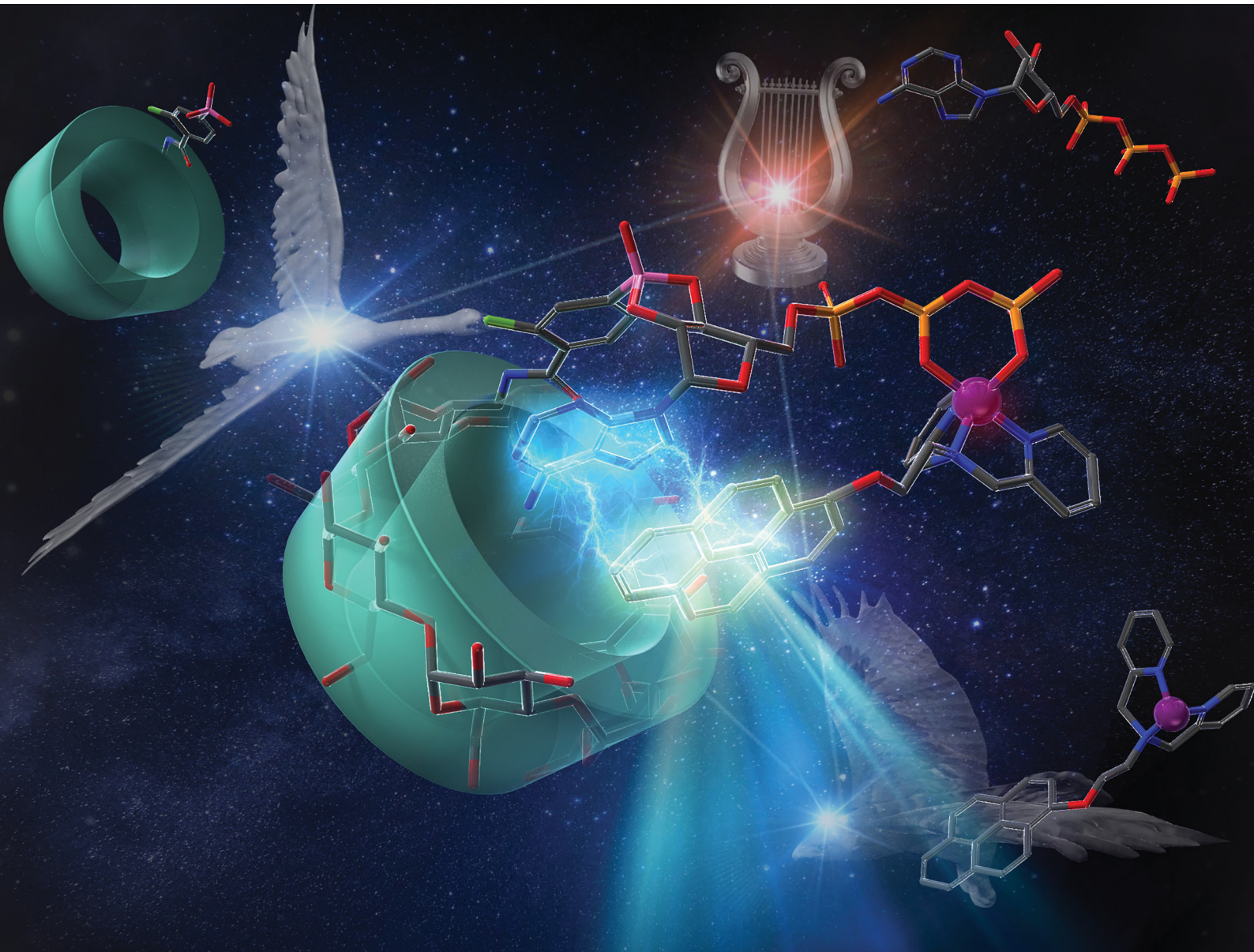


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PAPER

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Selective ATP recognition by boronic acid-appended cyclodextrin and a fluorescent probe supramolecular complex in water[†]

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Various organic compound-based chemosensors for adenosine triphosphate (ATP) have been reported; however, the conventional designs have problems with their water solubility, selectivity, and sensitivity. Herein, a supramolecular complex of boronic acid-appended γ -cyclodextrin and zinc-dipicolylamine-based pyrene probe (**Zn-1/FB γ CyD**) was designed based on the concept of a “cyclodextrin (**CyD**)-based supramolecular chemosensor”. As a proof-of-concept, we have demonstrated that this complex exhibits a turn-on fluorescence response to ATP in water at physiological pH with excellent selectivity and sensitivity, owing to the efficient self-assembly of **Zn-1/FB γ CyD** with ATP through synergistic interactions to afford a strongly fluorescent product. Remarkably, **Zn-1/FB γ CyD** demonstrated extremely strong affinity for ATP with the conditional formation constant of $(7.71 \pm 0.09) \times 10^6 \text{ M}^{-1}$, resulting in the low limit of detection of 18.9 nM. Furthermore, **Zn-1/FB γ CyD** can monitor changes in ATP concentration after the addition of apyrase, hydrolysing ATP to adenosine monophosphate and a phosphate ion. We believe that the presented approach of a **CyD**-based supramolecular chemosensor will propose a novel design that realises unexplored selectivity in the research field of fluorescent molecular recognition.

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Introduction

The field of fluorescence recognition for application to the fluorometric sensing of biologically important analytes has advanced tremendously.^{1,2} In particular, chemosensors for adenosine triphosphate (ATP) have attracted significant attention given the various important roles that ATP plays in biological processes as the energy currency of cells;³ for example, metabolism,⁴ cellular signalling,⁵ and protein synthesis.⁶ The chemosensing approach for the detection of ATP has several advantages, including ease of use, low cost, and high accuracy, overcoming the drawbacks of the conventional luciferin–luciferase assay for ATP sensing, which has low stability and accuracy and requires cumbersome operations.^{7,8}

The strategic design of ATP-selective chemosensors is mainly based on multitopic receptors that capture several recognisable moieties of ATP, namely, the ribose, triphosphate, and adenine

units.^{3,9,10} Despite extensive developments in this research area, most previously designed chemosensors are based on organic scaffolds and are sometimes poorly soluble in water requiring 10 vol% or more organic solvent.^{11–14} Moreover, designing highly selective ATP chemosensors has been challenging because of structural similarities among phosphate derivatives including polyphosphates and nucleotides.^{15,16} Selectivity is an important issue for applications in biological research; for example, chemosensors that allow for the discrimination of ATP from adenosine diphosphate (ADP), adenosine monophosphate (AMP), and the pyrophosphate ion (PPi) are useful in gaining a better understanding of cellular processes.^{17,18} However, conventional strategies require careful organic synthesis and given the difficulty in attaining high selectivity for ATP, reports of ideal chemosensors that exhibit a turn-on response to only ATP have been scarce.

We previously presented a concept of multidentate recognition by water-soluble supramolecular chemosensors based on binding unit-appended cyclodextrins (**CyDs**) with fluorescent and colorimetric probes.^{19–22} As the assembly behaviour of **CyD** is highly sensitive to the structures of the components, the **CyD**-based supramolecular chemosensors display a variety of responses depending on the analyte structure: this feature is advantageous because it provides excellent selectivity for the target analyte (Fig. 1a). Based on these previous studies, we report

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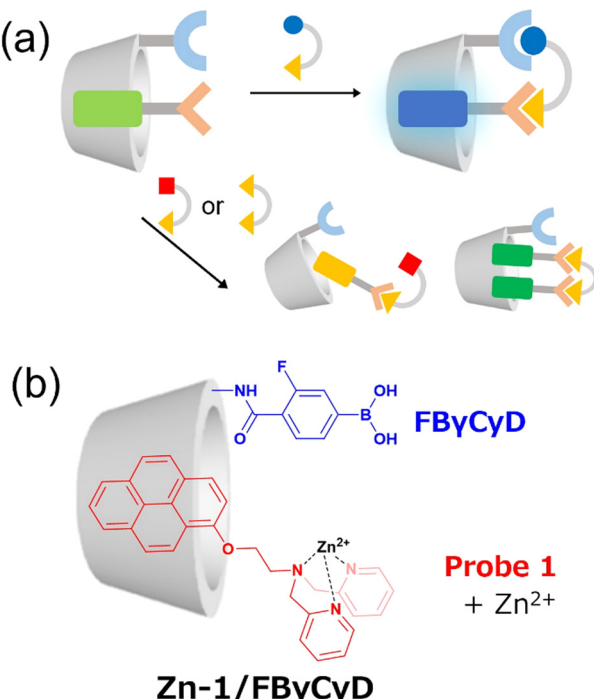


Fig. 1 (a) Cartoon representation of the concept of CyD-based supramolecular chemosensors based on a modified CyD with fluorescent probes. (b) Structure of Zn-1/FByCyD.

a supramolecular complex (Zn-1/FByCyD) of fluorophenylboronic acid-appended γ CyD (FByCyD) and a Zn(II)-dipicolylamine (Zn-dpa)-based fluorescent probe (Zn-1) as a proof-of-concept to design a multitopic CyD-based supramolecular chemosensor for ATP showing turn-on fluorescence response (Fig. 1b). FByCyD has one fluorophenylboronic acid moiety introduced at the C3 position of one 1,4-glycopyranoside unit on the secondary rim. The boronic acid and Zn-dpa moieties capture the ribose and triphosphate units of ATP, respectively.^{11,17} The electron-withdrawing fluorine group is introduced to increase the Lewis acidity of the boron centre; boronic acids with high acidity react with 1,2-diol compounds efficiently.²³ The Zn-1/FByCyD complex displays specific fluorescence enhancement in response to only ATP.

Experimental

Reagents

All reagents were purchased from commercial sources and used as received without further purification. The detailed information is summarised in the ESI.[†]

Instruments

All the nuclear magnetic resonance (NMR) spectra were measured by using a JEOL JNM-ECA 500 spectrometer (JEOL, Japan) at room temperature. For ¹H and ¹³C NMR spectroscopy, the solvent signals were used as ref. 24 The high resolution electrospray ionization mass spectra (HRMS-ESI) were recorded in methanol using a JEOL Accu-TOF JMS T100LC (JEOL, Japan). The high resolution fast atom bombardment mass spectra

(HRMS-FAB) were recorded using a MStation JMS-700 (JEOL, Japan). Solution pH was measured by using a HORIBA pH meter F-52 (Horiba, Japan). The UV-vis absorption spectra were measured using a Hitachi U-3900H absorption spectrophotometer (Hitachi, Japan) equipped with a temperature controller (Hitachi, Japan) to keep the temperature constant at 25 °C. The fluorescence spectra were measured using a Hitachi F-7000 fluorescence spectrophotometer (Hitachi, Japan) and a Hitachi F-7100 fluorescence spectrophotometer (Hitachi, Japan) equipped with a temperature controller (Hitachi, Japan) and an EYELA CCA-1111 (EYELA, Japan) to keep the temperature constant at 25 °C. The induced circular dichroism (ICD) spectra were measured using a JASCO J-820 spectrophotometer (JASCO, Japan) equipped with a Peltier temperature controller (JASCO, Japan) and an EYELA Cool Ace CA-1111 (EYELA, Japan) to keep the temperature constant at 25 °C under a nitrogen atmosphere.

Preparation of sample solutions

All sample solutions were prepared with Milli-Q water and spectroscopic-grade dimethyl sulfoxide (DMSO) using acid-washed glassware. Unless otherwise noted, a mixed solvent system of DMSO/water (1/99 in v/v) was used. The pH of the solutions for spectroscopic measurements was adjusted to 7.4 with diluted aq. HCl and aq. NaOH in the presence of 5 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) buffer. Appropriate amounts of DMSO and the following stock solutions: probe 1 DMSO solution (10 mM), an aq. zinc(II) ion (4 mM), aq. FByCyD (5 mM), a pH buffer solution (50 mM), and an aq. phosphate derivative, were added into a volumetric flask to adjust the concentrations for each experimental condition. Basically, the sample solutions contain 10 μ M of 1, 10 μ M of Zn²⁺, 0.1 or 0.5 mM of FByCyD, and 5 mM of HEPES in DMSO/water (1/99 in v/v). Before the measurements of the fluorescence and UV-vis absorption spectra, all the sample solutions were vortexed for 10 seconds.

Determination of acid dissociation constant

The absorbance of FByCyD at a specific wavelength was measured and analysed using the KaleidaGraph program based on the acid dissociation model of monobasic acids.

Determination of inclusion constants

Fluorescence intensity changes ($F - F_0$) of Zn-1 at various concentrations of cyclodextrin compounds were analysed using KaleidaGraph program to determine the inclusion constants.

Determination of conditional equilibrium constants

Multi-wavelength data (360–600 nm) of the fluorescence spectra at various concentrations of phosphate derivatives were analysed using the ReactLab EQUILIBRIA program to determine the conditional equilibrium constants.²⁵

Monitoring of the change in ATP concentrations

Fluorescence intensity changes at 386 nm were monitored for 1200 seconds with stirring using an Hitachi F-7100 fluorescence spectrophotometer after the apyrase addition (100 mU,



50 mU, and 10 mU) into **Zn-1/FByCyD-ATP** solution. During these measurements, the temperature was kept constant at 25 °C.

Results and discussion

Probe **1** and modified **CyDs** were readily synthesised (see ESI† and Fig. S1–S4). **Zn-1** was prepared by the addition of zinc nitrate (10 µM) to an equimolar amount of **1** ($C_1 = 10 \mu\text{M}$) in a mixed solvent of DMSO and water (1/99 in vol.). The acid dissociation constant (pK_a) of the boronic acid moiety was determined to be 7.92 ± 0.02 from the pH-dependence of the UV-vis absorption spectra of **FByCyD** (Fig. S5 and S6, ESI†), indicating that 23% of **FByCyD** is present as the conjugate tetrahedral boronate anion at pH 7.4 (Scheme S1, ESI†).²³

The fluorescence intensity of **Zn-1** was enhanced by increasing the concentration of coexisting **FByCyD** (Fig. S7, ESI†), suggesting that the supramolecular encapsulation of **Zn-1** by **FByCyD** suppresses the non-radiative decay of the excited state.²⁶ Although the γ **CyD** cavity forms a pyrene dimer that exhibits dimeric fluorescence at approximately 500 nm,²⁷ this was not observed for the supramolecular complex of **FByCyD** with **Zn-1**. This finding suggests that **FByCyD** and **Zn-1** form a supramolecular complex in a 1:1 stoichiometric ratio, *i.e.*, **Zn-1/FByCyD**. The changes in the fluorescence intensity of **Zn-1** at various **FByCyD** concentrations were well fitted to the 1:1 stoichiometric inclusion model, with an inclusion constant of $(3.08 \pm 0.08) \times 10^5 \text{ M}^{-1}$ (Fig. S7, ESI†). In contrast, the fluorescence enhancement observed with the addition of **FByCyD** was not apparent when native γ **CyD** was added to **Zn-1** (Fig. S8, ESI†); the inclusion constant was much smaller at $(4.25 \pm 0.73) \times 10^3 \text{ M}^{-1}$ (Fig. S9, ESI†). These findings suggest that the incorporated boronic acid moiety of **FByCyD** facilitates the tight supramolecular complexation of **Zn-1/FByCyD** via electrostatic attraction between the positive **Zn-dpa** moiety of **Zn-1** and the negative boronate species of **FByCyD** partially present in the solution.

To determine the selectivity of **Zn-1/FByCyD** for ATP, the UV-vis absorption and fluorescence spectra of **Zn-1/FByCyD** were measured in the absence and presence of various phosphate derivatives: phosphate ion (Pi), PPi, triphosphate ion (Tri), AMP, ADP, and ATP (Chart S1, ESI†). Whereas no differences in the UV-vis absorption spectra were observed (Fig. S10, ESI†), the fluorescence spectra varied depending on the added phosphate derivatives (Fig. 2). In the absence of the phosphate derivatives, **Zn-1/FByCyD** exhibited a vibronic-structured fluorescence band in the 370 to 450 nm region, which is a characteristic spectral pattern of pyrene compounds.²⁸ The addition of the inorganic polyphosphates (PPi and Tri) resulted in a clear reduction in fluorescence intensity. The PPi moiety efficiently coordinates to the Zn(II) centres of two **Zn-dpa** moieties to form a bis-dentate complex, *i.e.*, PPi-bis[Zn-dpa].¹⁵ As shown in the normalised spectra of Fig. 2a, the dimeric fluorescence of the pyrene moieties was clearly observed at approximately 500 nm in the presence of PPi and Tri (Fig. S11, ESI†). Thus, the decrease in fluorescence is attributable to the

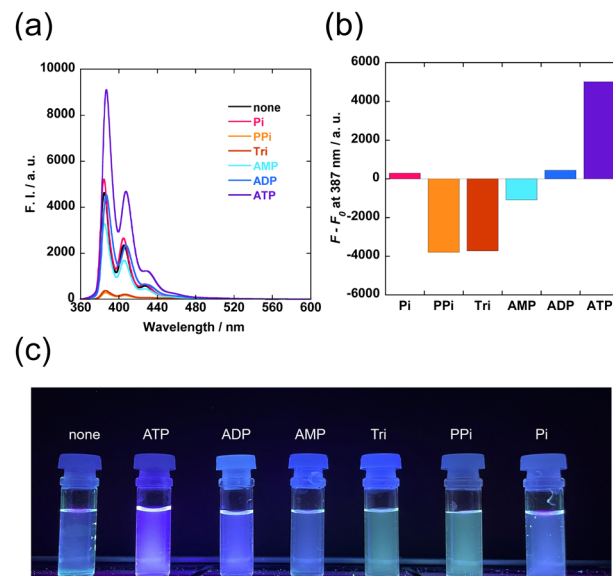


Fig. 2 The fluorescence spectra of **Zn-1/FByCyD** (a) and fluorescence enhancement ($F - F_0$) at 387 nm of **Zn-1/FByCyD** (b) in the absence and presence of various 1.0 mM phosphate derivatives in DMSO/water (1/99 in vol.): $C_1 = 10 \mu\text{M}$, $C_{\text{Zn}} = 10 \mu\text{M}$, $C_{\text{FByCyD}} = 0.5 \text{ mM}$, $C_{\text{HEPES}} = 5 \text{ mM}$, pH 7.4, $T = 25^\circ\text{C}$, and $\lambda_{\text{ex}} = 350 \text{ nm}$. Photographs of sample solutions with 365 nm UV irradiation at room temperature (c).

formation of the inclusion complex of the **Zn-1** dimer with **FByCyD** and the inorganic polyphosphates, PPi- and Tri-(**Zn-1**)₂/ **FByCyD** (Chart S2, ESI†), which quenches the monomeric fluorescence. Pi, AMP, and ADP caused almost no changes in the fluorescence spectra of **Zn-1/FByCyD**, suggesting that these phosphate derivatives inefficiently react with **Zn-1/FByCyD**. Surprisingly, **Zn-1/FByCyD** exhibited a fluorescence turn-on response to only ATP, demonstrating its excellent selectivity for ATP: this supramolecular complex can discriminate between ATP and other typical interfering phosphate derivatives, such as PPi and ADP, which give indistinguishable fluorescence spectra in many reported systems.¹⁵ Notably, the change in fluorescence colour was visible to the naked eye (Fig. 2c).

The fluorescence spectra of **Zn-1/FByCyD** were measured at various concentrations of the phosphate derivatives (Fig. 3 and Fig. S12–S17, ESI†). As shown in Fig. 2, the fluorescence enhancement of **Zn-1/FByCyD** was observed only with the addition of ATP. The change in fluorescence intensity reached saturation when the total concentration of ATP was 10 µM (the same as **Zn-1** concentration): this result signifies that **Zn-1/FByCyD** quantitatively reacts with ATP. The conditional equilibrium constant was determined to be $(7.71 \pm 0.09) \times 10^6 \text{ M}^{-1}$ according to the 1:1 binding model (Fig. S18, ESI†). The extremely large affinity gave rise to the excellent sensitivity to ATP, with a low limit of detection of 18.9 nM (Fig. S19, ESI†). For PPi and Tri, spectral changes were well fitted to the 2:1 binding model (probe: phosphate), supporting that **Zn-1** forms dimers with PPi and Tri inside the **FByCyD** cavity (Fig. S20 and S21, ESI†).

As competitive experiments, the fluorescence spectra of **Zn-1/FByCyD** with ATP (**Zn-1/FByCyD-ATP**) were measured in

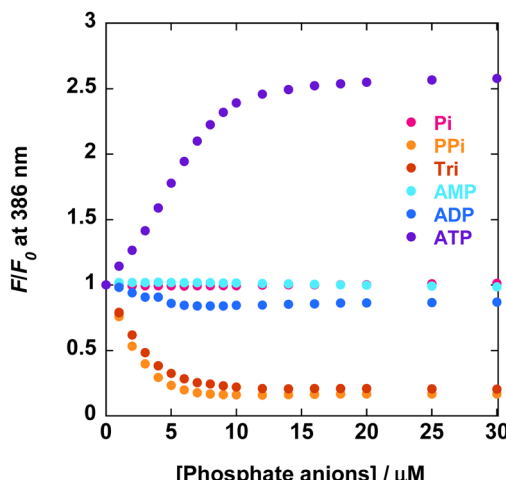


Fig. 3 Fluorescence intensity ratios (F/F_0) at 386 nm of **Zn-1/FByCyD** at various phosphate concentrations in DMSO/water (1/99 in vol.): $C_1 = 10 \mu\text{M}$, $C_{\text{Zn}} = 10 \mu\text{M}$, $C_{\text{FB}\gamma\text{CyD}} = 0.1 \text{ mM}$, $C_{\text{HEPES}} = 5 \text{ mM}$, pH 7.4, $T = 25^\circ\text{C}$, and $\lambda_{\text{ex}} = 350 \text{ nm}$.

the presence of the other phosphate derivatives as competitors (Fig. S22, ESI[†]). The fluorescence intensity of **Zn-1/FByCyD-ATP** remained unchanged by the addition of AMP and ADP, demonstrating the high selectivity of the complex for ATP even in the presence of other adenine nucleotides. Although the additions of PPi and Tri decreased the fluorescence intensity of **Zn-1/FByCyD-ATP**, the turn-on response of **Zn-1/FByCyD** to ATP was partially preserved. Given that ATP is present in large amounts among phosphate derivatives in both intracellular and extracellular environments,²⁹ the presence of competitors should not significantly affect the sensitivity to ATP. Taken together, **Zn-1/FByCyD** is potentially useful in practical situations where ATP measurement is performed in the presence of other phosphate derivatives.

To clarify the role of the boronic acid moiety of **FByCyD**, the fluorescence spectra of **Zn-1** with phosphate derivatives were measured in the absence of **FByCyD** (Fig. S23, ESI[†]) and the presence of native γCyD (Fig. S24, ESI[†]) and phenyl-appended γCyD (**PhyCyD**, Fig. S25, ESI[†]). In all the systems, fluorescence enhancement was observed in response to all phosphate derivatives except Pi, suggesting that **Zn-1/γCyD**, **Zn-1/PhyCyD**, or **Zn-1** alone shows no sensitivity to ATP. The observed fluorescence enhancement is likely due to the suppression of the photo-induced electron transfer quenching process by the complexation of the **Zn-dpa** moiety with the phosphate moieties, which decreases the cationic character of the zinc centre.¹⁵ These results clearly show that the boronic acid moiety incorporated into **FByCyD** plays an essential role in improving the selectivity for ATP, plausibly due to the binding of the boronic acid moiety to the 1,2-diol moiety of ATP. Therefore, the observed ATP selectivity of **Zn-1/FByCyD** is surely originated from the multiple bindings with ATP. In other words, **Zn-1/FByCyD** possesses binding sites that are structurally unsuitable for the multidentate recognition of AMP and ADP, and **FByCyD** facilitates the formation of PPi- and

Tri-(Zn-1)₂/FByCyD with PPi and Tri, respectively, which results in the quenching of the monomer fluorescence. Such dimer formation does not occur in the **Zn-1/γCyD** and **Zn-1/PhyCyD** systems because γCyD and **PhyCyD** do not tightly encapsulate **Zn-1**. As mentioned, multidentate recognition by the **Zn-dpa** moiety and the boronic acid moiety of **Zn-1/FByCyD** was suggested in binding to ATP. As controls, the fluorescence spectra of **Zn-1/γCyD** and **Zn-1/PhyCyD** were measured in the presence of ATP (Fig. S26, ESI[†]). Both supramolecular complexes showed much weaker fluorescence than **Zn-1/FByCyD-ATP**. This suggests that the boronic acid moiety rigidifies the pyrene fluorophore through multidentate recognition and therefore supports the expected multidentate recognition.

The supramolecular complex structures of **CyD** with a pyrene compound vary depending on the ring size of **CyD**; pyrene compounds form 2:1 and 1:1 (pyrene:**CyD**) stoichiometric supramolecular complexes with γCyD and βCyD , respectively.²⁷ Therefore, the effect of **CyD** ring size on selectivity was also investigated by using native βCyD and fluorophenylboronic acid-appended βCyD (**FBβCyD**) instead of **FByCyD** (Chart S3, ESI[†]). The corresponding βCyD -based supramolecular complexes, *i.e.*, **Zn-1/βCyD** (Fig. S27, ESI[†]) and **Zn-1/FBβCyD** (Fig. S28, ESI[†]), showed similar trends with **Zn-1** alone, **Zn-1/γCyD**, and **Zn-1/PhyCyD**, exhibiting fluorescence enhancement in response to all phosphate derivatives except Pi. This finding indicates that the signaling mechanisms of **Zn-1/βCyD** and **Zn-1/FBβCyD** are essentially the same as **Zn-1** alone, **Zn-1/γCyD**, and **Zn-1/PhyCyD**, clearly signifying that the **FByCyD** host provides overwhelmingly superior ATP selectivity compared with other **CyD** hosts.

The induced circular dichroism (ICD) spectra of **Zn-1/FByCyD** were measured in the absence and presence of phosphate derivatives to further extract structural information (Fig. S29, ESI[†]). The supramolecular complexation of **FByCyD** with **Zn-1** led to a strongly positive Cotton effect at 350 nm. This result empirically demonstrates that **Zn-1** is encapsulated by the cavity of **FByCyD**.³⁰ When the phosphate derivatives having pyrophosphate moieties (PPi, Tri, and ATP) were present, the intensity of ellipticity at 350 nm was markedly decreased. This implies that the pyrophosphate moieties coordinate to the **Zn-dpa** moiety of **Zn-1/FByCyD**.

The selectivity of **Zn-1/FByCyD** for ATP was further examined by screening other nucleotides (Chart S1, ESI[†]) including guanosine triphosphate (GTP), cytidine triphosphate (CTP), and uridine triphosphate (UTP). The additions of CTP and UTP attenuated the fluorescence intensity, whereas the addition of GTP enhanced the fluorescence intensity (Fig. 4). These results suggest that the purine bases play a vital role in triggering the turn-on response of **Zn-1/FByCyD**. The adenine moiety of ATP may form π - π stacking with the pyrene moiety of the **Zn-1** unit in the **CyD** cavity to further rigidify the **Zn-1** unit, given that **CyD** cavities encapsulate the adenine moiety in water.^{31,32} As **Zn-1/FByCyD-ATP** emits stronger fluorescence than **Zn-1/FByCyD-GTP**, the assembly with ATP is expected to be more efficient than with GTP.

The obtained results indicate that **Zn-1** and **FByCyD** form a rigid supramolecular complex that features three-point recognition



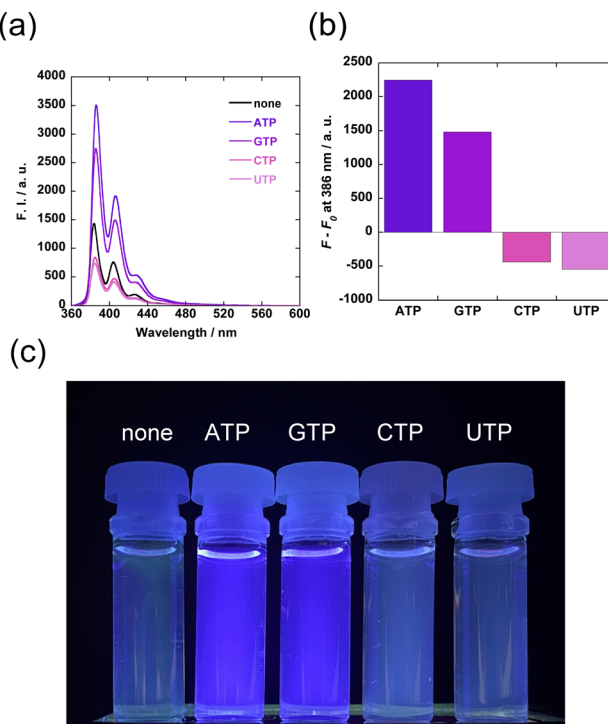


Fig. 4 The fluorescence spectra of **Zn-1/FB γ CyD** (a) and changes in the fluorescence intensity ($F - F_0$) at 386 nm of **Zn-1/FB γ CyD** (b) in the absence and presence of 1 mM of various nucleotides in DMSO/water (1/99 in vol.): $C_1 = 10 \mu\text{M}$, $C_{\text{Zn}} = 10 \mu\text{M}$, $C_{\text{FB}\gamma\text{CyD}} = 0.1 \text{ mM}$, $C_{\text{HEPES}} = 5 \text{ mM}$, pH 7.4, $T = 25^\circ\text{C}$, and $\lambda_{\text{ex}} = 350 \text{ nm}$. Photographs of the sample solutions with 365 nm UV irradiation at room temperature (c).

including (1) esterification of the boronic acid and ribose moieties, (2) complexation of the **Zn-dpa** and triphosphate moieties, and (3) π - π stacking between the pyrene and adenine moieties (Fig. 5). On the contrary, the additions of PPI and Tri result in fluorescence reduction, as the formation of the **Zn-1** dimer leads to quenching of the monomeric fluorescence. For the recognition of AMP and ADP, the binding sites of **Zn-1/FB γ CyD** are structurally mismatched; hence, the excellent selectivity for ATP is due to the various responses of the **CyD**-based supramolecular chemosensor to different analyte structures.

Zn-1/FB γ CyD was evaluated for potential application to the monitoring of ATP concentrations. Apyrase was added to a solution containing **Zn-1/FB γ CyD** and ATP (30 μM) in order to hydrolyse ATP *via* conversion into ADP and AMP, as follows: (1) ATP \rightarrow ADP + Pi, and (2) ADP \rightarrow AMP + Pi.³³ Changes in fluorescence intensity at 386 nm were monitored during this hydrolysis reaction (Fig. 6). **Zn-1/FB γ CyD** was robust to apyrase under these experimental conditions (Fig. S30, ESI[†]). After the addition of apyrase, the fluorescence intensity of **Zn-1/FB γ CyD**-ATP at 386 nm gradually decreased with time. The rate of hydrolysis of ATP increased with increasing apyrase concentration; this clearly shows that ATP is hydrolysed by apyrase. After the hydrolysis reaction was completed, the fluorescence of **Zn-1/FB γ CyD** was recovered (Fig. S31 and S32, ESI[†]). These results suggest that ATP is completely hydrolysed to AMP and Pi without any influence on the **Zn-1/FB γ CyD** fluorescence spectra

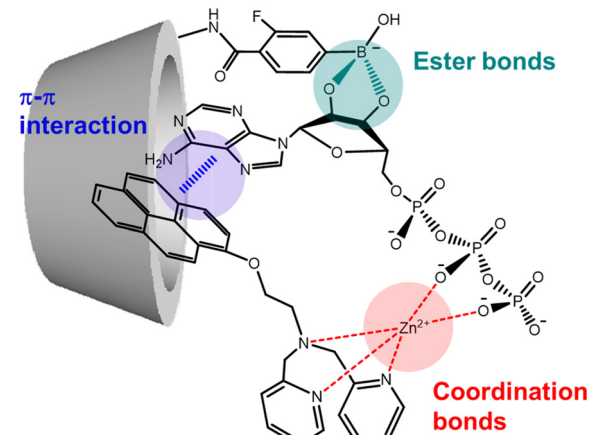


Fig. 5 Plausible structure of **Zn-1/FB γ CyD**-ATP.

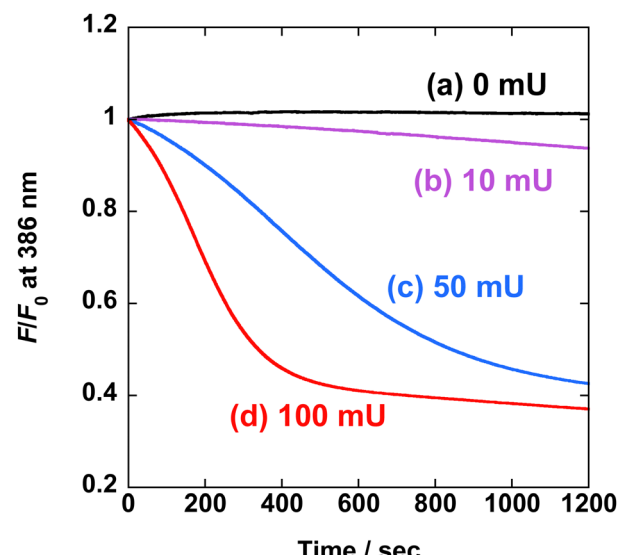


Fig. 6 Changes in relative fluorescence intensity (F/F_0) at 386 nm of **Zn-1/FB γ CyD** with ATP (30 μM) in DMSO/water (1/99 in vol.) following the addition of apyrase: (a) 0 mU, (b) 10 mU, (c) 50 mU, and (d) 100 mU: $C_1 = 10 \mu\text{M}$, $C_{\text{Zn}} = 10 \mu\text{M}$, $C_{\text{FB}\gamma\text{CyD}} = 0.1 \text{ mM}$, $C_{\text{HEPES}} = 5 \text{ mM}$, pH 7.4, $T = 25^\circ\text{C}$, and $\lambda_{\text{ex}} = 350 \text{ nm}$.

(Fig. 2). Consequently, **Zn-1/FB γ CyD** allows for the monitoring of changes in ATP concentration with time. This feature is potentially advantageous when applied to biological studies to examine cellular processes relating to ATP production and consumption.

Conclusions

We have designed a **CyD**-based supramolecular chemosensor, **Zn-1/FB γ CyD**, which selectively recognises ATP and allows for the monitoring of changes in ATP concentration with time. Its excellent selectivity is due to the efficient assembly of **Zn-1/FB γ CyD** with ATP, which is facilitated by synergistic multi-recognition to form a fluorescent supramolecular product.

Surprisingly, **Zn-1/FByCyD** demonstrated extremely strong affinity for ATP, achieving the detection of ATP with nM levels. In other words, we have demonstrated that the concept of a **CyD**-based supramolecular chemosensor has benefits providing various advantageous features compared to simple fluorescent probes, such as water solubility, excellent selectivity, and sensitivity. We believe that **Zn-1/FByCyD** not only has the potential to serve as a basic structure of ATP chemosensing systems but also presents a desirable approach to exploit the concept of a **CyD**-based supramolecular chemosensor, which will be useful in various fields relating to molecular recognition. This presented concept holds promise to pioneer novel chemosensors for unexplored analytes.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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