts CHI, CHP, and AMI and dyes DSMI, PSMI, DQMI, MSMI)⁶⁶ was applied to a series of 18 other nonclassical DNA folded structures, including hairpins, Hoogsteen triplets, and i-motifs, as well as imperfect G4s with bulges or vacancies. These targets ranged from entirely different folding motif structures, such as triplets, to those with very small differences, such as bulges or vacancies in G4 structure. Interrogating 18 DNAs with 16 array elements established a multidimensional data set with 1440 data points. This large and multidimensional data set illustrates the power and the limitations of the array-based analysis: it is not simple to quantify the classification effect, nor easy to pick out the most effective components from such a large and complex data set. This is where machine learning approaches can be exploited, to select the most optimal array elements, evaluate the classification effect, and even predict unknown DNA folding.

The fluorescence output from the array was analyzed by the support vector machine (SVM) algorithm,^{74,75} followed by a PCA step to minimize data dimensions, and visualized in a 2D SVM decision region boundary (DRB) plot (Figure 7b,c). Training the SVM-RFE algorithm with the data from the known DNA pool can lead to a classification model that permits the use of the fluorescence responses from an “unknown” DNA to predict its folding motif. To test this hypothesis, we chose 4 DNA strands with known folding motif as “unknown” targets: *c-myc* 2345 and *EAD4* (known to form a parallel G4 structure), *APE 1–4 track* (an i-motif), and *Telo24* (a hybrid G4). The resultant PC values were then projected to the original DRB plot by the classifiers resulting from running scaled 2D PCA-coupled SVM-RFE on the training data set. The location of the unknown DNA (shown as solid blocks in Figure 7b) shows successful prediction: all of the 10 repeats for each of the “unknown” targets located within the boundary for their corresponding folding. This is remarkable, because the DNA pool contained structures with high similarity, including parallel vs hybrid G4s and parallel G4s with bulges or vacancies that differ by only one base.

OUTLOOK

The outlook for biosensing with synthetic receptors is very exciting. The last four decades (or so) have seen a steady progression in the “difficulty” of application: the initial focus was on molecular recognition in organic solvents, then progressed to function in water and biological media. The complexity of targets has increased along with it, from the pioneering work on cation recognition, to anions and small neutral molecules and then to truly biologically important species such as amino acids, peptides, and nucleotides. Water-soluble deep cavitands are an enticing structural motif for biosensing, as they are high affinity hosts for important biological targets, can be easily tuned, and exploit multiple different recognition mechanisms for differential sensing of diverse targets in the presence of different

targets. As well as the targets described here, we have used cavitand/dye combinations to detect other relevant molecules, including protein isoforms,⁷⁶ biothiols,⁷⁷ and hydrophobic species such as steroids,⁷⁸ cannabinoids,⁷⁹ or insect pheromones.⁸⁰

There are a couple of important “next steps” for synthetic receptor-based biosensing. Differential sensing is a powerful tool, and can take advantage of Big Data to use machine learning and AI approaches to truly achieve the selectivity and sensitivity of antibody-based sensing. An important task is to selectively recognize and confirm target identity in a highly complex mixture of similar components, rather than detecting individual standards in pure solutions. This will require a combination of synthetic organic chemistry and statistical analysis to provide a large suite of host and dye combinations and the computing power to pick out essential elements for optimized sensing. Also, as has been elegantly described by Hof,⁵⁰ an important task is to move away from pure water as a medium and to focus on biorelevant conditions for recognition, including high salt concentration and the presence of lipid aggregates (membranes and liposomes) and other competitive species, as well as on function in biomedia such as urine, saliva, serum, cells, and living organisms. Some elegant work has been performed to illustrate the difficulties in molecular recognition in these types of media but also the opportunities.^{81,82} Obviously, as the application medium becomes more challenging, choice of receptor tends to revert to those with the most well understood mechanisms of action, such as calixarenes and cucurbiturils. Flexible deep cavitands are far more variable and complex in their behavior: this has allowed us to study a wide range of targets and show powerful selectivity, but we have only just started to apply them in complex media: we have shown that selectivity persists in serum,⁴ urine,^{78,79} and saliva⁷⁹ and applied them in human cells,⁴⁷ but happily, there is still much to be done.

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Notes

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Biographies

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Richard J. Hooley obtained his undergraduate degrees in Natural Sciences from Emmanuel College, Cambridge University, and his Ph.D. in the laboratory of Prof. Martin F. Semmelhack at Princeton. Following a postdoctoral position in the lab of Prof. Julius Rebek, Jr., at The Scripps Research Institute in La Jolla, CA, he began his independent career in the Department of Chemistry at the University of California—Riverside in 2008 and was promoted to full Professor in 2018. The Hooley group works on various aspects of organic, inorganic, and supramolecular chemistry, with a focus on the synthesis and applications of biomimetic self-assembled cages and hosts, and their functions in supramolecular catalysis, molecular recognition, and biosensing.

ACKNOWLEDGMENTS

The authors thank the National Science Foundation (CHE-1707347) for support.

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